Conversion of levoglucosan into glucose by the coordination of four enzymes through oxidation, elimination, hydration, and reduction

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Levoglucosan (LG) is an anhydrosugar produced through glucan pyrolysis and is widely found in nature. We previously isolated an LG-utilizing thermophile, *Bacillus smithii* S-2701M, and suggested that this bacterium may have a metabolic pathway from LG to glucose, initiated by LG dehydrogenase (LGDH). Here, we completely elucidated the metabolic pathway of LG involving three novel enzymes in addition to LGDH. In the S-2701M genome, three genes expected to be involved in the LG metabolism were found in the vicinity of the LGDH gene locus. These four genes including LGDH gene (*lgdA, lgdB1, lgdB2, and lgdC*) were expressed in *Escherichia coli* and purified to obtain functional recombinant proteins. Thin layer chromatography analyses of the reactions with the combination of the four enzymes elucidated the following metabolic pathway: *LgdA* (LGDH) catalyzes 3-dehydrogenation of LG to produce 3-keto-LG, which undergoes β-elimination of 3-keto-LG by *LgdB1*, followed by hydration to produce 3-keto-d-glucose by *LgdB2*; next, *LgdC* reduces 3-keto-d-glucose to glucose. This sequential reaction mechanism resembles that proposed for an enzyme belonging to glycoside hydrolase family 4, and results in the observational hydrolysis of LG into glucose with coordination of the four enzymes.

Monosaccharide anhydrides, such as levoglucosan (1,6-anhydro-β-d-glucopyranoside; LG), mannosan, and galactosan, are generated from the burning of cellulose and hemicellulose1,2. As the burning of biomass is a major source of particle matter, these anhydrosugars are expected to be indicators of air pollution3. It has been estimated that 90 million metric tons of anhydrosugars are produced every year, with the majority being LG (> 90%)4. As LG is also formed by pyrolysis of starch, food industries that process carbohydrates have the potential to produce LG as a by-product. We recently found that LG was produced during the industrial production process for novel water-soluble indigestible polysaccharide (resistant glucan) developed by our group, and isolated an LG-utilizing thermophile, *Bacillus smithii* S-2701M, which we used to construct an LG-removing bioprocess5.

Two types of microbial metabolisms of LG have been reported to date. The eukaryotic LG metabolic pathway, in fungi and yeasts, starts with LG phosphorylation with the cleavage of the 1,6-linkage by LG kinase to produce glucose 6-phosphate, which is then metabolized through the glycolytic pathway6,7. With respect to the bacterial metabolic pathway of LG, NAD-dependent LG dehydrogenase (LGDH) was found in *Arthrobacter* sp. I-5528. Recently, a gene encoding LGDH was found in the genome of *Pseudoarthrobacter phenanthenivorans* based on the partial amino acid sequences of the LGDH in *Arthrobacter* sp. I-552, and the crystal structure of LGDH was determined9. The biochemical and structural analyses of LGDH confirmed the C3-specific oxidation of LG that

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was completely converted to glucose with all four Lgd proteins (Fig. 2). However, lower level of the conversion with the crude extract, the reaction was conducted with the four recombinant Lgd proteins in various combinations. The growth on LG with NAD+, Mn²⁺ were consistent with the molecular weight expected from the corresponding nucleotide sequences. We estimated molecular masses of LgdA, LgdB₁, LgdB₂, and LgdC were 44, 30, 36, and 43 kDa, respectively, which agreed with those in a soluble fraction. Sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) indicated that the recombinant LgdA (BSM4216_1279), LgdB₁ (BSM4216_1280), and LgdB₂ were deleted in the same region in the genome of the B. smithii strain, which was consistent with the fact that DSM 4216 could not utilize anhydrosugars such as LG and cellobiosan in our previous study. However, no biochemical and genetic analyses have been performed to understand the reactions beyond the dehydrogenation of LG by LGDH in the speculated pathway.

Our previous study revealed that B. smithii S-2701M also possesses a thermophilic LGDH, and its glucose-forming activity was detected using the LGDH active fractions in an ion-exchange chromatography in a manner similar to that observed in Arthrobacter sp. 1-552. In the present study, we aimed to elucidate the LG metabolic pathway in S-2701M using the recombinant proteins including LGDH, which are expected to be involved in the pathway.

