Article

A Hybrid Microbial–Enzymatic Fuel Cell Cathode Overcomes Enzyme Inactivation Limits in Biological Fuel Cells

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Abstract: The construction of optimized biological fuel cells requires a cathode which combines the longevity of a microbial catalyst with the current density of an enzymatic catalyst. Laccase-secreting fungi were grown directly on the cathode of a biological fuel cell to facilitate the exchange of inactive enzymes with active enzymes, with the goal of extending the lifetime of laccase cathodes. Directly incorporating the laccase-producing fungus at the cathode extends the operational lifetime of laccase cathodes while eliminating the need for frequent replenishment of the electrolyte. The hybrid microbial–enzymatic cathode addresses the issue of enzyme inactivation by using the natural ability of fungi to exchange inactive laccases at the cathode with active laccases. Finally, enzyme adsorption was increased through the use of a functionally graded coating containing an optimized ratio of titanium dioxide nanoparticles and single-walled carbon nanotubes. The hybrid microbial–enzymatic fuel cell combines the higher current density of enzymatic fuel cells with the longevity of microbial fuel cells, and demonstrates the feasibility of a self-regenerating fuel cell in which inactive laccases are continuously exchanged with active laccases.

Keywords: oxygen reduction reaction; multi-functional catalysts for ORR; biocatalytic; biocathodic microbial communities; optimization of catalyst layers and electrode design

1. Introduction

Biological fuel cells (BFCs) use bioderived fuels and catalysts to produce electricity [1]. Most BFCs utilize the oxygen reduction reaction at the cathode, thus BFCs can be categorized by the type of catalyst used at the cathode: (I) metal catalysts, (II) enzymatic catalysts, and (III) microbial catalysts [2–4]. Platinum-group metal catalysts are most common due to their superb catalytic properties, but the high price of the catalysts prevents the large-scale adoption of biological fuel cells [5]. Enzymatic catalysts hold promise to replace metal catalysts, but suffer from short lifetimes and mass transport limitations [3,4,6,7]. Microbial fuel cells rectify the short lifetimes of enzymatic catalysts at the expense of additional mass transport limitations and lower current densities [2,8–10]. The construction of optimized biological fuel cells requires a cathode which combines the longevity of a microbial catalyst with the current density of an enzymatic catalyst [11,12].

Laccases are commonly used enzymatic catalysts in biological fuel cell cathodes because they demonstrate reversible adsorption to carbon electrodes and an ability to directly transfer electrons to carbon nanotubes [13–15]. Laccases catalyze the one-electron oxidation of diverse chemical substrates at a single-copper-containing site near the surface of the protein [16,17]. Concomitantly, laccases catalyze the four-electron reduction of
dioxygen to water [17,18] at a tri-copper site in the interior of the protein, without the production of superoxides or peroxides [19].

Substrate Oxidation Reaction

$$4\text{Substrate}^{\text{RED}} - 4e^- + \text{laccase}^{\text{OX}} \rightarrow \text{laccase}^{\text{RED}} + 4\text{Substrate}^{\text{OX}} + 4H^+$$

Oxygen Reduction Reaction

$$\text{laccase}^{\text{RED}} + 4e^- + O_2 + 4H^+ \rightarrow \text{laccase}^{\text{OX}} + 2H_2O$$

The current body of research establishes that laccase cathodes suffer from current density losses attributable to short enzyme lifetimes, suboptimal enzymatic reaction velocities, enzyme inactivation, and low enzyme adsorption [20–23]. In order to mitigate these deficiencies, attempts have been made to engineer laccases with faster reaction velocities [24–30]. However, few studies have investigated strategies to mitigate current density losses from these other factors [31,32]. In previous work, laccase cathode lifetimes were extended by the periodic exchange of laccase-containing electrolytes without the enzyme-producing microbe in the electrode [22,23]. To advance research in this area, filamentous fungi and rational cathode design can be used to overcome short enzyme lifetimes, enzyme inactivation, and low enzyme adsorption. Fungi secrete laccases to catalyze the breakdown of lignin in the normal metabolism [16,33,34]. Fungi naturally address the problem of laccase inactivation via specialized endocytic pathways which facilitate the absorption of inactive enzymes for breakdown and reuse [21–23,33,34].

