Lysine Decarboxylase Catalyzes the First Step of Quinolizidine Alkaloid Biosynthesis and Coevolved with Alkaloid Production in Leguminosae

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Lysine decarboxylase (LDC) catalyzes the first-step in the biosynthetic pathway of quinolizidine alkaloids (QAs), which form a distinct, large family of plant alkaloids. A cDNA of lysine/ornithine decarboxylase (L/ODC) was isolated by differential transcript screening in QA-producing and nonproducing cultivars of Lupinus angustifolius. We also obtained L/ODC cDNAs from four other QA-producing plants, Sophora flavescens, Echinosophora koreensis, Thermopsis chinensis, and Baptisia australis. These L/ODCs form a phylogenetically distinct subclade in the family of plant ornithine decarboxylases. Recombinant L/ODCs from QA-producing plants preferentially or equally catalyzed the decarboxylation of L-lysine and L-ornithine. L. angustifolius L/ODC (La-L/ODC) was found to be localized in chloroplasts, as suggested by the transient expression of a fusion protein of La-L/ODC fused to the N terminus of green fluorescent protein in Arabidopsis thaliana. Transgenic tobacco (Nicotiana tabacum) suspension cells and hairy roots produced enhanced levels of cadaverine-derived alkaloids, and transgenic Arabidopsis plants expressing (La-L/ODC) produced enhanced levels of cadaverine, indicating the involvement of this enzyme in lysine decarboxylation to form cadaverine. Site-directed mutagenesis and protein modeling studies revealed a structural basis for preferential LDC activity, suggesting an evolutionary implication of L/ODC in the QA-producing plants.

INTRODUCTION

Alkaloids are one of the most diverse groups of natural products and are found in ~20% of plant species. Many of the ~12,000 known alkaloids produced by plants display potent pharmacological activities and are widely used as pharmaceuticals (Croteau et al., 2000; De Luca and St Pierre, 2000). Alkaloids are derived from the products of primary metabolism with amino acids such as Phe, Tyr, Trp, Orn, and Lys serving as their main precursors (Facchini, 2001). Quinolizidine alkaloids (QAs) are derived from cadaverine, and several hundred structurally related compounds have been identified that are distributed mostly within the Leguminosae (Ohmiya et al., 1995; Michael, 2008). Some QAs exhibit beneficial pharmacological properties, such as cytotoxic, antiarrhythmic, oxytocic, hypoglycemic, and anti- pyretic activities, and thus can be used as drugs (Ohmiya et al., 1995). They can therefore serve as potential starting compounds in the development of new drugs and in pest control for plants (Saito and Murakoshi, 1995).

QAs are synthesized through the cyclization of the cadaverine unit (Golebiewski and Spenser, 1988), which is produced through the action of Lys decarboxylase (LDC; EC 4.1.1.18) (Figure 1), via a postulated enzyme-bound intermediate (Saito and Murakoshi, 1995). The various skeletons of QAs [e.g., bicyclic alkaloids such as (−)-lupinine and (+)-eplupinine or tetracyclic alkaloids such as (+)-multiflorine, (+)-lupanine, and (+)-matrine] are then further modified by tailoring reactions (e.g., dehydrogenation, oxygenation, esterification, and glycosylation) to yield hundreds of structurally related alkaloids (Ohmiya et al., 1995).

However, the molecular mechanism underlying QA biosynthesis is poorly understood, unlike the mechanisms of biosynthesis of other plant alkaloids, such as tropane, isoquinoline, and indole alkaloids (De Luca and St Pierre, 2000; Facchini, 2001; Sato et al., 2007; Kutchan et al., 2008). Very few studies have aimed to characterize the enzymes and regulatory mechanisms involved in QA biosynthesis. These include some preliminary biochemical characterizations performed using crude cell-free extracts (Strack et al., 1991; Saito et al., 1993; Hirai et al., 2000) or purified preparations from Lupinus plants (Suzuki et al., 1994) and the identification of the gene for an acetyltransferase that catalyzes the final step of QA biosynthesis (Okada et al., 2005).

LDC and Om decarboxylase (ODC; EC 4.1.1.17) are key enzymes involved in the formation of cadaverine and putrescine by the decarboxylation of Lys and Orn, respectively. Putrescine is also synthesized by an additional pathway via Arg decarboxylase (EC 4.1.1.19) as elucidated in Arabidopsis thaliana (Hanfrey et al., 2001). ODC is closely related to cell proliferation and is essential for normal cell growth (Pegg, 1988). Cadaverine and putrescine, and other aliphatic polyamines, such as spermidine and spermine,
are known to be involved in a number of growth and developmental processes (Bagni and Tassoni, 2001). Cadaverine is implicated as a precursor not only in the biosynthesis of QAs but also of piperine, lobeline, and other types of alkaloids having piperidine or pyridine rings (Leistner and Spenser, 1973). Unlike plant ODCs, plant LDCs have not been well studied at the molecular level despite their important role in the synthesis of a broad range of alkaloids. It has been suggested that ODC may also have LDC activity in plants that make these types of alkaloids (Sinclair et al., 2004). There are a few studies on crude protein extracts or partially purified LDC from soybean (Glycine max) (Kim et al., 1998; Ohe et al., 2009) and garden lupin (Lupinus polyphyllus) (Hartmann et al., 1980), but no molecular information is available so far for plant LDC.