**Results**

**B. smithii S-2701M genomic region is involved in LG metabolism.** Based on the nucleotide and amino acid sequences of LGDH from P. phenanthrenivorans, the putative gene encoding LGDH was found in the genome of B. smithii S-2701M. Around the putative LGDH gene (lgdA), three genes encoding enzymes (lgdB₁, lgdB₂, and lgdC) were found (Fig. 1). The gene product of lgdB₁ was annotated as a sugar phosphate isomerase/epimerase and the closest relative was that from Paenarthrobacter nicotinovorans with 71% similarity (GenBank ID: GAT87749.1). The gene product of lgdB₂ was also annotated as sugar phosphate isomerase, but showed relatively lower similarity (51%) to the closest relative from P. nicotinovorans (GenBank ID: GAT89806.1). Intriguingly, these closest relatives were encoded by the different gene loci in the genome of B. smithii and S-2701M. The annotations of a-f in S-2701M genome are as follows: a, 2-haloalkanoic acid dehalogenase; b, LacI family transcriptional regulator; c, sugar ABC transporter substrate-binding protein; d, sugar ABC transporter ATP-binding protein; e, ABC transporter permease; f, ABC transporter permease. The genes g (BSM4216_1279), h (BSM4216_1280), and i (BSM4216_1281) in the DSM 4216 genome are similar to the genes a, lgdC, lgdB₂, respectively, in the S-2701M genome. The DSM 4216 genome lacks regions from the gene b to 5-prime of lgdC in the S-2701M genome.

Figure 1. Organizations of the gene clusters involved in LG metabolism in the genomes of two B. smithii strains. The annotations of a-f in S-2701M genome are as follows: a, 2-haloalkanoic acid dehalogenase; b, LacI family transcriptional regulator; c, sugar ABC transporter substrate-binding protein; d, sugar ABC transporter ATP-binding protein; e, ABC transporter permease; f, ABC transporter permease. The genes g (BSM4216_1279), h (BSM4216_1280), and i (BSM4216_1281) in the DSM 4216 genome are similar to the genes a, lgdC, lgdB₂, respectively, in the S-2701M genome. The DSM 4216 genome lacks regions from the gene b to 5-prime of lgdC in the S-2701M genome.

produces 3-keto-LG. This finding led us to depict the possible LG metabolic pathway initiated by the LGDH reaction, followed by hydrolysis of 3-keto-LG and reduction of the hydrolysate to produce glucose. However, no biochemical and genetic analyses have been performed to understand the reactions beyond the dehydrogenation of LG by LGDH in the speculated pathway. Based on the biochemical investigation described below, we suggested that LgdC catalyzed the reduction of 3-keto-LG to glucose in the speculated LG metabolic pathway as described above.

Results

**B. smithii S-2701M genomic region is involved in LG metabolism.** Based on the nucleotide and amino acid sequences of LGDH from P. phenanthrenivorans, the putative gene encoding LGDH was found in the genome of B. smithii S-2701M. Around the putative LGDH gene (lgdA), three genes encoding enzymes (lgdB₁, lgdB₂, and lgdC) were found (Fig. 1). The gene product of lgdB₁ was annotated as a sugar phosphate isomerase/epimerase and the closest relative was that from Paenarthrobacter nicotinovorans with 71% similarity (GenBank ID: GAT87749.1). The gene product of lgdB₂ was also annotated as sugar phosphate isomerase, but showed relatively lower similarity (51%) to the closest relative from P. nicotinovorans (GenBank ID: GAT89806.1). Intriguingly, these closest relatives were encoded by the different gene loci in the genome of P. phenanthrenivorans genome, and there were no marked similarities between the amino acid sequences of lgdB₁ and lgdB₂, in spite of the same annotation. The gene product of lgdC was annotated as a Gfo/Idh/MocA family oxidoreductase. Gfo and Idh refer to glucose-fructose oxidoreductase and inositol 2-dehydrogenase, respectively, and MocA catalyzes a dehydrogenase reaction involved in the catabolism rhizopine (3-O-methyl-scyll-o-inosamine). Prior to commencing the biochemical investigation described below, we suggested that LgdC catalyzed the reduction of 3-keto-LG hydrolysate to glucose in the speculated LG metabolic pathway as described above. LgdA, lgdB₁, and a part of lgdC were deleted in the same region in the genome of the B. smithii type strain DSM 4216, which is consistent with the fact that DSM 4216 could not utilize anhydrosugars such as LG and cellobiosan in our previous study.

