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Full Communication

Direct electron transfer-type bioelectrocatalysis by membrane-bound aldehyde dehydrogenase from Gluconobacter oxydans and cyanide effects on its bioelectrocatalytic properties

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ABSTRACT

The bioelectrocatalytic properties of membrane-bound aldehyde dehydrogenase (AlDH) from Gluconobacter oxydans NBRC12528 were evaluated. AlDH exhibited direct electron transfer (DET)-type bioelectrocatalytic activity for acetaldehyde oxidation at several kinds of electrodes. The kinetic and thermodynamic parameters for bioelectrocatalytic acetaldehyde oxidation were estimated based on the partially random orientation model. Moreover, at the multi-walled carbon nanotube-modified electrode, the coordination of CN to AlDH switched the direction of the DET-type bioelectrocatalysis to acetate reduction under acidic conditions. These phenomena were discussed from a thermodynamic viewpoint.

1. Introduction

Acetic acid bacteria have a variety of membrane-bound dehydrogenases that work in their unique metabolism such as acetic acid fermentation [1]. The application of these bacteria in bioelectrochemical devices (biosensors, biofuel cells, biosupercapacitors, etc.) is significant for industry [1]. Specifically, membrane-bound dehydrogenases such as fructose dehydrogenase (FDH) [2], alcohol dehydrogenase (ADH) [3-5], gluconate dehydrogenase (GaDH) [6], and lactate dehydrogenase (LDH) [7] are known to directly communicate with electrodes and proceed electro-enzymatic reactions, which is called direct electron transfer (DET)-type bioelectrocatalysis. It is suggested that these dehydrogenases transfer electrons from their substrates to the electrodes via their catalytic centers and one or more hemes c in this order [1].

Herein, we focus on another membrane-bound dehydrogenase of acetic acid bacteria, namely, aldehyde dehydrogenase (AlDH). AlDH from Gluconobacter oxydans NBRC12528 is a heterotrimeric enzyme composed of a large subunit (86 kDa) containing a molybdopterin cofactor (Moco) as a catalytic center, a membrane-bound cytochrome c subunit (55 kDa) containing three hemes c, and a small subunit (the molecular mass of which is unknown) containing an iron-sulfur cluster [8,9]. In vivo, AlDH oxidizes acetaldehyde, and the extracted electrons are transferred to ubiquinone in the inner membrane [9]. However, the electrochemical properties and the bioelectrocatalytic performance of AlDH have not yet been examined.

In the present study, we report DET-type bioelectrocatalysis by AlDH at planar gold (Au), 2-mercaptoethanol (ME)-functionalized Au, and multi-walled carbon nanotube (MWCNT)-modified glassy carbon (GC) electrodes. Planar and ME-functionalized Au electrodes were used as platforms for DET-type bioelectrocatalysis by FDH [2,10]. MWCNTs are nanostructured electrode materials that promote various DET-type reactions [11] with the following effects: 1) the enlarged effective surface area of the electrode increases the amount of adsorbed enzymes [12]; 2) the curvature of the mesoporous electrode structure increases the probability of enzyme orientations suitable for DET-type reactions [13,14]; 3) the electric field strengthened by the expansion of the electric double layer accelerates the kinetics of heterogeneous electron transfer at the edge of the micropores [15].

The effects of the coordination of cyanide ions (CN ) to the hemes c on the bioelectrocatalytic properties of AlDH are also examined herein. Interestingly, CN -coordinated AlDH exhibits DET-type activity for acetate reduction.

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2. Experimental

2.1. Materials and chemicals

Protein markers for sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) were obtained from Nacalai Tesque Inc. (Japan). Water-dispersed MWCNTs were kindly donated by Nitta Co. (Japan). Other reagents were purchased from Wako Pure Chemical Industries, Ltd. (Japan). All solutions were prepared using ultrapure water.

2.2. Purification of AlDH

AlDH was purified from a variant of *Gluconobacter oxydans* NBRC12528 (∆aldhA::Km® [16]) according to the procedure in the literature [8], with minor modifications. Briefly, the process of removing ADH from the membrane fraction using multiple surfactants was omitted because the gene of ADH was knocked out. The purified enzyme solution contained 0.5% Triton X-100 as a solubilizer and 25 mM benzaldehyde and 10% sucrose as stabilizers. The results of SDS-PAGE analysis are shown in Fig. S1, which demonstrate that the subunit structure of AlDH from the variant was identical to the reported structure of that purified from the wild-type cells [8].