In this study, we cloned and characterized Lys/Orn decarboxylases (L/ODCs) from QA-producing plants and provide evidence from in vitro and in vivo experiments in tobacco (Nicotiana tabacum) and Arabidopsis that these enzymes are involved in the first step of QA biosynthesis. In addition, we identified an important residue for LDC activity (substantiated by protein modeling and mutation studies), implying an adaptive evolutionary link between the production of QAs and the occurrence of LDC in plants. This research is a breakthrough in the understanding of QA biosynthesis, a topic that has not received much attention, despite the importance of QAs as a large class of plant alkaloids.

RESULTS

Cloning of cDNAs Encoding LDC from QA-Producing Plants

To isolate cDNAs involved in QA biosynthesis, we conducted a differential PCR-select subtraction analysis using cDNAs from the seedlings of two different Lupinus angustifolius cultivars: a QA-producing bitter cultivar (cv Fest) and a sweet cultivar that does not produce QA (cv Uniharvest) (Bunsupa et al., 2011). The bitter and sweet cultivars are similar in flowering time and pod shattering but differ in alkaloid contents and the colors of the flower and seed coat (Oram, 1983). We obtained 71 cDNA fragments specifically expressed in the bitter cultivar and 43 fragments from the sweet cultivars. Among those fragments specifically expressed only in the QA-producing bitter cultivar, three exhibited sequence similarity with plant ODC and were identified as putative LDCs. A full-length cDNA clone L. angustifolius L/ODC (La-L/ODC) was obtained by 5’- and 3’-rapid amplification of cDNA ends. The L/ODC cDNA contained an open reading frame (ORF) of 1320 bp encoding 440 amino acids.

Figure 1. Biosynthetic Pathway of QAs.
Biosynthetic pathway of QAs from L-Lys via cadaverine. LDC is responsible for the first step of biosynthesis.
A phylogenetic tree was constructed by the neighbor-joining method for plant L/ODCs and ODCs from bakers' yeast (Saccharomyces cerevisiae) (Fonzi and Sypherd, 1987), human (Homo sapiens) (Dufe et al., 2007), frog (Xenopus laevis) (Bassez et al., 1990), and a protist (Trypanosoma brucei) (Grishin et al., 1999). All plant L/ODCs were in the same branch, which was further divided into three subgroups: L/ODCs from mosses, monocots, and dicots. La-L/ODC and other L/ODCs from QA-producing plants are closely related to one another and form a distinct clade inside ODCs from other Leguminosae (Figure 2; see Supplemental Figure 1 and Supplemental Data Set 1 online), suggesting a close evolutionary relationship between L/ODCs from QA-producing plants.

L/ODC from QA-Producing Plants Exhibit L/ODC Activity in Vitro

To determine the biochemical function of La-L/ODC, Sf-L/ODC, and Ek-L/ODC, the ORFs of these genes were inserted into a glutathione S-transferase (GST)-tag Escherichia coli expression vector as an N-terminal fusion. The observed molecular masses of tag-purified/cleaved recombinant proteins on SDS-PAGE matched well with the predicted 47-, 49-, and 49-kD sizes for La-L/ODC, Sf-L/ODC, and Ek-L/ODC, respectively (see Supplemental Figure 2A online). Decarboxylation activities were assayed with these recombinant proteins using L-Orn and L-Lys as substrates. The optimal pH for both ODC and LDC activities was evaluated, and pH 7.5 was determined to be the optimal value for both reactions.

The three recombinant L/ODCs exhibited decarboxylase activities using both substrates with relatively similar kinetic properties. This is in contrast with the reported decarboxylase activity of N. glutinosa, which exhibits an extremely low activity to L-Lys (Lee and Cho, 2001). The apparent $k_{cat}$ values ranged from 0.73 to 1.16 s$^{-1}$ for L-Orn and 1.18 to 2.32 s$^{-1}$ for L-Lys, while the $K_m$ values ranged from 1.05 to 1.53 mM for L-Orn and 2.10 to 3.85 mM for L-Lys (Table 1). Comparison of the catalytic efficiency ($k_{cat}/K_m$) of these L/ODCs with respect to the substrates revealed that they preferentially or equally catalyzed the decarboxylation of L-Lys over L-Orn. Because the results of our previous experiment showed that La-L/ODC possesses decarboxylase activity toward both L-Lys and L-Orn, we asked if they compete with each other. A competitive assay was performed by varying the concentrations of L-Lys (0.5 to 3.0 mM) in the presence (2mM) and absence of L-Orn. The double-reciprocal plots of 2.0 mM L-Lys and L-Orn at 1/V$_{max}$ on the y axis, showing a competitive reaction pattern (see Supplemental Figure 3A online). In addition, we also tested if the decarboxylase activity of La-L/ODC was inhibited by α-difluoromethylornithine (α-DFM0), an ODC suicide inhibitor. The observed LDC activity was distinctly inhibited by α-DFM0 in a dose-dependent manner (see Supplemental Figure 3B online). These results demonstrate that the catalytic site of La-L/ODC for the two substrates is identical.

To obtain kinetic information on ODCs from non-QA-producing leguminous plants, recombinant proteins Gm-ODC from soybean and Lj-ODC from Lotus japonicus were assayed for L/ODC activity. The recombinant Gm-ODC and Lj-ODC exhibited ODC and LDC activities with optimal pH at 7.5 to 8.0. However, Lj-ODC exhibited very weak LDC activity. The activities of Gm-ODC and Lj-ODC were very low compared with La-L/ODC, presumably because of the insoluble nature of the recombinant proteins.