Thus, the gene products of lgdB₁, lgdB₂, and lgdC were expected to be sugar-related enzymes and involved in LG metabolism. Therefore, we attempted to express these genes and lgdA in Escherichia coli to obtain purified preparations of each protein.

**Recombinant Lgd proteins are functional in converting LG to glucose.** Each lgd gene (lgdA, lgdB₁, lgdB₂, and lgdC) was expressed in E. coli as a GST-fusion protein, followed by removal of the GST moiety with a protease (Fig. S1). Only for the expression of lgdB₂, N-laurolsarcosine was required to obtain the gene product in a soluble fraction. Sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) indicated that the estimated molecular masses of LgdA, LgdB₁, LgdB₂, and LgdC were 44, 30, 36, and 43 kDa, respectively, which were consistent with the molecular weight expected from the corresponding nucleotide sequences.

Our preliminary examination indicated that LG was converted to glucose by the crude extract of S-2701M grown on LG with NAD⁺, Mn²⁺, and 2-mercaptoethanol (2-ME) as cofactors. Based on the reaction conditions with the crude extract, the reaction was conducted with the four recombinant Lgd proteins in various combinations using LG as the substrate with above cofactors, followed by thin layer chromatography (TLC) analysis. LG was completely converted to glucose with all four Lgd proteins (Fig. 2). However, lower level of the conversion

![Figure 1](https://www.nature.com/scientificreports/)
could still be observed even without LgdB1, suggesting that LgdB2 was a partially bifunctional enzyme that could replace the function of LgdB1. NAD+ and Mn2+ were necessary in the reaction mixture for glucose formation, whereas no difference was observed with or without 2-ME (data not shown).

LgdA is confirmed to be LGDH. The purified LgdA had a NAD-dependent dehydrogenase activity against LG (1.32 U/mg), whereas a higher activity (6.0 U/mg) was observed using 3-keto-LG, which is the reaction product obtained after LG oxidation, and NADH as the substrates. These enzyme activities were calculated by initial velocity of the change of absorbance at 340 nm due to NADH. Next, we examined the effect of the other Lgd proteins on the LGDH activity of LgdA. As shown in Table 1, the addition of LgdB1 or LgdB2 increased the specific activity (calculated by initial velocity) of LgdA. Intriguingly, these additional effects were notable when the absorbance of the reaction mixture was measured after a long time (20 min), indicating that the enzyme reaction with LgdA alone has a lower saturation point that could be prolonged by the addition of LgdB1 or LgdB2 with a consequently higher maximum saturation velocity. These results suggest that the conversion of 3-keto-LG to another compound alleviates the reaction product inhibition on LGDH.

Figure 2. Enzyme reactions with Lgd proteins using LG as the substrate. The reaction mixture containing 50 mM HEPES-NaOH (pH 7.5), 0.1% LG, 1 mM β-NAD+, 1 mM MnCl2·4H2O, 10 mM 2-ME, and appropriate amount of the recombinant protein(s) were incubated at 30 °C for 15 h. Ten microliters of the reaction mixture were used for TLC analysis. A, B1, B2, and C in the figure represent LgdA, LgdB1, LgdB2, and LgdC, respectively.

| Additional enzyme(s) | Specific activity (unit/mg-protein) | A420 After 20 min |
|----------------------|-----------------------------------|------------------|
| None                 | 0.44 ± 0.033                      | 0.04 ± 0.004     |
| LgdB1                | 0.98 ± 0.002                      | 0.37 ± 0.002     |
| LgdB2                | 0.52 ± 0.008                      | 0.23 ± 0.004     |
| LgdC                 | 0.35 ± 0.037                      | 0.04 ± 0.003     |
| LgdB1, LgdB2         | 0.86 ± 0.004                      | 0.37 ± 0.005     |
| LgdB1, LgdC          | 0.94 ± 0.003                      | 0.36 ± 0.004     |
| LgdB2, LgdC          | 0.45 ± 0.010                      | 0.05 ± 0.004     |
| LgdB1, LgdB2, LgdC   | 0.68 ± 0.017                      | 0.04 ± 0.004     |

