Roles of two glutathione S-transferases in the final step of the \(\beta\)-aryl ether cleavage pathway in Sphingobium sp. strain SYK-6

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Sphingobium sp. strain SYK-6 is an alphaproteobacterial degrader of lignin-derived aromatic compounds, which can degrade all the stereoisomers of \(\beta\)-aryl ether-type compounds. SYK-6 cells convert four stereoisomers of guaiacylglycerol-\(\beta\)-guaiacyl ether (GGE) into two enantiomers of \(\alpha\)-(2-methoxyphenoxy)-\(\beta\)-hydroxypropiovanillone (MPHPV) through GGE \(\alpha\)-carbon atom oxidation by stereoselective \(\text{Ca} \)-dehydrogenases encoded by \(\text{ligD, ligL, and ligN}\). The ether linkages of the resulting MPHPV enantiomers are cleaved by stereoselective glutathione (GSH) S-transferases (GSTs) encoded by \(\text{ligF, ligE, and ligP}\), generating (\(\beta\R/\beta\S\))-\(\alpha\)-glutathionyl-\(\beta\)-hydroxypropiovanillone (GS-HPV) and guaiacol. To date, it has been shown that the gene products of \(\text{ligG}\) and SLG_04120 (\(\text{ligQ}\), both encoding GST, catalyze GSH removal from (\(\beta\R/\beta\S\))-GS-HPV, forming achiral \(\beta\)-hydroxypropiovanillone. In this study, we verified the enzyme properties of LigG and LigQ and elucidated their roles in \(\beta\)-aryl ether catabolism. Purified LigG showed an approximately 300-fold higher specific activity for (\(\beta\R\))-GS-HPV than that for (\(\beta\S\))-GS-HPV, whereas purified LigQ showed an approximately six-fold higher specific activity for (\(\beta\S\))-GS-HPV than that for (\(\beta\R\))-GS-HPV. Analyses of mutants of \(\text{ligG, ligQ, and both genes revealed that SYK-6 converted}\) (\(\beta\R\))-GS-HPV using both LigG and LigQ, whereas only LigQ was involved in converting (\(\beta\S\))-GS-HPV. Furthermore, the disruption of both \(\text{ligG}\) and \(\text{ligQ}\) was observed to lead to the loss of the capability of SYK-6 to convert MPHPV. This suggests that GSH removal from GS-HPV catalyzed by LigG and LigQ, is essential for cellular GSH recycling during \(\beta\)-aryl ether catabolism.

Lignin is one of the major components of plant cell walls, accounting for 15–40%\(^1\). The industrial production of useful substances from lignin is desired, since it is the most abundant aromatic resource and the second most abundant bioresource on Earth after cellulose. In recent years, microbial conversion processes known as “biological funneling” have attracted attention for their ability to upgrade heterogeneous mixtures of low-molecular-weight aromatic compounds obtained through chemical lignin depolymerization into platform chemicals\(^2\-5\). Based on these facts, elucidating the bacterial lignin catabolic system has become more critical.

Lignin is formed through the dehydrogenative polymerization of hydroxycinnamic alcohols, resulting in various C–C and C–O–C bonds between the phenylpropane units\(^6,7\). Although lignin has asymmetric carbons, it is considered optically inactive because of its equal amounts of optical isomers\(^8\). Among the intermonomer linkages in lignin, \(\beta\)-aryl ether is the most abundant, comprising 45%–50% of all linkages in softwood lignin and 60–62% in hardwood lignin\(^9\). Accordingly, cleavage of the \(\beta\)-aryl ether linkage is considered a crucial step in lignin biodegradation. Furthermore, \(\beta\)-aryl ether-type biaryls have two distinct isomeric forms, erythro and threo, and each form has enantiomeric forms\(^10,11\).