Site-Directed Mutation and Protein Modeling Locates a Critical Amino Acid Residue for Substrate Specificity

Sequence alignment of eukaryotic L/ODCs indicated that the amino acid sequences (positions 339 to 344) of presumable substrate binding sites responsible for holding the substrate/product were completely conserved in L/ODCs from QA-producing plants (Figure 2; see Supplemental Figure 1 online). Three-dimensional structures of ODCs and earlier site-directed mutation studies (Grishin et al., 1999; Lee and Cho, 2001; Jackson et al., 2004) suggested the catalytic mechanism and substrate binding sites. Two amino acid residues (Met 341 and Phe 344) that are conserved within L/ODCs from QA-producing plants were characteristically altered from the consensus residues of other eukaryotic ODCs. These two amino acid residues of La-L/ODC were mutated to the other eukaryotic ODC consensus ones, thus generating the mutants La-L/ODC-M341L, La-L/ODC-F344H, and La-L/ODC-F344Y (Figure 2). The purified mutant proteins (see Supplemental Figure 2B online) were tested for decarboxylase activity (Table 1). Of these, the mutation of Phe 344 proved to be the most critical for substrate binding. The mutations of La-L/ODC-F344H resulted in an increase of $K_m$ for L-Lys by 41-fold (97.84 mM/2.37 mM) and only by 1.5-fold (1.53 mM/1.05 mM) for L-Orn, whereas the mutation of La-L/ODC-F344Y resulted in an increased $K_m$ for L-Lys by 2.2-fold (5.26 mM/2.37 mM) and by 1.5-fold (1.59 mM/1.05 mM) for L-Orn compared with the wild-type La-L/ODC, indicating the importance of Phe 344 for acceptance of L-Lys as the substrate. The mutations of La-L/ODC-M341L caused only minor changes to the $K_m$ for both substrates (Table 1). Interestingly, the mutation of La-L/ODC-F344H not only increased $K_m$ value but also the turnover number for L-Lys. The mutation in this position may affect the overall structure of La-L/ODC protein, thus leading to changes in overall enzymatic properties, especially in the catalytic step, of which the precise mechanism(s) still needs to be clarified.

The tertiary structure of La-L/ODC was predicted by SWISS-MODEL (http://swissmodel.expasy.org/) using human ODC (Hs-ODC) (Dufe et al., 2007), with 45% amino acid identity to La-L/ODC as a template for protein modeling. Overall, the predicted protein structure of La-L/ODC was similar to the eukaryotic ODC structures that have been determined previously, including T. brucei (Tb-ODC) (Jackson et al., 2000) and mouse (Mus musculus) (Kern et al., 1999). The predicted La-L/ODC structure was a symmetrical homodimer, formed by a head-to-tail interaction between the barrel of one domain and the sheet domain of the other (see Supplemental Figure 4A online). Conserved amino acid residues of the binding sites of pyridoxal 5'-phosphate (PLP) and the substrate were superimposed well on to the Hs-ODC crystal structure with the exception of the 310-helix at the active site, which is responsible for holding the PLP-bound putrescine/cadaverine (see Supplemental Figure 4B online). This 310-helix
was slightly shorter in the La-L/ODC structure compared with Tb-ODC and Hs-ODC (Figures 3A and 3B).

We also generated a structural model of the La-L/ODC mutants and *N. glutinosa* ODC (Ng-ODC) (Lee and Cho, 2001) by the same method. The predicted La-L/ODC-F344H and Ng-ODC protein structures showed a longer 310-helix like Hs-ODC, while the other two mutants showed a shorter 310-helix as for the wild-type La-L/ODC (Figures 3C and 3D; see Supplemental Figure 4C online). These results suggest a critical role for the length of the 310-helix in accepting L-Lys as the substrate. The length of the 310-helix is determined by the residue at 344, and it presumably changes the cavity size of the substrate binding pocket (see Discussion).

### Tissue-Specific and Cultivar-Specific Expression of *L. angustifolius* L/ODC

The L/ODC transcript level was examined in different tissues (young and mature leaves, cotyledons, hypocotyls, and roots) of bitter and sweet cultivars of *L. angustifolius* using quantitative RT-PCR (Figure 4A). In the bitter cultivar, the L/ODC transcript level was the highest in the young leaves and was barely detected.

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**Figure 2.** Phylogenetic Tree of Eukaryotic L/ODCs.

Unrooted neighbor-joining phylogenetic tree of selected eukaryotic L/ODCs. The active-site amino acid residues that differ between LDC and ODC proteins are indicated for each sequence of the tree. Residue number is based on La-L/ODC numbering. Green boxes and red triangles indicate L/ODC and ODC enzymes, respectively, whose biochemical properties have been investigated. Magenta lines indicate quinolizidine alkaloid–producing plants. Bootstrap values (1000 replicates) are shown above each branch. Accession numbers of enzymes are listed in Supplemental Table 2 online.
in other organs. No detectable level of expression was observed in the tissues of the sweet cultivar (Figure 4A).