This work presents a hybrid microbial–enzymatic cathode leveraging fungal metabolism to overcome short enzyme lifetimes, enzyme inactivation, and low enzyme adsorption associated with pure laccase cathodes (Figure 1). Five different filamentous fungi were screened for their ability to produce extracellular laccases. Three inducers were analyzed to maximize laccase secretion while maintaining the medicinal and/or nutritional value of the resulting fungal biomass. To increase enzyme loading at the cathode, a functionally graded coating was applied to the cathode prior to introducing the fungus. The laccase-secreting fungus was grown directly on the coated cathode and the exchange of inactive enzymes with active enzymes was monitored electrochemically.

Figure 1. Illustration of (a) bare carbon electrode with no fungus incorporated and no functionally graded coating to increase laccase loading, (b) bare carbon electrode with laccase recycling fungus incorporated but no functionally graded coating, and (c) bare carbon electrode with laccase recycling fungus incorporated and an optimized functionally graded coating to increase laccase loading. The black structures represent the carbon electrode and carbon nanotubes (in functionally graded coating), blue circles represent TiO₂ nanoparticles, white circles represent fully active laccase, light grey circles represent partially active laccase, and dark grey circles represent inactive laccase. The orange region indicates the fungal cytoplasm and the dashed line represents the semipermeable membrane of the fungus.

2. Results

2.1. Assessment of Laccase Secretion

The abilities of five different fungi to secrete enzymatically active laccases were compared with and without induction (Figure 2). In the absence of a laccase-inducing substance, all tested fungi produced less than 10 U·mL⁻¹ of laccase. When copper sulfate, a common laccase inducer, was added, all fungi increased the amount of extracellular...
laccase. In 200 mL culture bottles, the Pleurotus ostreatus cultures demonstrated the highest extracellular laccase activity. The genetic differences between the subspecies of P. ostreatus had a notable effect on extracellular laccase production, with the commercial strain P. ostreatus N001 producing the least active extracellular laccases on day 4 and the in-house cultured P. ostreatus White producing the highest.

![Laccase Activity Graph](image)

**Figure 2.** Five different fungi were screened for their ability to produce laccase. Means that do not differ from *P. ostreatus* White grown for 4 days after induction with CuSO₄ by a p-value of 0.05 are annotated with “a”. Means that do not differ from *P. ostreatus* Blue grown for 4 days after induction with CuSO₄ by a p-value of 0.05 are annotated with “b”. Means that do not differ from *P. ostreatus* Blue grown for 2 days after induction with CuSO₄ by a p-value of 0.05 are annotated with “c”. Means that do not differ from *P. ostreatus* Blue measured immediately after induction with CuSO₄ by a p-value of 0.05 are annotated with “d”.

When smaller reaction vessels (20 mL) were compared to larger reaction vessels (200 mL), *Fusarium oxysporum* and *Trametes versicolor* produced extracellular laccases with the highest activity (Figure 3). The high activities of the extracellular laccases produced by *F. oxysporum* are likely due to *F. oxysporum*’s role as a plant pathogen [35,36], while the high activity of the cellular laccases produced by the *T. versicolor* and the *P. ostreatus* strains can be attributed to their roles as lignin-degrading saprotrophs [36,37]. Despite the ecological relevance of laccases to all tested species [36], it remains unclear why smaller culturing vessels significantly improved the activity of the secreted laccases in the *F. oxysporum* and *T. versicolor* cultures, while having less of an influence over *P. ostreatus* cultures. While the mechanism for this is unclear, it may be related to the ability of laccases to degrade phenolic compounds that arrest fungal growth [38]. The 20 mL setup was used for all subsequent studies, as it provided higher laccase activity, enough biomass for analysis, and was more manageable for subsequent trials.
Figure 3. Laccase induction was compared between large and small culture vessels. Means that do not differ from *T. versicolor* induced with CuSO₄ grown in 20 mL reaction vessels by a p-value of 0.05 are annotated with “a”. Means that do not differ from *T. versicolor* induced with CuSO₄ grown in 200 mL reaction vessels by a p-value of 0.05 are annotated with “b”. Means that do not differ from *T. versicolor* induced with water grown in 200 mL reaction vessels by a p-value of 0.05 are annotated with “c”.