Table 1. Effect of addition of the other Lgd proteins on the LGDH activity. All examinations were carried out in triplicate and the values are represented in the form of mean ± standard deviation.
Glucose is formed by subsequent hydration and reduction. We also attempted to identify the structure of the compound expected to be the reaction product of LgdB2. However, the compound was unstable and degraded rapidly during the extraction and purification of the compound. Hence, we used another approach to identify the compound. As described above, LgdC was expected to be an oxidoreductase using NAD(P) as the cofactor. When NADH was added to the reaction mixture containing 3-keto-LG as the substrate and containing LgdB1, LgdB2, and LgdC as the enzymes, the violet spot that was considered to be the LgdB2 reaction product became weak and the spot corresponding to glucose was observed (Fig. 3). These result suggest that LgdC reduces the C3-keto group of the LgdB2 reaction product to form glucose; accordingly, the reaction product of LgdB2 is presumed to be 3-keto-d-glucose. Indeed, when LgdC was reacted with glucose and NAD+ as the substrate and the cofactor, respectively, the violet spot showed the same migration as that corresponding to the LgdB2 reaction product (Fig. 4). Furthermore, marked reduction of NAD+ in the reaction mixture was also confirmed by spectrophotometry (data not shown).

In 1969, Fukui and Hayano reported the method to determine 3-keto-sucrose and 3-keto-d-glucose. They showed that an absorption maximum at 310 nm was observed when 3-keto-d-glucose was incubated in potassium phosphate buffer (pH 7.0). In this study, the violet spot expected to be the LgdB2 reaction product was extracted from a TLC plate and immediately incubated with potassium phosphate buffer. Consequently, an absorption maximum of the solution was observed at 310 nm as shown in Fig. 5. Furthermore, mass spectrum analysis showed that the LgdB2 reaction product extracted from the TLC spot had a $m/z$ of 379.08 (calculated for $C_{12}H_{20}O_{12}$ [2M + Na+]), suggesting a keto-glucose (Fig. S10). Taken together, these findings suggest that the LgdB2 reaction product is 3-keto-d-glucose.

Accordingly, we speculate that LgdB2 catalyzes the hydration of the 1,2-double bond of 2-hydroxy-3-keto-d-glucal to produce 3-keto-d-glucose, and LgdC is glucose-3-dehydrogenase (G3DH), through which 3-keto-d-glucose is reduced to glucose.