A genomic PCR was performed using the genomic DNA extracted from young leaves of both bitter and sweet cultivars of *L. angustifolius* using the same specific primers used for quantitative RT-PCR. The fragment obtained from genomic PCR was 1.3 kb in both the bitter and sweet cultivars, indicating that there is no intron inside the *L/ODC* gene in the *L. angustifolius* genome (Figure 4B). The sequences of these genomic fragments from both the bitter and sweet cultivars were identical with that of the cDNA from the bitter cultivar. A DNA gel blot analysis to look for nucleotide identities showed that there is at least one copy of *L/ODC* with identical nucleotide sequence (of at least 1.3-kb span) in both bitter and sweet cultivars of *L. angustifolius* (Figure 4C).

**L/ODC Is Targeted to Plastids**

The probable localization of *L. angustifolius* L/ODC protein was analyzed using the program WoLF PSORT (http://wolfpsort.org/), which predicted that this protein is most likely found in the chloroplast (probability score for chloroplast: 10.0, nucleus; 2.0, cytosol; 2.0). However, analysis of the N-terminal sequence for chloroplast targeting using ChloroP v1.1 (http://www.cbs.dtu.dk/services/ChloroP/), SignalP v3.0 (http://www.cbs.dtu.dk/services/SignalP/), and TargetP v1.1 (http://www.cbs.dtu.dk/

### Table 1. Kinetic Parameters of L/ODCs and La-L/ODC Mutant Proteins

<table>
<thead>
<tr>
<th>Protein</th>
<th>LDC</th>
<th>ODC</th>
<th>LDC</th>
<th>ODC</th>
<th>LDC</th>
<th>ODC</th>
<th>LDC</th>
<th>ODC</th>
<th>Ratio of $k_{cat}/K_m$</th>
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<tbody>
<tr>
<td>La-L/ODC_wild type</td>
<td>2.37</td>
<td>1.05</td>
<td>1.49</td>
<td>1.14</td>
<td>1.180</td>
<td>0.91</td>
<td>433</td>
<td>859</td>
<td>0.500</td>
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<tr>
<td>Stf-L/ODC</td>
<td>2.10</td>
<td>1.53</td>
<td>2.94</td>
<td>1.46</td>
<td>2.320</td>
<td>1.16</td>
<td>1108</td>
<td>755</td>
<td>1.470</td>
</tr>
<tr>
<td>Ek-L/ODC</td>
<td>3.85</td>
<td>1.32</td>
<td>2.42</td>
<td>0.76</td>
<td>1.910</td>
<td>0.73</td>
<td>469</td>
<td>454</td>
<td>1.030</td>
</tr>
<tr>
<td>Ng-ODC*</td>
<td>1.59</td>
<td>0.56</td>
<td>1.52×10⁻⁴</td>
<td>1.25×10⁻²</td>
<td>0.007</td>
<td>0.58</td>
<td>4.4</td>
<td>1035</td>
<td>0.004</td>
</tr>
<tr>
<td>La-L/ODC-M341L</td>
<td>2.49</td>
<td>1.04</td>
<td>4.90</td>
<td>0.37</td>
<td>3.880</td>
<td>0.29</td>
<td>1558</td>
<td>279</td>
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<tr>
<td>La-L/ODC-F344Y</td>
<td>5.26</td>
<td>1.59</td>
<td>7.99</td>
<td>0.51</td>
<td>6.330</td>
<td>0.40</td>
<td>1203</td>
<td>253</td>
<td>4.750</td>
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<tr>
<td>La-L/ODC-F344H</td>
<td>97.84</td>
<td>1.53</td>
<td>181.82</td>
<td>2.90</td>
<td>143.980</td>
<td>2.29</td>
<td>1471</td>
<td>1496</td>
<td>0.980</td>
</tr>
</tbody>
</table>

All experiments were performed in 50 mM potassium phosphate buffer, pH 7.5. Kinetic parameters were calculated from mean values ($n = 3$ to 4).

*Recalculated from reported values for *N. glutinosa* L/ODC (Lee and Cho, 2001).

A genomic PCR was performed using the genomic DNA extracted from young leaves of both bitter and sweet cultivars of *L. angustifolius* using the same specific primers used for quantitative RT-PCR. The fragment obtained from genomic PCR was ~1.3 kb in both the bitter and sweet cultivars, indicating that there is no intron inside the L/ODC gene in the *L. angustifolius* genome (Figure 4B). The sequences of these genomic fragments from both the bitter and sweet cultivars were identical with that of the cDNA from the bitter cultivar. A DNA gel blot analysis to look for nucleotide identities showed that there is at least one copy of L/ODC with identical nucleotide sequence (of at least 1.3-kb span) in both bitter and sweet cultivars of *L. angustifolius* (Figure 4C).

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services/TargetP/) programs showed that no transit peptides were present.