2.1.1. Inducing Laccase Secretion with a Biocompatible Inducer

*Trametes versicolor* has medicinal properties that make the biomass a useful byproduct of this biological fuel cell [39]. The use of copper sulfate as an inducer leads to concerns of heavy metal sequestration by *T. versicolor*, precluding its use for medicinal purposes. We explored Tween20 as an alternative laccase inducer, as the detergent causes oxidative stress to the fungus and is non-toxic to humans and animals.

When copper sulfate, Tween20, and a mixture of the two were compared, it was discovered that the mean of extracellular laccase activity was not significantly higher in copper sulfate-induced cultures than in Tween20-induced cultures for any tested fungus (Figure 4). This experiment did not detect any evidence against the use of Tween20 as a laccase inducer that does not contaminate edible or medicinal biomass with heavy metal inducers. No significant differences between biomass were observed between any of the treatments. Despite the absence of heavy metal contamination in Tween20-induced cultures, copper sulfate was chosen as the inducer for subsequent studies because proteomics is needed to provide a deeper insight into the mechanistic differences between crude filtrate and pure laccases in fuel cell cathodes. Tween20 can cause signal loss in proteomic characterization, and contaminates the mass spectrometry instruments used in proteomics.
2.2. Trametes Versicolor Secretome Analysis

The secretome of *T. versicolor* was analyzed for protein composition under copper sulfate induction and compared to an uninduced culture. Of particular interest are laccases, peroxidases and proteases. Laccases facilitate the four-electron reduction of oxygen to water [18], while peroxidases facilitate the two-electron reduction of oxygen to peroxide [17], which can kill the fungus [40] and inactivate functional laccases [41]. Proteases hydrolyze the peptide bonds between amino acids in a protein, rendering the protein inactive and the amino acids usable for metabolism or protein synthesis [42].

Copper sulfate induction raised the laccase composition of the secretome 21%, from 33% to 54% (Figure 5). Additionally, the peroxidase composition of the secretome was decreased by 8%, to 14% of total protein. Proteases were not found in abundance in the uninduced culture, but comprised 1% of the total protein in the copper sulfate-induced culture. The small number of proteases supports our hypothesis that *T. versicolor* is able to break down and recycle inactive laccases in situ, without disturbing active laccases at the electrode surface. In addition to the proteins of interest, glucose oxidase and glycosidase enzymes were detected in the *T. versicolor* secretome. These metabolic enzymes do not prohibit the laccase function at the cathode, but can occupy sites otherwise usable by laccases for oxygen reduction [43].

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**Figure 4.** Three laccase inducers, copper sulfate, Tween20 and a mixture of copper sulfate and Tween20, were compared to assess the viability of harvesting the fungal biomass of *T. versicolor* and *P. ostreatus* for secondary applications. Means that do not differ from *P. ostreatus* N001 induced with CuSO₄ by a p-value of 0.05 are annotated with “a”. Means that do not differ from *P. ostreatus* Blue induced with CuSO₄ by a p-value of 0.05 are annotated with “b”. Means that do not differ from *P. ostreatus* N001 induced with water by a p-value of 0.05 are annotated with “c”.

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2.3. Comparison of Electrode Coatings

The carbon cathodes used in this study do not provide a large enough surface area to demonstrate an increase in current density due to the recycling of laccase at the cathode. We therefore developed a functionally graded cathode coating with the goal of improving enzyme loading on the cathodes, while facilitating the time-dependent change of active laccases adsorbed onto the electrode. The coating was composed of titanium dioxide nanoparticles (TiO$_2$) to increase porosity [44], carbon nanotubes to increase the electrical conductivity of the coating [45], and Nafion to favorably orient laccase adsorption towards the composite coating [46]. Choi et al. showed that adding carbon nanotubes to the Nafion–TiO$_2$ mixture improved response time, sensitivity, and long-term stability in glucose oxidase-based biosensors [47]. Water was used as a negative control to account for any structural changes to the electrodes during the coating process; an open circuit potential of 52mV vs. SCE was measured. An optimized ratio of 0.2% titanium dioxide nanoparticles to 0.7% carbon nanotubes (w:w) was found to increase the open circuit potential from 52mV vs. SCE (water coating) to 584mV vs. SCE (Table 1 and Figure 6).