Sphingobium sp. strain SYK-6 is an alphaproteobacterium that exhibits the best-characterized catabolic system for lignin-derived aromatic compounds\(^3,12\). SYK-6 can utilize lignin-derived biaryls, including \(\beta\)-aryl ether, phenylcoumaran, biphenyl, and diarylpropene, and monoaryls, including ferulate, vanillin,
and syringaldehyde, as its sole sources of carbon and energy. In SYK-6 cells, four stereoisomers of a model β-aryl ether compound, guaiacylglycerol-β-guaiacyl ether (GGE), are converted into two enantiomers of α-(2-methoxyphenoxy)-β-hydroxypropiovanillone (MPHPV) through the oxidation of the α-carbon, which is catalyzed by Cα-dehydrogenases, LigD, LigL, and LigN (Fig. 1)\textsuperscript{13}. LigD oxidizes (α\textsubscript{R},β\textsubscript{S})-GGE and (α\textsubscript{R},β\textsubscript{R})-GGE into (β\textsubscript{S})-MPHPV and (β\textsubscript{R})-MPHPV, respectively, whereas LigL/LigN converts (α\textsubscript{S},β\textsubscript{R})-GGE and (α\textsubscript{S},β\textsubscript{S})-GGE into (β\textsubscript{R})-MPHPV and (β\textsubscript{S})-MPHPV, respectively. The ether linkage in the resulting MPHPV is cleaved by enantioselective glutathione (GSH) S-transferases (GSTs), LigF, LigE, and LigP (β-etherases), to produce α-glutathionyl-β-hydroxypropiovanillone (GS-HPV) and guaiacol through the nucleophilic attack of GSH on the MPHPV β-carbon atom (Fig. 1)\textsuperscript{14–16}. LigF and LigE/LigP attack (β\textsubscript{S})-MPHPV and (β\textsubscript{R})-MPHPV, subsequently producing (β\textsubscript{R})-GS-HPV and (β\textsubscript{S})-GS-HPV, respectively\textsuperscript{14–18}. Recently, X-ray crystal structures of LigE and LigF were determined, suggesting that LigE was most similar to the GSTFuA fungal class, and LigF was classified in a new structural class closely related to GSTFuAs or fungal Ure2p-like GST\textsuperscript{19}. LigF and LigE cleave the β-aryl ether linkage through an \textit{S}_\text{N}^2\textsuperscript{18,19} nucleophilic attack of GSH on the β-carbon of the substrates\textsuperscript{8,19}. Another GST, LigG, catalyzes the cleavage of the thioether linkage in (β\textsubscript{R})-GS-HPV by transferring the (β\textsubscript{R})-GS-HPV GSH moiety to another GSH molecule, producing achiral β-hydroxypropiovanillone (HPV) and oxidized glutathione (GSSG)\textsuperscript{14,16}. Meux et al. and Pereira et al. independently solved the X-ray crystal structure of LigG\textsuperscript{20,21}. LigG belongs to the \textit{omega}-class GST, and Cys15 is a catalytic residue\textsuperscript{20}. LigG had little to no activity for (β\textsubscript{S})-GS-HPV, suggesting the involvement of an alternative GST in (β\textsubscript{S})-GS-HPV conversion\textsuperscript{14}. A recent study reported that a \textit{Nu}-class GST, NmGST3 of \textit{Novosphingobium} sp. strain MBES04, exhibited GSH-removing activity toward both GS-HPV isomers\textsuperscript{22,23}. Kontur et al. reported that NaGST\textsubscript{val}, belonging to the \textit{Nu}-class GST, showed activity toward both GS-HPV isomers, and it was the only enzyme essential for converting both GS-HPV isomers in \textit{Novosphingobium aromaticivorans} DSM 12444\textsuperscript{24}. Furthermore, they showed that the gene product of SLG_04120, a putative \textit{Nu}-class GST in SYK-6, had GSH-removing activity for both GS-HPV isomers.

In this study, we identified the GS-HPV-converting GST genes in SYK-6 and uncovered the roles of these genes in β-aryl ether catabolism, since the genes involved in GS-HPV catabolism have not been clarified in SYK-6 to date.

\begin{figure}
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\includegraphics[width=\textwidth]{Figure1.png}
\caption{Catabolic pathway of guaiacylglycerol-β-guaiacyl ether in \textit{Sphingobium} sp. strain SYK-6. Stereoisomers of β-aryl ether model compound, GGE, are stereospecifically converted to an achiral C6–C3 monomer, HPV\textsuperscript{13–16}. The resulting HPV is oxidized to vanillol acetic acid via vanillol acetaldehyde and further catabolized through vanillate\textsuperscript{13,37}. Enzymes: LigD, LigL, and LigN, Ca-dehydrogenases; LigF, LigE, and LigP, β-etherases; LigG and LigQ, GSH-removing enzymes; HpvZ, HPV oxidase; ALDHs, aldehyde dehydrogenases; SLG_20400, vanillyl acetaldehyde dehydrogenase. \textit{Abbreviations}. GGE, guaiacylglycerol-β-guaiacyl ether; MPHPV, α-(2-methoxyphenoxy)-β-hydroxypropiovanillone; GS-HPV, α-glutathionyl-β-hydroxypropiovanillone; HPV, β-hydroxypropiovanillone; GS\textsuperscript{−}, reduced glutathione; GSSG, oxidized glutathione.}
\end{figure}
Results