To determine the subcellular localization of L/ODC, we examined the transient expression of a green fluorescent protein (GFP) fused with the N-terminal 100 amino acids of \textit{L. angustifolius} L/ODC protein (La-L/ODC100-GFP) in \textit{Arabidopsis} plants using the particle bombardment method. Two plastid-targeting fusion proteins, a fusion of ribulose-1,5-bis-phosphate carboxylase/oxygenase (Rubisco) transit peptide (Krebbers et al., 1988) with GFP (Rubisco-TP-GFP) and a fusion of transit peptide of rice (\textit{Oryza sativa}) \textit{Waxy} (Kitajima et al., 2009) with red fluorescent protein from \textit{Discosoma} sp (Wx-TP-DsRed), were used as references for plastid localization. A Bsas 3:1 (spinach mitochondrial b-substituted Ala synthase) transit peptide fused to GFP (Bsas 3:1-TP-GFP) (Noji et al., 1998) and the pTH2 vector (Chiu et al., 1996) were used as references for mitochondrion and cytosol localizations, respectively. La-L/ODC100-GFP and Rubisco-TP-GFP were simultaneously expressed with Wx-TP-DsRed in \textit{Arabidopsis} leaves. As shown in Figure 5A, L/ODC100-GFP fluorescence overlapped with the plastid fluorescence visualized by Wx-TP-DsRed. This localization pattern was identical with Rubisco-TP-GFP, which overlapped with Wx-TP-DsRed (Figure 5B). Thus, the N-terminal 100 amino acids of L/ODC was capable of translocating the passenger protein to plastids, indicating the plastidic localization of L/ODC, in agreement with the previous study of localization of LDC enzymatic activity in \textit{L. polyphyllus} (Hartmann et al., 1980).

**Overexpression of \textit{L. angustifolius} L/ODC Enhances Production of Cadaverine-Derived Alkaloids in Tobacco Hairy Roots and BY-2 Cells**

To confirm whether L/ODC functions as an LDC in alkaloid production in vivo, we overexpressed \textit{L. angustifolius} L/ODC in stable transformants under the control of the 3SS promoter in tobacco hairy root cultures and BY-2 suspension cells. RT-PCR analysis confirmed the expression of L/ODC in all overexpressing lines. The alkaloid levels in the transgenic tobacco lines expressing La-L/ODC were analyzed by HPLC-photodiode array detection and HPLC–mass spectrometry (MS). In the hairy roots, the contents of anabasine and anatalline (Figure 6) increased by 28.9% (P = 0.023) and 25.1% (P = 0.015), respectively, compared with the corresponding levels in the control lines (\textit{b-glucuronidase [GUS]}-expressing lines). By contrast, the level of nicotine remained constant or even decreased slightly (Figure 6). In addition, the amounts of L-Lys, L-Orn, cadaverine, and putrescine in the hairy roots of transgenic tobacco expressing La-L/ODC were analyzed by capillary electrophoresis (CE)-MS. The

![](image)

**Figure 4. Molecular Analysis of \textit{L. angustifolius} L/ODC.**

(A) Accumulation of La-L/ODC mRNA in various tissues of two cultvars, bitter (cv Fest) and sweet (cv Uniharvest), of \textit{L. angustifolius} plants by quantitative RT-PCR analysis. (B) Genomic PCR analysis of La-L/ODC. DNA was extracted from young leaves of both cultivars and subjected to genomic PCR analysis using the same primer pair that was used for quantitative RT-PCR analysis. Lane M, \textit{PstI} marker. (C) DNA gel blot analyses of La-L/ODC. Fifteen micrograms of genomic DNA of both cultivars was digested with EcoRl (lane I), EcoRV (lane II), and \textit{HindIII} (lane III). The La-L/ODC ORF was used as a probe for hybridization. In both cultivars, at least one EcoRI- and \textit{HindIII}-digested band was hybridized at 4.4 and 4.2 kb, respectively.

![](image)

**Figure 5. Plastid Localization of \textit{L. angustifolius} L/ODC N-Terminal 100 Amino Acids Fused with GFP in \textit{Arabidopsis} Leaves.**

(A) \textit{Arabidopsis} leaves expressing \textit{L. angustifolius} L/ODC100-GFP and Wx-TP-DsRed. L/ODC100-GFP, green (left); Wx-TP-DsRed, red (middle); GFP and DsRed merged image (right). (B) Control experiments. Rubisco-TP-GFP (left), Wx-TP-DsRed (middle), and GFP and DsRed merged image (right). Bars = 10 \textmu m.
levels of cadaverine and putrescine slightly increased in the transgenic lines by 24.1% (P = 0.126) and 12.1% (P = 0.098), respectively (see Supplemental Figure 5 online).

The alkaloid levels in the transformed tobacco BY-2 suspension cells expressing La-L/ODC were determined after treatment with methyl jasmonate. Compared with the levels of anabasine and anatalline in the control lines, the levels in the La-L/ODC expressing cells increased by 102.9 and 66.4% (P = 0.0437), while the those of nicotine even decreased by 29.1% (Figure 6).

**Expression of *L. angustifolius* L/ODC in *Arabidopsis* Promotes Production of Cadaverine**

In transgenic *Arabidopsis* plants expressing *L. angustifolius* L/ODC, the contents of L-Lys, L-Orn, cadaverine, and putrescine were analyzed by CE-MS. Ectopic expression in several

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**Figure 6. Overexpression of *L. angustifolius* L/ODC in Tobacco Hairy Roots and BY-2 Suspension Cells.**