**Table 1.** Summary of the different coating compositions designed to increase laccase loading at the cathode, their open circuit potentials (mV vs. SCE) and current density at −200 mV vs. SCE (mA·cm$^{-2}$).

| Coating Composition | Open Circuit Potential (mV vs. SCE) | Current Density at −200mV vs. SCE |
|---------------------|------------------------------------|-----------------------------------|
| 0.9% TiO$_2$        | 128.0                              | –68                               |
| 0.7% TiO$_2$–0.2% CNT | 134                                | –152                              |
| 0.45% TiO$_2$–0.45% CNT | 189                                | –196                              |
| 0.2% TiO$_2$–0.7% CNT | 584                                | –216                              |
| 0.9% CNT            | 117                                | –119                              |
| 18MΩ H$_2$O         | 52                                 | –101                              |

Figure 5. Radar plot of the composition of the of *T. versicolor* secretome under standard (blue) and copper sulfate-induced conditions (orange).
Comparing the current densities of laccase electrodes at potentials less than +400 mV vs. SCE quantifies their performance while excluding oxygen transport limitations [23]. The 0.7% TiO₂–0.2% CNT coating had a current density of ~100mA/cm² at +400 mV vs. SCE, 3% higher than a similar setup utilizing pure laccase adsorbed onto a buckypaper cathode [23]. For all other coating compositions, low laccase loading or poor electrical contact between the enzyme and the electrode lowered the open circuit potential to near +100 mV vs. SCE, so the current densities at ~200 mV vs. SCE were reported in Table 1. Table 1 and Figure 6 show a trend whereby increasing the weight percent of CNT in the CNT–TiO₂–Nafion coating raised the OCP except for when the coating was composed entirely of CNT, TiO₂ or Nafion.

At lower carbon nanotube loading, the number of direct electron transfer sites accessible to laccases is insufficient to promote efficient oxygen reduction, resulting in a lower open circuit potential and low current density (Figure 6). However, at higher CNT loading, the nanotubes form aggregates, significantly diminishing the effective surface area of the coating [48–50]. This phenomenon is observable in the linear sweep voltammograms, wherein a coating composed entirely of CNT has nearly the same open circuit potential as one composed entirely of TiO₂. In these cases, the nanostructures appear to have increased the porosity of the coating, but have not created a significant number of sites where laccase can directly transfer electrons to the cathode. Equal amounts of CNT and TiO₂ increased the OCP to 250 mV, but the best performance was from 0.7% CNT and 0.2% TiO₂.
Figure 7. Scanning electron micrograph of the optimized coating with *T. versicolor* growing on top of the coating.

A scanning electron micrograph of the optimized coating taken 4 days after inoculation is depicted in Figure 7. The large ~10μm structures are the fibrous, highly porous TiO2–CNT–Nafion coating [51], and the smaller ~5μm structures are the fungal hyphae [52]. The larger coating structures spatially distribute the fungal hyphae, while the nanoscale components promote enzyme exchange between the fungus and the electrode, and facilitate high enzyme loading with electrical contact to the electrode.

### 2.4. Comparison of Electrochemical Properties with and Without Enzyme Regeneration

The current density of pure laccase in sterile electrolyte without fungus was 20% higher than that of the crude laccase from the *T. versicolor* culture without fungus (Figure 8a). This discrepancy is supported by the proteomics data (Figure 5), which confirm the presence of electrochemically inert enzymes in the crude filtrate. Inert enzymes such as glucose oxidase and manganese peroxidase can occupy direct electron transfer sites that are otherwise used by laccase to produce current [43]. Incorporating the fungus at the cathode lowered the open circuit potential of the pure laccase treatment by 24 mV (Figure 8b). The presence of the fungus raised the current density and open circuit potential of the crude laccase treatment to ~52 mA/cm² and 524 mV, respectively. The higher performance of crude filtrate over pure laccase is in agreement with the findings of Sane et al. (2013) [23], and could be associated with the preservation of posttranslational protein modifications, such as glycosylation [37], or the secretion of uncharacterized mediators [53], which may be lost in the protein purification process [54].
Forty-eight hours after introducing electrolyte to the fuel cells, linear sweep voltammetry was performed again. In cultures without *T. versicolor* (Figure 8c), the open circuit potential and current density were significantly diminished. However, in cultures with *T. versicolor* growing on the cathode, current density and open circuit potential were aligned (Figure 8d). Interestingly, the performance of the media control, containing no laccase at initiation, was “rescued” by the presence of the fungus. The higher bioelectrocatalytic activity of the electrodes indicates that the active laccases produced by the fungus are successfully exchanged with the inactive laccases previously adsorbed onto the cathode.