Preparation of GS-HPV isomers. Racemic MPHVP (400 μM) was reacted with purified LigF and LigE (Fig. S1) with 5 mM GSH in 50 mM Tris–HCl buffer (pH 7.5; buffer A) to obtain enantiopure GS-HPV isomers24. High-performance liquid chromatography (HPLC) analysis of the reaction mixtures showed that MPHVP was converted into GS-HPV by purified LigF and LigE at 60 min (Fig. S2A–C). The conversion of MPHVP by LigF and LigE stopped at approximately 50%, and approximately equimolar GS-HPV to the converted MPHVP was generated (Fig. S2D and E). Chiral HPLC analysis was conducted to confirm whether LigF and LigE enantioselectively convert MPHVP. The MPHVP preparation that was used as the substrate was an equimolar mixture of (βR)-MPHPV and (βS)-MPHPV (53:47) (Fig. S3A). LigF and LigE completely converted (βS)-MPHPV and (βR)-MPHPV, respectively, indicating (βR)-GS-HPV and (βS)-GS-HPV generation in each reaction (Fig. S3B and C). We attempted to remove the remaining MPHVP and guaiacol from the reaction mixtures by ethyl acetate extraction; however, we found that the prepared GS-HPV isomers racemized after incubation in buffer A at 30 °C for 24 h (data not shown). Hence, we prepared each GS-HPV isomer by incubating 400 μM MPHVP with purified LigF or LigE and 5 mM GSH in buffer A before usage.

Conversion of GS-HPV isomers by Sphingobium sp. strain SYK-6. In order to verify the ability of SYK-6 to convert GS-HPV isomers, the extracts of SYK-6 cells grown in Wx minimal medium containing 10 mM sucrose, 10 mM glutamate, 0.13 mM methionine, and 10 mM proline (Wx-SEMP) were incubated with 100 μM (βR)-GS-HPV or (βS)-GS-HPV and 2.4 mM GSH in buffer A. HPLC analysis indicated that SYK-6 converted both GS-HPV isomers into HPV after incubation for 5 min (Fig. S4). The activities of the extracts of SYK-6 cells grown in a Wx-SEMP with or without 5 mM GGE supplementation were measured to evaluate the inducibility of GS-HPV conversion. The specific activity of the extracts from the cells grown in Wx-SEMP for (βS)-GS-HPV (142 ± 3 nmol·min⁻¹·mg⁻¹) was approximately six-fold higher than that for (βR)-GS-HPV (24.5 ± 0.2 nmol·min⁻¹·mg⁻¹). The specific activities for (βS)-GS-HPV (150 ± 19 nmol·min⁻¹·mg⁻¹) and (βR)-GS-HPV (31.4 ± 1.1 nmol·min⁻¹·mg⁻¹) of the extracts from the cells grown in Wx-SEMP supplemented with 5 mM GGE were not significantly higher than those of the extracts from the cells grown in Wx-SEMP. These results suggest that GSH-removing enzyme genes are constitutively expressed.

Identification of the GST genes involved in GS-HPV isomer conversion. Figure 2 shows a phylogenetic tree of 18 putative GSTs from SYK-6, including LigF, LigE, LigP, LigG, and SLG_04120, with GSTs conserved in SLG_29340 (Cys10 and Pro11) (Fig. S5). On the basis of these findings, we selected ligG, SLG_00360, SLG_04120, and SLG_29340 as candidates for GS-HPV-converting GST genes. To examine whether the gene products of ligG, SLG_00360, SLG_04120, and SLG_29340 exhibit activities toward GS-HPV, these genes fused with a His-tag at the N-terminal were expressed in E. coli. Sodium dodecyl sulfate–polyacrylamide gel electrophoresis (SDS-PAGE) showed the protein production of 33, 28, 36, and 33 kDa, appearing as products of ligG, SLG_00360, SLG_04120, and SLG_29340 (Fig. S6). The cell extracts of these E. coli transformants were incubated with 100 μM (βR)-GS-HPV or (βS)-GS-HPV and 2.4 mM GSH in buffer A. HPLC analysis showed that ligG gene product completely converted (βR)-GS-HPV into HPV after incubation for 30 min, whereas it only slightly converted (βS)-GS-HPV (Fig. 3A,B). This result confirmed that LigG is highly specific for (βR)-GS-HPV, which is similar to that of previous reports4,14,21,26. By contrast, the SLG_04120 gene product completely converted both GS-HPV isomers into HPV, verifying its specificity, as previously reported (Fig. 3C,D)24. However, SLG_00360 and SLG_29340 gene products showed no activity for both GS-HPV isomers (Fig. 3E–H). Given these results, we designated SLG_04120 as ligQ and investigated the role of ligG and ligQ in converting GS-HPV isomers in SYK-6, as well as the enzymatic properties of these gene products.