Biosynthetic pathway of tobacco alkaloids with the average contents of tobacco alkaloids in La-L/ODC-overexpressing (LDC) hairy roots and BY-2 cells and compared with the corresponding levels in the control lines expressing a bacterial GUS gene. For hairy roots, each bar represents the mean ± SE of average contents of tobacco alkaloids in six independent L/ODC-overexpressing lines (biological replicates; n = 4 for each line) and four independent GUS-expressing lines (biological replicates; n = 4 for each line). For tobacco BY-2 cells, each bar shows the mean ± SE of average contents of tobacco alkaloids in eight independent L/ODC-overexpressing BY-2 cells and five independent control lines expressing GUS gene; for BY-2 cells, both cell lines were treated with methyl jasmonate. Student’s one-tailed t test, *P value < 0.05. Broken lines indicate unresolved reactions. DW, dry weight; FW, fresh weight.
overexpressing lines of La-L/ODC Arabidopsis showed the accumulation of cadaverine, which was not detected in the control plants (wild-type and GUS-overexpressing lines) (Figure 7A). La-L/ODC expression was positively correlated with the levels cadaverine (P = 0.000015 and r = 0.745) but showed no correlation with putrescine (P = 0.316 and r = 0.103) (Figures 7A and 7B; see Supplemental Figures 6A and 6B online). The levels of L-Lys and L-Orn were negatively correlated with the expression levels of La-L/ODC (P = 0.008, r = −0.485 and P = 0.030, r = −0.388, respectively) (see Supplemental Figures 6C and 6D online). Furthermore, the levels of L-Lys and L-Orn in the La-L/ ODC–overexpressing lines were substantially higher than the corresponding levels in the control (Figure 7C). No other remarkable changes were observed in the rest of metabolites detected by CE-MS (see Supplemental Figure 7 online).

The Patterns of Cadaverine-Related Metabolites Differ in the Bitter and the Sweet Cultivars of L. angustifolius

A study by Hirai et al. (2000) showed no QAs in young leaves (3 weeks old, at the same stage used in this study) of the sweet cultivar of L. angustifolius. In this study, we set out to investigate the contents of L-Lys and L-Orn, the potential biosynthetic precursors of QAs, and putative cadaverine conjugates (their storage form) in the sweet and bitter cultivars. To reveal the relationship between L-Lys and cadaverine and the production of QAs, the contents of L-Lys, L-Orn, putrescine, and cadaverine were determined by CE-MS in both the bitter and sweet cultivars of L. angustifolius. The amounts of L-Lys and L-Orn in the bitter cultivar were ~1.6 and 1.8 times higher than those of sweet cultivar (Table 2). By contrast, the sweet cultivar contained ~1.7 times more putrescine than the bitter one. It is noteworthy that cadaverine was not found in either the bitter or the sweet cultivars; however, the content of putative cadaverine conjugates in the bitter cultivar was 3.2 times higher than in the sweet cultivar, while the amount of putative putrescine conjugates was similar in both cultivars. It is evident that the bitter cultivar more actively accumulated the potential biosynthetic precursor (L-Lys) and its storage form (the cadaverine conjugates) than the sweet cultivar along with the production capability of QAs.

DISCUSSION

Physiological Roles of LDC in QA-producing Plants

In this study, we identified three L/ODCs from QA-producing plants and found that these L/ODCs show similar LDC and ODC activities, unlike the known ODCs that accept only Orn (Boeker and Fischer, 1983). Although these L/ODCs from QA-producing plants could catalyze the decarboxylation of both L-Lys and L-Orn with a nearly equal efficiency in vitro, the availability of substrates and their compartmentalization might explain the in vivo function of this enzyme as an LDC. The amount of L-Lys is 45 times higher than L-Orn in L. angustifolius (Table 2). Since protein–amino acid biosynthesis is strictly regulated in the plant cell (Saito et al., 1993; Stepansky et al., 2006), the controlled high content of L-Lys in the bitter cultivar of L. angustifolius may facilitate an efficient formation of cadaverine leading to alkaloid biosynthesis in vivo.

Our results indicate that L. angustifolius L/ODC is localized in the chloroplast, where diaminopimelate decarboxylase, the enzyme involved in the last step of Lys biosynthesis, is thought to be localized (Mazelis et al., 1976). This result is in concurrence with previous studies in Lupinus plants showing that the activity of the Bitter and the Sweet Cultivars of L. angustifolius

Figure 7. Overexpression of L. angustifolius L/ODC in Arabidopsis Plants.

(A) The contents of cadaverine in wild-type and transgenic Arabidopsis plants expressing GUS or L/ODC (LDC). FW, fresh weight.
(B) Quantitative RT-PCR analysis of La-L/ODC transcript in wild-type (WT), GUS-expressing (GUS), and La-L/ODC–overexpressing (LDC) Arabidopsis plants. The average contents of Lys, Orn, cadaverine, and putrescine in wild-type and transgenic Arabidopsis plants expressing GUS or La-L/ODC (LDC). Expression of La-L/ODC was measured by quantitative RT-PCR, and the expression ratio was calculated based on the β-tubulin gene. For (A) and (B), each bar represents the mean ± SE of four to six biological replicates for each independent line. For (C), each bar represents the mean ± SE of the wild type (biological replicates; n = 4), six independent La-L/ODC–overexpressing lines (biological replicates; n = 4 to 6 for each line), and four independent GUS-expressing lines (biological replicates; n = 6 for each line). Student’s one-tailed t test, *P value < 0.05. ND, not detected.
of LDC is associated with chloroplasts (Wink and Hartmann, 1982) and that the majority of QAs are synthesized in shoot tissues (Lee et al., 2007a). This localization of La-L/ODC is in agreement with the localization of ODC in chloroplasts (at least partial) (Andreadakis and Kotzabasis, 1996; Tassoni et al., 2003), supporting a hypothesis that L/ODC evolved from an ancestor ODC (as discussed below). Additionally, it was reported that the metabolic action of bacterial LDC in transgenic plants was improved by its fusion to a chloroplast-targeting signal sequence (Herminghaus et al., 1996).