3. Discussion

Biological fuel cells continue to gain recognition as a solution to global energy challenges due to their multifunctionality [2]. However, neither enzymatic nor microbial fuel cells have achieved commercial viability due to their short lifetimes and low current densities [31,32,55,56]. The hybrid microbial-enzymatic fuel cell combines the higher current density of enzymatic fuel cells with the longevity of microbial fuel cells. Additionally, the hybrid microbial-enzymatic fuel cell demonstrates the feasibility of a self-regenerating fuel cell cathode, in which an enzyme-producing microbe is
incorporated into the electrode by continuously recycling inactive laccases to active laccases.

In comparison to the recently presented microbial cathode, in which *T. versicolor* was grown in the catholyte, our approach is advantageous, as it incorporates the fungus at the cathode, precluding the need for mediators, which have a limited lifetime [22,57]. This proof-in-concept work demonstrated the feasibility of using unpurified fungal culture broth as an electrolyte in a laccase-based cathode. Directly incorporating the laccase-producing fungus at the cathode overcomes the limitations imposed by the short lifetime of laccases, while eliminating the need for the frequent replenishment of the electrolyte. However, the current density at +400 mV vs. SCE was −112 mA/cm², 13% lower than a laccase fuel cell without fungus, which utilized crude laccase adsorbed onto a buckypaper cathode, as operated by Sane et al. [23]. The hybrid microbial–enzymatic cathode addresses the issue of enzyme inactivation by using the natural ability of fungi to exchange inactive laccases with active laccases, at the cost of slightly lower current density.

The production of edible and medicinal biomass was used as the metric to analyze the secondary benefits of this system. *T. versicolor* has shown promise in boosting the immune response in cancer patients, and increasing lymphocyte counts, natural killer cell functional activity, CD8+ T cell counts, and CD19+ B cell counts [58]. These benefits have come from taking capsules of freeze-dried fungal biomass produced for consumption. *Pleurotus ostreatus* is a gourmet mushroom with a highly sought-after taste and high mineral content [59,60]. The increasing adoption of plant-based proteins provides a growing market for *P. ostreatus* biomass [61,62], which is produced in the hybrid microbial–enzymatic cathode. The harvesting of fungal biomass for edible and medicinal purposes represents a new area of high-profit secondary benefits [63,64] that should be explored as an avenue to amend the functionality of microbial fuel cells. Temperature and pH impact laccase activity [65–68] and fungal biomass production [35–37]. Once a fungal byproduct is decided upon, variable pH and temperature studies should be carried out to optimize the production of electricity and biomass for the intended application. Fungi are also adept at heavy metal sequestration and waste remediation, which presents opportunities to customize and combine secondary benefits based on application [69,70].

The various ecological roles played by filamentous fungi in nature allow the hybrid microbial–enzymatic fuel cell to be tuned to provide a variety of secondary and tertiary benefits during operation. Future research will explore the use of additional fungi, and the genetic and cultural conditions that can increase the laccase composition of the secretome in the fungi of interest, as well as reducing the number of detrimental (peroxidase) and metabolic (glucose oxidase and glycosidase) enzymes in the extracellular environment.