Roles of ligG and ligQ in GS-HPV isomer catalysis. We then created ligG mutant (ΔligG), ligQ mutant (ΔligQ), and ligG ligQ double mutant (ΔligG ligQ) (Fig. S7) to determine whether ligG and ligQ are indeed involved in converting GS-HPV in SYK-6. The cell extract of each mutant was reacted with 100 μM (βR)-GS-HPV or (βS)-GS-HPV and 2.4 mM GSH in buffer A, and their ability to convert GS-HPV isomers was compared with SYK-6. ΔligG and ΔligQ both showed a decreased conversion of (βR)-GS-HPV, but only ΔligQ exhibited the loss of (βS)-GS-HPV conversion (Fig. 4A,B). Furthermore, ΔligQ ligQ had little ability to convert both GS-HPV isomers (Fig. 4A,B). We also examined the activities of ΔligG ligQ cells harboring each of pqF carrying ligG (pQPqFligG) and ligQ (pQPqFligQ) to confirm whether this activity defect was due to the disruption of ligG and ligQ. A cell extract of ΔligG ligQ harboring pQPqFligG showed higher specific activity for (βR)-GS-HPV than that of the wild type (Fig. 4C). Also, a cell extract of ΔligQ ligQ harboring pQPqFligQ presented almost the same specific activity for (βR)-GS-HPV as the wild type (Fig. 4C) and higher specific activity for (βS)-GS-HPV than the wild type (Fig. 4D). These results indicated that both ligG and ligQ participate in the conversion of (βR)-GS-HPV in SYK-6 cells, whereas only ligQ is involved in (βS)-GS-HPV conversion.

Enzyme properties of LigG and LigQ. LigG and LigQ were purified to near-homogeneity through nickel affinity chromatography from the cell extracts of E. coli expressing His-tagged ligG and ligQ, respectively (Fig. S8). Purified LigG and LigQ were incubated with 100 μM (βR)-GS-HPV or (βS)-GS-HPV and 2.4 mM GSH in buffer A. LigG showed approximately 300-fold higher specific activity for (βR)-GS-HPV (33 ± 1 μmol·min⁻¹·mg⁻¹) than

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for (βS)-GS-HPV (0.11 ± 0.01 µmol·min⁻¹·mg⁻¹). By contrast, LigQ showed approximately six-fold higher specific activity for (βS)-GS-HPV (110 ± 14 µmol·min⁻¹·mg⁻¹) than for (βR)-GS-HPV (17 ± 2 µmol·min⁻¹·mg⁻¹). The specific activity of LigG for (βR)-GS-HPV was approximately two-fold higher than that of LigQ. These results indicated that LigG is highly specific for (βR)-GS-HPV, and LigQ prefers (βS)-GS-HPV more than (βR)-GS-HPV. The $K_m$ values of LigG and LigQ for GSH were determined to be 1.19 ± 0.22 and 0.34 ± 0.10 mM, respectively (Fig. S9).

**Disruption of both ligG and ligQ affects the MPHPV catabolism.** The resting cells of each mutant were reacted with 200 µM racemic MPHPV in buffer A to investigate whether the disruption of ligG and ligQ in SYK-6 affects MPHPV conversion. Here ΔligG and ΔligQ cells converted MPHPV similar to the wild type, whereas ΔligG ΔligQ cells lost the ability to convert MPHPV (Fig. 5A). By contrast, a cell extract of ΔligG ΔligQ was able to convert MPHPV in the presence of 5 mM GSH in buffer A at comparable levels with that of the wild type (Fig. S10). The introduction of pQF ligG or pQF ligQ into ΔligG ΔligQ cells restored their MPHPV-converting ability, indicating that the disruption of both ligG and ligQ led to the loss of the ability to convert MPHPV in vivo (Fig. 5B).

**Discussion**

We confirmed that LigG is highly specific for (βR)-GS-HPV, and LigQ converts both GS-HPV isomers. Our study further elucidated that both ligG and ligQ participate in converting (βR)-GS-HPV in SYK-6 cells, whereas only ligQ is involved in converting (βS)-GS-HPV (Figs. 1, 4A,B). Considering that the rate of (βR)-GS-HPV conversion.