The ODC transcript levels in plants such as tomato (Solanum lycopersicum) (Kwak and Lee, 2001), soybean (Dels et al., 2005), and devil’s trumpet (Michael et al., 1996) showed high transcript levels in developing tissues and expanding cells, such as those in hypocotyls and roots. The transcripts of L. angustifolius L/ODC accumulated coincidently with the production of QAs, exclusively in the young leaves of the bitter cultivar (Figure 4A); this accumulation coincidently with the production of QAs, excluded from the control plants, thus confirming the involvement of this enzyme in Lys decarboxylation to form cadaverine. In the literature (Shoji and Hashimoto, 2008; and references cited therein), anabasine has been confirmed to be derived from cadaverine. The in vivo experiments with transgenic tobacco (hairy roots and BY-2 suspension cells) showed the enhanced production of anabasine in the LDC-overexpressing tobacco cells, clearly indicating that L/ODC acts as an LDC to produce cadaverine from L-Lys in vivo, thus leading to the formation of cadaverine-derived alkaloids.

Taken together, these results support the conclusion that the identified L/ODC catalyzes the decarboxylation of L-Lys to form cadaverine leading to the synthesis of QAs in Lupinus plants.

### Substrate Preference and Active Site Structure

The competition and inhibition studies suggest that the catalytic site of L/ODC for the recognition of both substrates is identical (see Supplemental Figure 3 online). The evidence for both LDC and ODC activities being ascribed to the same active site have also been reported in a L/ODC from the bacterium Selenomonas ruminantium, which is capable of decarboxylating both L-Lys and L-Orn with similar kinetic properties, and in the ODC from N. glutinosus (Takatsuka et al., 1999; Lee and Cho, 2001). Unlike other eukaryotic ODCs that are almost exclusively specific for L-Orn over L-Lys (Lee et al., 2007b), the three L/ODCs from QA-producing plants proteins exhibited decarboxylase activities toward both L-Lys and L-Orn, with similar kinetic properties. These results imply that the active site of L/ODC is relatively tolerant to variable side chains of substrate (C5 for Lys and C4 for Orn) for the intermediary Schiff-base formation.

Evaluation of the L/ODC-F344H mutant enzyme activity points out that Phe-344 residue is essential for the expansion of substrate acceptability to L-Orn and L-Lys. Phe-344 is positioned next to the essential Asp-343 residue, which interacts with N(ε) of PLP-bound putrescine as indicated in the human ODC (Grishin et al., 1999). In the authentic ODCs accepting only Orn, this Phe-344 is replaced with His (Figure 2B). Because the decarboxylation reaction needs a fixed quinoid intermediate, the size and polarity of the cavity of the reaction intermediate is very important for forming the correct C(α)-N configuration (Grishin et al., 1999). The amino acid residue (Phe or His-344) adjacent to the Asp-343 responsible for the correct positioning of the quinoid intermediate must have a critical role for the substrate acceptability (i.e., Lys and Orn). In fact, our results show that the mutation of L/ODC-F344H resulted in a remarkable increase of K<sub>m</sub> for L-Lys but only in a minimal change for L-Orn, indicating the importance of Phe-344 for the acceptance of L-Lys as a substrate. However, the L/ODC-F344Y mutant has a lesser effect on the K<sub>m</sub> values for these two substrates. Interestingly, Gm-ODC has Tyr instead of zipper protein, and coronatine-insensitive 1 (Bunsupa et al., 2011). Since the enzyme activity and the transcript level of an acyltransferase committed in the terminal step of QA biosynthesis do not differ in bitter and sweet cultivars (Hirai et al., 2000; Okada et al., 2005), this finding is interesting in terms of the regulation of the whole biosynthetic pathway.

In the in vivo experiment with Arabidopsis plants expressing L. angustifolius L/ODC showed the accumulation of cadaverine, which was not detected in the control plants, thus confirming the involvement of this enzyme in Lys decarboxylation to form cadaverine. In the literature (Shoji and Hashimoto, 2008; and references cited therein), anabasine has been confirmed to be derived from cadaverine. The in vivo experiments with transgenic tobacco (hairy roots and BY-2 suspension cells) showed the enhanced production of anabasine in the LDC-overexpressing tobacco cells, clearly indicating that L/ODC acts as an LDC to produce cadaverine from L-Lys in vivo, thus leading to the formation of cadaverine-derived alkaloids.

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### Table 2. Levels of Polyamines and QAs in Leaves of the Bitter and Sweet Cultivars of L. angustifolius

<table>
<thead>
<tr>
<th>Compound</th>
<th>Bitter Cultivar</th>
<th>Sweet Cultivar</th>
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</thead>
<tbody>
<tr>
<td>L-Orn</td>
<td>4.91 ± 0.12</td>
<td>2.79 ± 0.15</td>
</tr>
<tr>
<td>L-Lys</td>
<td>220.53 ± 10.02</td>
<td>141.18 ± 12.44</td>
</tr>
<tr>
<td>Putrescine</td>
<td>5.28 ± 0.46</td>
<td>9.04 ± 0.74</td>
</tr>
<tr>
<td>Cadaverine</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>Putrescine conjugates&lt;sup&gt;a&lt;/sup&gt;</td>
<td>9.84 ± 7.51</td>
<td>10.62 ± 6.24</td>
</tr>
<tr>
<td>Cadaverine conjugates&lt;sup&gt;b&lt;/sup&gt;</td>
<td>13.34 ± 1.81</td>
<td>4.17 ± 1.03</td>
</tr>
<tr>
<td>QAs&lt;sup&gt;c&lt;/sup&gt;</td>
<td>1218 (µg g&lt;sup&gt;-1&lt;/sup&gt; FW&lt;sup&gt;-1&lt;/sup&gt;)</td>
<td>&lt;1.0 (µg g&lt;sup&gt;-1&lt;/sup&gt; FW&lt;sup&gt;-1&lt;/sup&gt;)</td>
</tr>
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</table>

Data represent means ± so the polyamines in the mixture of leaves from 10 L. angustifolius plants (3 weeks old) in each cultivar (analytical replicate; n = 6). FW, fresh weight; ND, not detected.