4. Materials and Methods

Unless otherwise stated, all chemicals were of reagent grade and purchased from Sigma Aldrich (St. Louis, MO, USA). Bacto yeast extract was obtained from Fisher Scientific (San Diego, CA, USA); *T. versicolor* laccase standard (13.6 U/mg) was purchased from MyBioSource (Cerritos, CA, USA). *P. ostreatus* var. pearl, *P. ostreatus* var. blue, and *T. versicolor* were purchased from Fungi Perfecti (Olympia, WA, USA); *P. ostreatus* var. N001 was purchased from American Type Culture Collection (http://www.atcc.org//ps/32783.ashx); *F. oxysporum* f.s. lycopersici was obtained from Dr. Hans VanEtten (The University of Arizona, Tucson, AZ, USA). The 500 mL media bottles and 50 mL miniature bioreactors were purchased from Dow Corning (Midland, MI, USA); single-wall carbon nanotubes (95 wt.%; OD: 10–20 nm; ID: 5–10 nm; L: 10–30 μm) were provided by Dr. Mike Foley (CTI Materials, Grafton, VT, USA). Spectroscopic carbon rods were purchased from Ted Pella Inc (Redding, CA, USA). PTFE heat shrink tubing was purchased from McMaster Carr Inc. (Elmhurst, IL, USA). Difco yeast extract was purchased from Fisher Scientific (Waltham, MA, USA).
All fungi were maintained on 100 mm × 15 mm petri dishes containing glucose yeast extract agar (GYEA) composed of 1% glucose, 1% yeast extract, and 1.5% agar. In total, 10 mL of acidified glucose yeast extract broth at pH 4.5 (1% glucose, 1% yeast extract, 1% succinic acid) was dispensed onto a 7-day-old culture to create pre-inoculum. The mycelium was dislodged from the plate using a 10 mL wide-bore pipette and the resulting slurry was added to 190 mL of AGYEB in a sterile 500 mL reagent bottle. The reagent bottles were kept shaking at 250 rpm and 27 °C for 7 days. The resulting inoculum was used at 10% volume-to-volume ratio to inoculate all experimental cultures. Experimental culture conditions were as follows: 500 mL media bottles and 50 mL bioreactors were filled with 200 mL and 20 mL AGYEB, respectively. Cultures were either kept shaking at 250 rpm or 0 rpm. All fungi were grown at 27 °C for the predetermined time course. After 72 h of growth an inducer was added aseptically to promote laccase secretion. After the appropriate time course, cultures were filtered through pre-weighed cheesecloth, then cell-free filtrate was collected using 0.45 μm low-protein binding cellulose acetate filters. After straining, the fungal biomass and the cheesecloth were placed into beakers, frozen at −80 °C, and lyophilized under a 200 mitorr vacuum at −60 °C. Dry biomass was determined by weighing the biomass–cheesecloth combination and subtracting the weight of the cheesecloth.

Laccase activity in the cell-free filtrate was determined spectrophotometrically using a Bio Tek Synergy II microplate reader with microinjector [71]. Total protein was quantified using the Bradford assay [72]. Briefly, one-part cell-free filtrate was used to one-part Bradford reagent, and absorbance was monitored at 595 nm after 10 min. Then, 100 μL of sample, standard, or blank was pipetted into the corresponding well, and 100 μL of 2 mM ABTS (2,2-azino-bis(3-ethylbenzthiazoline-6-sulphonic acid)) was injected into each well of the plate and absorbance was read at 410 nm continuously after injection. The reaction was allowed to proceed for 10 min at 27 °C under the medium shaking condition. Data reduction was performed according to the standard fluorometric protein quantification parameters in the Gen5 software (Version 2.09, BioTek, Winooski, VT, USA, 2012) provided with the reader. ANOVA and a multiple comparison mean tests were performed in MATLAB 2020b (Mathworks, Natick, MA, USA) according using Sidák’s multiple comparisons test for one way ANOVA (Figures 2 and 4) and two-stage linear step-up procedure of Benjamini, Krieger and Yekutieli for two-way ANOVA (Figure 3) [73].