The AlDH activity was spectrophotometrically measured using potassium ferricyanide as an electron acceptor and ferric sulfate-Dupanol reagent [8]. Here, one unit (U) of AlDH activity is defined as the amount of the enzyme that oxidizes 1 μmol of acetaldehyde per minute at pH 4.0. The protein concentration was estimated using a DC protein assay kit (Bio-Rad, USA) with bovine serum albumin as the standard. The specific activity (approximately 600 U mg⁻¹) was slightly better than that described in the literature (430 U mg⁻¹ [8]), plausibly due to the shortening of the purification process, as described above.

2.3. Electrode preparation

Au and GC electrodes (3 mm in diameter, BAS, Japan) were polished with 1.0 μm and 0.05 μm alumina powder. The electrodes were subsequently rinsed by sonication in distilled water. Self-assembled monolayers of 2-mercaptoethanol (ME) were then formed on the Au electrodes by immersing the Au electrodes in an ethanol solution containing 2 mM ME for 2 h. Subsequently, the ME-functionalized Au electrodes were washed with ethanol and distilled water. MWCNT-modified GC electrodes were also prepared by applying 60 μL of 0.1 wt% MWCNT dispersion to the surface of each GC electrode, followed by drying at 70 °C.

2.4. Electrochemical measurements

All electrochemical measurements were performed at 25 °C with electrochemical analyzers (CV-50 W (BAS, USA) or ALS 701E (ALS Co. Ltd., Japan)). Anaerobic measurements were carried out in a nitrogen (N₂) chamber filled with a mixture of 96% N₂ and 4% H₂. A platinum wire and a homemade Ag/AgCl|sat. KCl electrode were used as the counter and reference electrodes, respectively. In this study, all potentials are referred to the reference electrode.

3. Results and discussion

3.1. DET-type bioelectrocatalytic acetaldehyde oxidation by AlDH

Fig. 1 shows cyclic voltammograms (CVs) recorded at planar and ME-functionalized Au electrodes in a McIlvaine buffer (McB) (pH 4.0) containing 0.1 M acetaldehyde under Ar atmospheric conditions. After adding AlDH at a final concentration of 2 μg mL⁻¹ into the electrolysis solution, clear sigmoidal waves were observed at both electrodes (Fig. 1A). As shown in Fig. S2, the anodic current density gradually increased with time and finally reached the maximum value. Thus, these waves are ascribed to DET-type bioelectrocatalysis of acetaldehyde oxidation by AlDH physically adsorbed on the electrodes. The bioelectrocatalytic current density reached approximately 20 μA cm⁻² at the planar Au electrode and 30 μA cm⁻² at the ME-functionalized Au electrode at 0.4 V. On the other hand, the slightest anodic current was also observed in the absence of acetaldehyde (Fig. S3), which seems to be due to the DET-type oxidation of benzaldehyde contained in an enzyme solution.

In this study, kinetic and thermodynamic analysis of the DET-type bioelectrocatalytic waves at the AlDH-modified electrodes was performed using a partially random orientation model in which it was assumed that the enzymes were adsorbed on the electrode surface in a homogeneously distributed orientation [17,18]. In the model, the steady-state current density (j) is expressed by the following equation [17]:

\[
  j = \frac{j_{\text{cat}}}{\beta \Delta d (1 + \eta^2)} \ln \left( \frac{k_c (1 + \eta)}{k_c (1 + \eta) + \eta^a \exp(-\beta \Delta d) + \eta^a} \right)
\]

where \( j_{\text{cat}} \) is the limited steady-state catalytic current density, \( k_c^{\text{max}} \) is...
finally disappeared (Fig. S4). The results can be explained by the following mechanisms:

1) The coordination of CN⁻ to the hemes c in AlDH and the adsorption of AlDH on an electrode proceed simultaneously, where the former is slower than the latter because the dissociation of a proton from HCN (pKₐ ~ 9.2) is unfavorable under acidic conditions. (Caution: addition of KCN to acidic solutions must be performed with good ventilation as the process generates HCN).

2) Intramolecular electron transfer in AlDH is inhibited by CN⁻ coordination. This means that the electrons are transferred from the catalytic center to the electrode through one or more hemes c in the native enzyme.

Fig. 2 presents the schematics of the electron transfer pathway of AlDH. The approximated values of the formal potential at pH 4.0 were evaluated based on the following considerations: a) the E°' value of one of hemes c obtained in this work, b) the formal potential of the acetate/acetalddehyde redox couple (E°') is −0.78 V at pH 7.0 [23] and shifts by −89 mV pH⁻¹ based on the following and the Nernst equations:

$$\text{CH}_3\text{COO}^- + 3\text{H}^+ + 2e^- \rightleftharpoons \text{CH}_3\text{CHO} + \text{H}_2\text{O} \quad (4)$$

3) The formal potential of the Moco (E°'Moco) in AlDH is assumed to be close to that of xanthine oxidase (−0.5 V [24]), which is a typical molybdenum enzyme. The involvement of other electron-mediating redox centers was ignored here. The downhill property is well explained by electron transfer from acetaldehyde to the electrode-active heme c at pH 4.0. However, the CN⁻ coordination to the heme c results in an uphill barrier in the intramolecular electron transfer process, which inhibits the enzyme activity for acetaldehyde oxidation (Fig. 2A).