**Figure 2.** Phylogenetic tree of GSTs from SYK-6 and other bacterial GSTs involved in β-aryl ether conversion. The phylogenetic tree was constructed by neighbor-joining with 1,000 bootstrap replicates. Bootstrap values are indicated at the nodes, and the scale corresponds to 0.2 amino acid substitutions per position. GSTs from SYK-6 are shown in red. Nu-class GSTs and Omega-class-like GSTs are shown with blue and green backgrounds, respectively. GSTs: LigF (SLG_08650), LigE (SLG_08660), and LigP (SLG_32600), β-etherases of *Sphingobium* sp. strain SYK-6; LigQ (SLG_04120) and LigG (SLG_08670), GSH-removing enzymes of SYK-6; SLG_00360 (BAK64711), SLG_06390 (BAK65314), SLG_07230 (BAK65398), SLG_11530 (BAK65828), SLG_24810 (BAK67156), SLG_28090 (BAK67484), SLG_28330 (BAK67508), SLG_29340 (BAK67699), SLG_31200 (BAK67795), SLG_33010 (BAK67976), SLG_33060 (BAK67981), SLG_34480 (BAK68123), and SLG_36080 (BAK68283), putative GSTs from SYK-6; NmGST4 (GM05530) and NmGST5 (GM05531), β-etherases of *Novosphingobium* sp. strain MBES04; NmGST3 (GM05529) and NmGST6 (GM05532), GSH-removing enzymes of MBES04; NsLigF (CCA92087) and NsLigE (CCA92088), β-etherases of *Novosphingobium* sp. strain PIPY; NsLigG (CCA92089), GSH-removing enzyme of PIPY; NaLigE (ABD26841), NaLigF1 (ABD26530), and NaLigF2 (ABD27301), β-etherases of *Novosphingobium aromaticivorans* DSM 12444; NaGSTNu (ABD27031), GSH-removing enzyme of DSM 12444; TdLigG (AAZ97003), GSH-removing enzyme of *Thiobacillus denitrificans* ATCC 25259.
Figure 3. GS-HPV enantiomer conversion by cell extracts of *E. coli* carrying GST genes of SYK-6. (βR)-GS-HPV (A,C,E,G; 100 µM) and (βS)-GS-HPV (B,D,F,H; 100 µM) were incubated with cell extracts of *E. coli* (100 µg protein/ml) harboring each of pET08670 (A,B), pET04120 (C,D), pET00360 (E,F), and pET29340 (G,H) in the presence of 2.4 mM GSH, respectively. Reaction mixtures were collected after incubation for 30 min and analyzed through HPLC.

Figure 4. GS-HPV enantiomer conversion by Δ*ligG*, Δ*ligQ*, and Δ*ligG ligQ*. (A,B) Conversion of 100 µM (βR)-GS-HPV (A) and (βS)-GS-HPV (B) by cell extracts (100 µg protein/ml) of SYK-6 (circles), Δ*ligG* (triangles), Δ*ligQ* (squares), and Δ*ligG ligQ* (diamonds) in the presence of 2.4 mM GSH. (C,D) Complementation of Δ*ligG ligQ* using pQF*ligG* and pQF*ligQ*, respectively. Gene expression was induced for 24 h at 30 °C by adding 1 mM cumate. The cell extracts of SYK-6(pQF), Δ*ligG* ligQ(pQF), Δ*ligG ligQ*(pQF*ligG*), and Δ*ligG ligQ*(pQF*ligQ*) were incubated with 100 µM (βR)-GS-HPV (C) and (βS)-GS-HPV (D) in the presence of 2.4 mM GSH. All experiments were conducted in triplicate, and each value represents the mean ± standard deviation. ND, not detected.
ligG that of ligQ influence on (βR) had more ligG conversion than LigQ and NaGSTNu, respectively. These genes may also be involved in GS-HPV isomer conversion.

Moreover, we found PP1Y_AT11650 (accession no. CCA92086) in the genome of this strain, which showed 80% S-class GSTs have higher specificity for (βS) NsLigG (PP1Y_AT11674), exhibiting 63% identity with LigG, was highly specific for (βS)-GS-HPV in this study. Kontur et al. reported that LigQ showed approximately 11-fold higher specific activity for (βS)-GS-HPV than for (βR)-GS-HPV21,22. Conversely, LigQ showed approximately six-fold higher specific activity for (βS)-GS-HPV than for (βR)-GS-HPV (0.11 ± 0.01 μmol·min⁻¹·mg⁻¹) was only 0.33% of that for (βR)-GS-HPV (33 ± 1 μmol·min⁻¹·mg⁻¹). This result is similar to previous reports demonstrating that LigG had little to no activity for (βS)-GS-HPV derivatives21,26. By contrast, Rosini et al. reported that 500 μM racemic GGE was fully converted into HPV and guaiacol by purified LigD, LigL, LigF, LigE, and LigG in 2 h27, suggesting that LigG catalyzed the thioether cleavage of both (βS)- and (βR)-GS-HPV. However, the conversion of (βS)-GS-HPV by LigG was likely to have proceeded because a large amount of purified LigG (200 μg/ml) was used for this conversion.

Conversely, LigQ showed approximately six-fold higher specific activity for (βS)-GS-HPV than for (βR)-GS-HPV in this study. Kontur et al. reported that LigQ showed approximately 11-fold higher kcat/Km value for (βS)-GS-HPV than for (βR)-GS-HPV (Table S2). Also, NaGSTnu of DSM 12444 showed approximately four-fold higher kcat/Km value for (βS)-GS-HPV than for (βR)-GS-HPV23, and NmGST3 of MBES04 had approximately two-fold higher specific activity for (βS)-GS-HPV than for (βR)-GS-HPV (Table S2). Therefore, all these Nm GSTs have higher specificity for (βS)-GS-HPV than for (βR)-GS-HPV.