Cell-free filtrates were also prepared for proteomics, then passed through 0.22 μm low-protein binding syringe filters (Fisher Scientific, San Diego, CA, USA). The resulting solution was ultrafiltered using an Amicon stirred cell concentrator (MerckMillipore, Temecula, CA, USA) [74]. After ultrafiltration, the concentrated protein solution was dialyzed against 50 mM ammonium bicarbonate pH 7.5. The dialysates were frozen until they were sent to the University of Arizona’s proteomics core facility. Tandem mass spec analysis was performed using a LTQ Orbitrap Velos mass spectrometer (Fisher Scientific, Waltham, MA, USA) equipped with a nano electrospray ion source; the instrument was operated in data-dependent acquisition mode. Raw data were processed using Scaffold Proteome Software (Version 3.6.5, Proteome Software Inc., Portland, OR, USA, 2012).

The tops of the spectroscopic carbon rods were passivated using PTFE heat shrink tubing ( McMaster Carr, Cleveland, OH, USA). The surface area was normalized to 10 cm² after modification with PTFE. Dip coating solutions of 0.9 wt.% were created with varying amounts of titanium dioxide and CNT and cured at 120 °C for three minutes. Single-walled carbon nanotubes (SWCNTs) have been used in the past to impart electronic conductivity to fuel cells, sensors, and bio-electronics [75,76]. The dimensions of SWCNTs facilitate the direct physical connections between the electrode and enzyme while the inherent conductivity of SWCNTs establishes an electrical connection [45]. Though other carbon nanostructures are more sustainable in terms of cost and environmental toxicity [77–79], SWCNTs were selected to facilitate direct electron transfer between the electrode and enzymatic catalyst by decreasing the electron tunneling distance [12].
Titanium dioxide nanoparticles were synthesized by adding tetramethylammonium hydroxide to titanium isopropoxide in a 1:1 ratio, then 3 parts 18MΩ deionized water was added, and the mixture was stirred at 95 °C for 2 h, then the solution was peptized at 70 °C for 48 h [80]. The resulting solution was passed through a 0.2 μm filter and 1 mL aliquots were placed in a roto evaporator to determine the final concentration of nanoparticles. The stock solution was diluted to 20 wt.% using 18 MΩ deionized water, and the resulting working solution was stored at 2 °C. Electrodes were dipped into the same solution three times; a constant pull rate of 2 mm/s was used for all coatings. The electrodes were then dip coated in 0.1% Nafton under sterile conditions at the same pull rate. For microbe-containing coatings, functionalized electrodes were placed on 1 mm × 3 mm round carbon supports in the 20 mL bioreactors and fungi were allowed to colonize for two days before electrochemical measurements were taken.

All electrochemical measurements were carried out on a Biologic VSP potentiostat (Biologic, Seyssinet-Pariset, Grenoble, France) with a saturated calomel reference electrode (Gamry Instruments, Warminster, PA, USA) and a 10 cm × 0.5 cm carbon counter electrode (Sigma Aldrich, St. Louis, MO, USA). Functionalized 0.32 cm × 10 cm spectroscopic carbon rods were used as working electrodes. The catholyte was stirred for 10 min under air before open circuit potential was determined and linear sweep voltammetry began. Catholytes were either sterile AGYEB (media), crude *T. versicolor* filtrate (crude), or sterile AGYEB spiked with purified *T. versicolor* laccase (pure). Sweeps were started at −200mV vs. SCE and run to the open circuit potential of the electrode at a scan rate of 50 mV/s.

5. Conclusions

Laccase-secreting fungi were grown directly on the cathode of a biological fuel cell constructed with a multifunctional cathode coating of titanium dioxide nanoparticles and carbon nanotubes. Culturing studies show that smaller reaction vessels increased the amount of laccase produced by most of the tested species. In *T. versicolor*, copper sulfate increased the laccase proportion of the secretome by 21% compared to the negative control. Electrochemical experiments demonstrated improvements in enzyme loading and cell longevity when the fungus was grown on the cathode. The ten-fold increase in open circuit potential demonstrated the ability of the optimized cathode coating to increase enzyme loading compared to the other tested coating compositions. The initial current density was higher without the fungus, but the presence of the fungus increased both the current density and the open circuit potential of the laccase cathodes at 48 h of continuous operation. The hybrid microbial–enzymatic fuel cell combines the higher current density of enzymatic fuel cells with the longevity of microbial fuel cells, and demonstrates the feasibility of a self-regenerating fuel cell cathode in which inactive laccases are continuously exchanged with active laccases.

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