Discussion

In this study, we completely elucidated the metabolism of LG to glucose and showed that it involved four enzymes, namely, LgdA, LgdB1, LgdB2, and LgdC. As shown in Fig. 6, our proposed pathway involves C3-oxidation by LgdA (LGDH), cleavage of the C-O bond of the 1,6-linkage through β-elimination catalyzed by LgdB1, and hydration of the 2-hydroxy-3-keto-d-glucal to produce 3-keto-d-glucose by LgdB2, followed by reduction to glucose by LgdC. This reaction mechanism resembles those proposed for the enzymes belonging to the glycosyl hydrolase family 4 (GH4). Prior to elucidating the LG metabolic pathway, we had expected similar reaction mechanisms between LG degradation and GH4-type hydrolysis. Most GH4 enzyme reactions also require NAD+, Mn2+, and relatively large amounts of a reducing reagent such as dithiothreitol or 2-ME. The reaction mechanism of GH4 enzyme has been well-studied in α-galactosidase (MelA) from Citrobacter freundii. In this...
mechanism, NAD\textsuperscript{+} requires the oxidation of the C3-hydroxy group in a galactose moiety, and Mn\textsuperscript{2+} is necessary for increasing polarization at the position of C3 to form an enolate anion before β-elimination. Reportedly, the reducing agent protects the oxidation of reactive Cys residue\textsuperscript{13}. Recently, Bell et al. reported a novel GH4 type hydrolase (Rg\textsubscript{NanOx}) from the human gut symbiont Ruminococcus gnavus which converted 2,7-anhydro-N-acetylneuraminic acid into N-acetylneuraminic acid\textsuperscript{18}. The reaction mechanism is thought to be close to that of LG conversion into glucose in S-2701M as anhydrosugars are the substrates in both reactions. Thus, in GH4 type hydrolases, a single polypeptide catalyzes four reactions, namely, dehydrogenation, β-elimination, hydration, and reduction, whereas the four enzymes examined in this study together act as a GH4-type enzyme and Mn\textsuperscript{2+} was required after β-elimination for the hydration of 3-keto-d-glucose in the LG degradation pathway in S-2701M. When LG was used as the substrate, the reaction product of LgdB1 was not detected on TLC, whereas the weak spots corresponding to the LgdB2 reaction product was observed when LgdA, LgdB1, and LgdB2 were used as the enzymes (Fig. 2), suggesting that LGDH reaction catalyzed by LgdA is rate-limiting in the in-vitro reconstitution reaction using LG (6 mM) as the substrate. As described above, the LGDH activity of LgdA in reverse reaction (reduction of 3-keto-LG) was higher than that in forward reaction (dehydrogenation of LG),
which was consistent with our previous result in LGDH of \textit{P. phenanthrenivorans} \cite{9}. However, the addition of LgdC caused the marked degradation of LG (Fig. 2), which may be due to NAD$^+$ regeneration accompanying glucose formation by LgdC reaction. Furthermore, in the presence of Mn$^{2+}$ ion, the second violet spot expected to correspond to 3-keto-\textit{d}-glucose was observed from 3-keto-LG using only LgdB2 at the same level as that when LgdB1 and LgdB2 were used (Fig. 3), suggesting that LgdB2 is a bifunctional enzyme that catalyzes the $\beta$-elimination of 3-keto-LG with low affinity and hydration of 2-hydroxy-3-keto-\textit{d}-glucal. This suggestion is also consistent with the results using LG as the substrate and it seemed that glucose formation was weak only when LgdB2 was used, which was probably due to a low concentration of 3-keto-LG (Fig. 2). The additional effect of LgdB1 and LgdB2 on LgdA activity (Table 1) also supported the bifunctional characteristic of LgdB2.

Only a few enzymes capable of cleaving glycosyl linkages via $\beta$-elimination have been reported; these are known as polysaccharide lyases and include pectin lyase \cite{19}, alginate lyase \cite{20}, and glucuronan lyase \cite{21}. Polysaccharide lyases catalyze the cleavage of a glycoside linkage of each polysaccharide from the non-reducing end in a $\beta$-elimination manner. The LgdB1 catalyzes an intramolecular glucoside linkage in LG; thus, LgdB1 is thought to be a novel enzyme, which can be called 3-keto-LG decyclase or 2-hydroxy-3-keto-\textit{d}-glucal synthase (EC 5.5). LgdB2 also has a unique characteristic and catalyzed the hydration of 2-hydroxy-3-keto-\textit{d}-glucal, and hence, it can be called 3-keto-\textit{d}-glucose-1,2-dehydratase. It is known that several sugar dehydratases catalyze 4,6-hydration of hexoses to form 6-deoxysugars, and are members of the short chain dehydrogenase/reductase protein superfamily \cite{22, 23, 24}. The reaction of the sugar 4,6-dehydratases also involves $\beta$-elimination in the 4,6-dehydration to eliminate the C6-hydroxyl group, whereas the dehydration in the LG metabolic pathway occurs in a compound after $\beta$-elimination.