The K values of LigG and LigQ for GSH were determined to be 1.19 ± 0.22 and 0.34 ± 0.10 mM, respectively (Table S1, Table S2, and Fig. S9). These values were considerably higher than K of LigG for (βR)-GS-HPV (0.016 ± 0.001 mM)21 and K of LigQ for (βR)-GS-HPV (0.055 ± 0.007 mM) and (βS)-GS-HPV (0.011 ± 0.002 mM). The K of LigG for GSH is supported by the observation of Picart et al. that the specific activity of LigG was halved when reacted in the presence of 0.4 mM GSH compared to 1 mM GSH24. Based on the quantum mechanics/molecular mechanics simulations, Prates et al. suggested that HPV leaves the active site after GS- of GS-HPV is transferred to Cys15 of LigG, giving enough space to accommodate a second GSH for a subsequent thiol-disulfide reaction, which forms GSH disulfide and restores the catalytic ability of LigG18. The lack of specific interactions to bind the GSH in the active site of LigG was suggested to be associated with the low affinity of LigG for GSH.
with high $K_m$ value for GSH estimated by the results of Picart et al\textsuperscript{18}. In addition, the $K_m$ values for GSH of *E. coli* Nu-class GSTs (YghU and YfgG) have been reported to be on the order of mM when using 2-hydroxyethyl disulfide as a substrate ($K_m$ of YghU, ca. 1.2 mM; $K_m$ of YfgG, ca. 1.6 mM)\textsuperscript{23,29}.

The mechanism for the cleavage of the thioether bond in GS-HPV by NaGST\textsubscript{Nu}, was proposed based on X-ray crystal structure, modeling of substrate binding, and mutant analysis. Thr51 and Asn53 of NaGST\textsubscript{Nu} provide hydrogen bonds that stabilize a reactive GSH thiolate anion, which attacks the GS− moiety of GS-HPV to form GS−-SG disulfide. The rupture of the thioether bond is facilitated by the formation of a transient enolate intermediate. This intermediate is stabilized by the interactions between GS-HPV and Tyr166 and Tyr224 hydroxy groups. The capture of a solvent-derived proton by the carbanion collapses the enolate to form HPV\textsuperscript{24}. All these residues are conserved in LigQ, GAM07399, and CCA92884, suggesting that these GSTs catalyze GSH removal from GS-HPV using the same reaction mechanism as that of NaGST\textsubscript{Nu}.

Finally, we observed that ΔligG ligQ lost the ability to convert MPHPV in vivo (Fig. 5A). In ΔligG ligQ cells, intracellular GSH was seemingly depleted due to GS-HPV accumulation during MPHPV catabolism. Consequently, the reactions that are catalyzed by β-etherases (LigF, LigE, and LigP) could not proceed (Fig. 1). Conversion of GS-HPV by LigG and LigQ produces HPV and GSSG; the latter of which must be reduced to GSH by NAD(P)H-dependent GSSG reductase (Fig. 1)\textsuperscript{30}. Hence, it can be concluded that GS− removal from GS-HPV by LigG and LigQ is essential for maintaining intracellular GSH concentration during β-aryl ether catabolism.

**Methods**

**Bacterial strains, plasmids, and culture conditions.** Table S3 lists the strains and plasmids that were used in this study. *Sphingobium* sp. strain SYK-6 and its mutants were grown in lysogeny broth (LB), Wx-SEMP, and Wx-SEMP containing 5 mM GGE at 30 °C. When necessary, 50 mg kanamycin/liter, 100 mg streptomycin/liter, or 12.5 mg tetracycline/liter were added to the cultures. *E. coli* strains were grown in LB at 37 °C. The media for *E. coli* transformants were supplemented with 100 mg ampicillin/liter, 25 mg kanamycin/liter, or 12.5 mg tetracycline/liter.

**Preparation of substrates.** MPHPV and HPV were prepared as described previously\textsuperscript{17,31}. In preparing (βR)-GS-HPV and (βS)-GS-HPV, purified LigF (40 µg protein/ml) and LigE (150 µg protein/ml) were incubated in 1 ml reaction mixtures containing 400 µM racemic MPHPV, 5 mM GSH, and buffer A, respectively, for 60 min at 30 °C\textsuperscript{14}. The resulting mixtures were used as preparations of 200 µM GS-HPV isomers, including 4.8 mM GSH. GGE and guaiacol were purchased from Tokyo Chemical Industry, Co., Ltd.