At lower pH (pH 2.5), E°' shifts toward positive potential and becomes more positive than E°'Moco. Therefore, it was expected that electron transfer from the CN⁻-coordinated heme c to acetate via Moco would be thermodynamically favorable, as shown in Fig. 2B. This hypothesis was tested by using a MWCNT-modified GC electrode as a platform for AlDH because the electrode provided a relatively smaller background current probably ascribed to proton reduction under strongly acidic conditions, compared to the planar Au and GC electrodes. In addition, DET-type bioelectrocatalytic acetaldehyde oxidation was observed at the AlDH-adsorbed MWCNT-modified GC electrodes, with a larger current density than that observed at AlDH-modified Au electrodes (Fig. S5). Since the AlDH solution at pH 2.5 kept the activity for acetaldehyde oxidation, there was no fatal change in the enzyme conformation under acidic conditions.

Fig. 3 shows the CVs recorded at the MWCNT-modified GC electrodes in an MeCl (pH 2.5) containing 0.1 M acetic acid and 1 mM KCN under anaerobic conditions. After addition of AlDH at a final concentration of 2 µg mL⁻¹ into the electrolysis solution, the cathodic DET-type bioelectrocatalytic current for acetate reduction by CN⁻-coordinated AlDH was observed. An increase in the current was only observed in the presence of AlDH, KCN, and acetate. Here, it is noteworthy that the current density increased over many hours, as demonstrated by the multi-scanned CVs in Fig. S6. The current decrease in the first 2 h is plausibly due to adsorption of AlDH on the electrodes, and CN⁻ was gradually coordinated to AlDH, which seemed to proceed very slowly, as described above. Only the current decrease was observed in the absence of KCN (Fig. S7), which also supports the above process. In addition, the onset potential (approximately −0.6 V) is somewhat more negative than that expected from the formal potential of the CN⁻-coordinated electrode-active heme c. The potential shift induced by CN⁻ coordination might be larger than 0.4 V. However, DET-type acetate reduction cannot be observed in the presence of high concentrations (≈mM) of acetaldehyde (data not shown), which is suggested to be due to the biased catalytic property and the product inhibition.

### 4. Conclusions

DET-type bioelectrocatalysis of acetaldehyde oxidation by AlDH
from *Gluconobacter oxydans* at several types of electrodes was examined, and the kinetics and thermodynamics of the bioelectrocatalytic reaction were evaluated. The effects of CN⁻ on the DET-type bioelectrocatalytic activity of AlDH were also evaluated, and the reverse DET-type bioelectrocatalytic activity of CN⁻-coordinated AlDH, i.e., acetal reduction, was observed under strongly acidic conditions. Further studies are required for detailed clarification of the characteristics of bioelectrocatalysis by AlDH. The reaction can be applied to detecting acetaldehyde in food and environment analyses, utilizing acetaldehyde as a biofuel, and producing acetic acid from acetaldehyde without any additional redox mediators.

**CRediT authorship contribution statement**

**Taiki Adachi:** Investigation, Conceptualization, Writing - original draft, Visualization. **Yuki Kitazumi:** Conceptualization. **Osamu Shirai:** Conceptualization, Writing - review & editing. **Kenji Kano:** Conceptualization, Writing - review & editing, Supervision, Project administration.

**Declaration of Competing Interest**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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**Appendix A. Supplementary data**

Supplementary data to this article can be found online at https://doi.org/10.1016/j.tecom.2020.106911.

**References**


**Fig. 2.** Proposed potential profiles for acetaldehyde oxidation at pH 4.0 (A) and acetate reduction at pH 2.5 (B) in DET-type bioelectrocatalysis by AlDH. Heme c indicates the electrode-active heme c.

**Fig. 3.** CVs for acetate reduction at the AlDH-adsorbed MWCNT-modified GC electrode in an McE (pH 2.5) containing 0.1 M acetic acid and 1 mM KCN in the presence (solid line) and absence (dotted line) of 2 μg mL⁻¹ AlDH at 25 °C under quiescent and anaerobic conditions, at v = 10 mV s⁻¹.