We speculate that LgdC catalyzes the oxidation of C3 position of glucose. Only a few enzymes oxidize a hydroxyl group at C3 position in sugars, and G3DHs have been found in \textit{Cytophaga marinoflava} \cite{25}, \textit{Halomonas sp.} \cite{25}, and \textit{Agaricus bisporus} \cite{25}. It is known that these G3DHs cannot use NAD$^+$ as the cofactor, but can use artificial cofactors such as 2,6-dichloroindophenol. LgdC was also thought to be G3DH and could use NAD$^+$ as the cofactor in this study. LGDH can also oxidize the C3-hydroxyl group of LG, which is the key reaction in the LG metabolic pathway. However, its activity toward d-glucose is very low; 0.05% and 5% activities of LG have been found in \textit{P. phenanthrenivorans} which was consistent with our previous result in LGDH of \textit{P. phenanthrenivorans} \cite{9}. However, the addition of LgdC caused the marked degradation of LG (Fig. 2), which may be due to NAD$^+$ regeneration accompanying glucose formation by LgdC reaction. Furthermore, in the presence of Mn$^{2+}$ ion, the second violet spot expected to correspond to 3-keto-\textit{d}-glucose was observed from 3-keto-LG using only LgdB2 at the same level as that when LgdB1 and LgdB2 were used (Fig. 3), suggesting that LgdB2 is a bifunctional enzyme that catalyzes the $\beta$-elimination of 3-keto-LG with low affinity and hydration of 2-hydroxy-3-keto-\textit{d}-glucal. This suggestion is also consistent with the results using LG as the substrate and it seemed that glucose formation was weak only when LgdB2 was used, which was probably due to a low concentration of 3-keto-LG (Fig. 2). The additional effect of LgdB1 and LgdB2 on LgdA activity (Table 1) also supported the bifunctional characteristic of LgdB2.

In conclusion, this study revealed that four enzymes, including three novel ones coordinated the conversion of LG to glucose in \textit{B. smithii} S-2701M. LG is known to be produced by pyrolysis or supercritical water treatment of lignocellulosic biomass \cite{29, 30}. Typically, saccharification of lignocellulose is a bottleneck for its utilization as feedstock for subsequent fermentation process. Recently, researchers have paid considerable attention to microbial conversion of LG as a biomass-derived compound \cite{31, 32, 33}. Thus, effective utilization of LG would lead to a novel and an effective utilization of cellulosic biomass. However, only fungal LG kinases have been used to construct the recombinant fermentation systems. It is expected that the bacterial LG metabolism involving the \textit{lgl} genes will be applied in the recombinant fermentation system for LG. Alternatively, S-2701M could utilize LG at the same level as glucose, which is free from catabolite repression \cite{34}, suggesting that this bacterium can be an effective host for production of useful compounds from LG. There are reports indicating another application of LG as a useful starting material, and that the ring-opening polymerization of 1,6-anhydrohexopyranose can be used for the synthesis of stereoregular polysaccharides with biological activities, such as anti-HIV reagents and artificial biocompatible materials \cite{35, 36}. The reaction mechanism of LG degradation elucidated in the present study may also contribute to the synthesis of such anhydrosugars.

**Methods**

**Materials.** LG was purchased from MSD K.K., Tokyo, Japan. 3-keto-LG was enzymatically synthesized as described previously \cite{27}.

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**Figure 6.** Proposed LG metabolic pathway in \textit{B. smithii} S-2701M.
Expression of Lgd proteins in Escherichia coli. The region involved in LG metabolism was searched in the draft genome of B. smithii S-2701M (accession no. LC529899) according to the amino acid sequence of LGDH from P. phenanthrenivorans. Lgd genes in this region were amplified from the S-2701M genome by PCR using primer sets with 5'-BamHI and 3'-SalI restriction sites for lgaA, lgdB1, and lgdB2, and 5'-EcoRI and 3'-SalI restriction sites for lgcC, respectively. The primers used in this study are listed in Table S2. Each PCR fragment was introduced into pGEX-6P-1 digested with appropriate restriction enzymes. E. coli DH5α [F’ phi80lacZΔM15 M1lacZYA-argF-U169 de3 recA1 hsdR17(k+ mcrB)] pBR322 thr1 gyrA96 relA1 was used for expression of each plasmid.