**HPLC analysis.** HPLC analysis was conducted using the ACQUITY UPLC system (Waters). The MPHPV and GS-HPV reaction products were analyzed using a TSKgel ODS-140TP column (2.1 x 100 mm; Tosoh). Additionally, a CHIRALPAK IE column (4.6 x 250 mm; DAICEL) was used to analyze MPHPV reaction products. All analyses were conducted at a flow rate of 0.5 ml/min. The mobile phase was a mixture of solution A (acetonitrile containing 0.1% formic acid) and B (water containing 0.1% formic acid) under the following conditions: detection of MPHPV reaction products using TSKgel ODS-140TP: 0–25.0 min, 30% A. Detection of GS-HPV reaction products using CHIRALPAK IE: 0–25.0 min, 30% A. Detection of MPHPV reaction products using CHIRALPAK IE: 0–25.0 min, 30% A. Detection of MPHPV reaction products using TSKgel ODS-140THP: 0–5.0 min, 10% A. MPHPV, GS-HPV, guaiacol, and HPV were detected with 280, 286, 276, and 276 nm, respectively.

**Enzyme assays for cell extracts of SYK-6 and its mutants.** GS-HPV-transforming activities of the cell extracts of SYK-6 and its mutants were determined by measuring HPV production using HPLC. The MPHPV-transforming activities of the cell extracts of SYK-6 and its mutants were determined by measuring the decrease in MPHPV using HPLC. The reaction products were also detected using HPLC analysis. SYK-6 and its mutant cells (ΔligG, ΔligQ, and ΔligG ligQ) grown in LB were washed with Wx medium, resuspended in Wx-SEMP medium to an optical density at 600 nm (OD\textsubscript{600}) of 0.2, and incubated for 16 h. The resultant cells were washed twice with buffer A. The cells that were resuspended in the same buffer were then broken by an ultrasonic disintegrator (QSonica Q125; WakenBtech Co., Ltd.), and the supernatants were obtained as cell extracts after centrifugation (19,000 x g for 15 min at 4 °C). Protein concentration was determined using the Bradford method, using bovine serum albumin as the standard (Bio-Rad Laboratories).

Cell extracts (5–1000 µg protein/ml) were incubated with 100 µM (βR)-GS-HPV, (βS)-GS-HPV, or 200 µM racemic MPHPV in the presence of 2.4 mM (for GS-HPV) or 5.0 mM GSH (for MPHPV) in buffer A at 30 °C. The reactions were stopped by adding acetonitrile (final concentration of 50%) after 3 min (measurement of the specific activities for GS-HPV), 5 min (identification of GS-HPV metabolites), 20 min (measurement of the specific activities for GS-HPV in mutant-complemented strains), or 60 min (for MPHPV). Precipitated proteins were removed by centrifugation at 19,000 x g for 15 min, and the resultant supernatants were diluted with water to a final acetonitrile concentration of 12.5%, filtered, and analyzed using HPLC. The specific activity of GS-HPV conversion was expressed in moles of HPV produced per min per milligram of protein. The cells of SYK-6 grown in LB were washed with Wx medium, resuspended in Wx-SEMP medium to an OD\textsubscript{600} of 0.2, and incubated at 30 °C to examine the effects of GGE on enzyme induction. When the cultures reached an OD\textsubscript{600} of 0.4–0.5, we added 5 mM GGE to them. After 12 h of further incubation, cell extracts were prepared and used for the enzyme assay. For complementing ΔligG ligQ, pQFligG and pQFligQ were constructed by In-Fusion cloning (Takara Bio) of ligG and ligQ fragments amplified by PCR using SYK-6 total DNA and their corresponding primer pairs (Table S4) into pQF. After confirming the nucleotide sequences of the inserts, each plasmid was introduced into...
ΔligG ligQ cells through electroporation. The transformed cells were grown in Wx-SEMP containing 100 µM cumate and tetracycline for 16 h, and the cell extracts were prepared and used for the enzyme assay.

**Sequence analysis.** Nucleotide sequences were determined using Eurofins Genomics. Sequence analysis was conducted using the MacVector program (MacVector, Inc.). Sequence similarity searches, pairwise alignments, and multiple alignments were conducted using the BLASTP program, the EMBOSS Needle program through the EMBL-EBI server, and the Clustal omega program, respectively. For phylogenetic analysis, multiple alignments were conducted using the Clustal omega program, and then, phylogenetic trees were generated using the neighbor-joining algorithm of the MEGA 7 software ver. 7.0.26, applying 1,000 bootstrap replicates.

**Expression of SYK-6 GST genes in E. coli and enzyme purification.** DNA fragments carrying SLG_00360, ligQ (SLG_08660), ligF (SLG_08650), ligE (SLG_08660), ligG (SLG_08670), and SLG_29340 were amplified through PCR using SYK-6 total DNA and the primer pairs listed in Table S4, and each amplified fragment was cloned into pET-16b by In-Fusion Cloning. The nucleotide sequences of the inserts were confirmed by sequencing. Each expression plasmid was introduced into E. coli BL21(DE3), and the transformed cells were grown in LB. Gene expression was induced for 4 h at 30 °C by adding 1 mM isopropyl-β-D-thiogalactopyranoside when the OD600 of the cultures reached 0.5. The cells were then harvested by centrifugation at 5000×g for 5 min at 4 °C and washed with buffer A. The cells were then resuspended in the same buffer and broken using an ultrasonic disintegrator. The supernatants were obtained as cell extracts after centrifugation at 19,000×g for 15 min at 4 °C. For purification of LigF, LigE, LigG, and LigQ, the cell extracts of E. coli BL21(DE3) harboring pET08650, pET08660, pET08670, and pET04120, respectively (Table S3), were transferred onto a His SpinTrap column (GE Healthcare). The purified fractions were subjected to desalting and concentration using an Amicon Ultra centrifugal filter unit (30 kDa cutoff; Merck Millipore), and the enzyme preparations were stored at −80 °C. Gene expressions and the purity of the preparations were examined using SDS-12% PAGE. The protein bands in gels were stained with Coomassie Brilliant Blue.

**Enzyme assays for cell extracts of E. coli transformants and purified enzymes.** GS-HPV-transforming activities were measured by measuring HPV production using HPLC. The reaction products were also detected through HPLC analysis. The cell extracts (100 µg protein/ml) and purified enzymes (LigG, 1.25 and 250 µg protein/ml for the conversion of (βR)- and (βS)-GS-HPV, respectively; LigQ, 0.2 and 1.0 µg protein/ml for the conversion of (βS)- and (βR)-GS-HPV, respectively) were incubated in the 100 µl reaction mixtures containing 100 µM (βR)-GS-HPV or (βS)-GS-HPV, 2.4 mM GSH, and buffer A at 30 °C. The reactions were stopped by adding acetonitrile or methanol to a final concentration of 50% after 30 min of incubation (for cell extracts) and 15 s (for purified enzymes). Protein precipitates were removed through centrifugation at 19,000×g for 15 min. The resulting supernatants were diluted with water to a final concentration of acetonitrile or methanol of 12.5%, filtered, and analyzed using HPLC. Specific activities for GS-HPV conversion of purified enzymes were calculated after 15 s incubation and expressed in moles of HPV produced per min per milligram of protein. The $K_m$ values of LigG and LigQ for GSH were determined using 100 µM (βR)-GS-HPV under the following protein and GSH concentrations: LigG, 1.25 µg/ml protein and 0.25–4.9 mM GSH; LigQ, 5.0 µg/ml protein and 0.1–1.9 mM GSH. The $K_m$ values were calculated through non-linear regression analysis using the GraphPad Prism 7 software (GraphPad Software Inc.; https://www.graphpad.com/scientific-software/prism/) fitted to the Michaelis–Menten equation and expressed as means ± standard deviations of three independent experiments.

**Construction of mutants.** To construct the ligG and ligQ mutants, the upstream and downstream regions of the genes were amplified through PCR from SYK-6 total DNA using the primer pairs listed in Table S4. The resulting fragments were cloned into pAK405 by In-Fusion Cloning. Each of the resulting plasmids was introduced into the SYK-6 cells by triparental mating, and the resulting mutants were selected as described previously. Gene deletion was confirmed through colony PCR using the primer pairs listed in Table S3.

**Resting cell assay.** The cells of SYK-6, ΔligG, ΔligQ, and ΔligG ligQ were grown in Wx-SEMP for 16 h. The cells were then collected by centrifugation, washed twice with buffer A, and resuspended in the same buffer. After adding 200 µM of MPHPV, the resting cells (OD600 of 1.0) were incubated at 30 °C with shaking for 24 h. Portions of the cultures were periodically collected, and the amounts of the MPHPV were measured using HPLC. For complementation analysis of ΔligG ligQ, the resting cells (OD600 of 5.0) of SYK-6 harboring pQF, ΔligG ligQ harboring pQFligG, or ΔligG ligQ harboring pQFligQ, grown in Wx-SEMP containing cumate (100 µM) and tetracycline, were incubated with 200 µM MPHPV at 30 °C with shaking for 24 h.

**Data availability**
All data supporting this study are available within the article and its Supplementary Information or are available from the corresponding author upon request.

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Author contributions
E.M. supervised the project. Y.H., N.K., and E.M. designed the study and wrote the manuscript. Y.H. and D.S. performed the experiments, analyzed the data, and equally contributed to this study. All authors reviewed the paper.

Competing interests
The authors declare no competing interests.

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