The recombinant E. coli was cultivated at 30°C in LB medium containing 50 μg/ml ampicillin, and isopropyl-β-d-thiogalactopyranoside (0.1 mM) was added to the medium when OD600 reached 0.5. The overnight culture was centrifuged, washed with 0.85% KCl, and suspended with PBS buffer (140 mM NaCl, 2.7 mM KCl, 10 mM Na2HPO4, 1.8 mM KH2PO4, 1 mM dl-dithiothreitol (DTT), pH 7.3). For purification of LgdB2, the cell extract was obtained by centrifugation at 20,400 × g for 10 min. Each recombinant protein fused with a glutathione S-transferase was recovered from the cell extract by using a Glutathione Sepharose 4B resin (GE Helthcare Japan, Tokyo). The resin was transferred to a column and washed with the cleavage buffer (50 mM Tris-HCl, 1 mM β-NAD+, 15 μg/ml LgdA, and 20 μg/ml LgdB1). Total 24 ml of the reaction mixture was subjected to TLC analysis as described above. High-resolution mass spectrometry was conducted on a Bruker Compact (ESI-Q-TOF). Nuclear magnetic resonance (NMR) spectra were recorded on a Bruker AVANCE III HD with CryoProbe Prodigy (1H 400 MHz, 13C 100 MHz) spectrometer in D2O–H2O (10:90) at 298 K. Chemical shifts for 1H NMR analysis as described above. High-resolution mass spectrometry was conducted on a Bruker Compact (ESI-Q-TOF). Nuclear magnetic resonance (NMR) spectra were recorded on a Bruker AVANCE III HD with CryoProbe Prodigy (1H 400 MHz, 13C 100 MHz) spectrometer in D2O–H2O (10:90) at 298 K. Chemical shifts for 1H NMR are expressed in parts per million (ppm) relative to MeOH (δ 3.34 ppm). Chemical shifts for 13C NMR are expressed in ppm relative to MeOH (δ 49.5 ppm).

Identification of reaction products. The reaction product of LgdB1 was obtained from LG with LgdA and LgdB1 and NAD regenerative reaction with lactate dehydrogenase (LDH) was used to enhance the LgdA reaction. The reaction was carried out at 30 °C for 17 h in the reaction mixture consisting of 50 mM HEPES-NaOH, 1% LG, 1 mM β-NAD+, 31 mM sodium pyruvate, 1 U/ml LDH (from pig heart, TOYOBO Co., Ltd., Osaka), and LgdB1. The reaction mixture was incubated at 30°C for 30 min. The reaction mixture (40 μl) was subjected to TLC analysis as described above. High-resolution mass spectrometry was conducted on a Bruker Compact (ESI-Q-TOF). Nuclear magnetic resonance (NMR) spectra were recorded on a Bruker AVANCE III HD with CryoProbe Prodigy (1H 400 MHz, 13C 100 MHz) spectrometer in D2O–H2O (10:90) at 298 K. Chemical shifts for 1H NMR were measured spectrophotometrically at 25 °C by following the increasing absorbance at 340 nm in the reaction mixture containing 50 mM Tris-HCl, 1 mM β-NAD+, and appropriate enzyme solution. No difference was observed in each enzyme assay with the other kinds of buffer (phosphate and HEPES).

Identification of reaction products. The reaction product of LgdB2 was determined according to the method of Fukui and Hayano. The reaction mixture containing 30 mM HEPES-NaOH (pH 7.5), 1% LG, 1 mM β-NAD+, 31 mM sodium pyruvate, 1 U/ml LDH (from pig heart, TOYOBO Co., Ltd., Osaka), 15 μg/ml LgdA, and 20 μg/ml LgdB1. Total 24 ml of the reaction mixture was subjected to TLC analysis as described above. High-resolution mass spectrometry was conducted on a Bruker Compact (ESI-Q-TOF). Nuclear magnetic resonance (NMR) spectra were recorded on a Bruker AVANCE III HD with CryoProbe Prodigy (1H 400 MHz, 13C 100 MHz) spectrometer in D2O–H2O (10:90) at 298 K. Chemical shifts for 1H NMR are expressed in parts per million (ppm) relative to MeOH (δ 3.34 ppm). Chemical shifts for 13C NMR are expressed in ppm relative to MeOH (δ 49.5 ppm).

The reaction product of LgdB2 was determined according to the method of Fukui and Hayano. The reaction mixture containing 30 mM HEPES-NaOH (pH 7.0), 0.3% 3-keto-LG, 1 mM MnCl2·4H2O, and 15 μg/ml LgdB2 was incubated at 30 °C for 30 min. The reaction mixture (40 μl) was subjected to TLC analysis as described above.

The reaction product was extracted at 4 °C for 10 min from the corresponding spot with 60 μl of distilled water. Four microliters of the extract were mix with 1 μl of 1 M potassium phosphate buffer (pH 7.0) and the absorption spectrum was measured after 3 min.

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**Competing interests**

The authors declare no competing interests.

**Additional information**

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