<sup>a</sup>Putative putrescine conjugates are predicted based on mass-to-charge ratio (m/z) from CE-MS, including cinnamoyl-, caffeoyl-, dicumaroyl-, dicaffeoyl- and diferuloyl-putrescine with the corresponding m/z 219.15, 251.15, 381.19, 413.18, and 441.19, respectively. The total amount is calculated on the mass response factor of putrescine as the standard.

<sup>b</sup>Putative cadaverine conjugates are predicted based on m/z from CE-MS, including cinnamoyl-, coumaroyl-, sinapoyl-, dicumaroyl-, dicaffeoyl- and diferuloyl-cadaverine with the corresponding m/z 233.16, 249.15, 309.18, 395.19, 427.19, and 455.24, respectively. The total amount is calculated on the mass response factor of cadaverine as the standard.

<sup>c</sup>Reported values of QAs in leaves of the bitter and sweet cultivars of L. angustifolius (3 weeks old) from a previous study (Hirai et al., 2000).
His at position 344 (La-L/ODC numbering). This suggests that Gm-ODC might catalyze both L-Lys and L-Orn with similar extent. Indeed, this speculation is supported by the study of recombinant Gm-ODC protein, which exhibits both LDC and ODC activities. By contrast, another leguminous plant enzyme LJ-ODC having His-344 exhibited only ODC activity, indicating the diversification of L/ODC in Leguminosae plants.

The molecular modeling data further explain why L/ODCs from QA-producing plants are able to efficiently catalyze the reaction for L-Lys. Predicted protein structures of the L/ODCs showed that the length of the $3_{10}$-helix of La-L/ODC-WT is shorter than those of La-L/ODC-F344H, Ng-ODC, and Hs-ODC (Figure 3; see Supplemental Figure 4C online). The $3_{10}$-helix is critical in determining the size of the cavity of catalytic site because Asp-343 adjacent to the $3_{10}$-helix forms a hydrogen bond with the pytrecine bound to PLP. Phe-344 does not form an extra helix next to the $3_{10}$-helix as His-344 does, thereby accommodating the one-carbon larger side chain of L-Lys. The La-L/ODC-F344H mutant, as with Hs-ODC and Ng-ODC, has a longer $3_{10}$-helix and was incapable of accepting L-Lys, but had a minimal effect on ODC activity. A similar mechanism of acceptance of substrates by varied cavity size of substrate pocket was reported by x-ray crystal structure studies in Vibrion vulnificus L/ODC (Vv-L/ODC), which decarboxylases both L-Lys and L-Orn with similar $K_m$ and $V_{\text{max}}$ values (Lee et al., 2007b). The ability of Vv-L/ODC to interact with L-Orn appears to be due to the positioning of a bridging water molecule that participates in the binding of the shorter putrescine ligand, but this water molecule is absent in the cadaverine-bond structure. A similar situation may be conceivable for La-L/ODC to accept both Lys and Orn substrates. Further studies on x-ray structural analysis of La-L/ODC should provide more precise insight into the structural basis for this dual function.

The results of the mutation study indicate that Phe-344 is a key residue for L. angustifolius L/ODC to exhibit substrate promiscuity for Lys and Orn. These results reveal how plants have been opportunistic in adapting primary metabolic processes to permit the diversification of secondary metabolism, in this case, substrate promiscuity. Accordingly, the mutation in this position might possibly be found not only in QA-producing Leguminosae plants but also in other plant species that produce a variety of cadaverine-derived alkaloids, such as pomegranate (Punica granatum), lobelia (Lobelia inflata), and black pepper (Piper nigrum), which produce pelletierine, lobeline, and piperine, respectively. This is an interesting question on possible convergent evolution of plant species modifying their synthesis of specialized metabolites for adaptation to their environment (as discussed in Pichersky and Lewinsohn, 2011) and needs to be addressed in future studies.

Evolution of LDC in QA-Producing Plants

Identification of the La-L/ODC answers a longstanding question about the molecular entity of LDC activity in plants. There was only a speculation that a member of group IV of PLP-dependent amino acid decarboxylases might be responsible for the production of cadaverine from Lys (Sandmeier et al., 1994). Our study provides the evidence for the molecular entity of plant LDC from several QA-producing plants.

QAs are typically found in some phylogenetically related tribes of the Leguminosae (Wink, 1992), but they have also been found in unrelated genera of Chenopodiaceae, Berberidaceae, Ranunculaceae, Scrophulariaceae, and Solanaceae (Kinghorn and Balandrin, 1984). The occurrence of LDC in the plant species of this study coincides with the production of QAs. Many studies have reported that duplication of essential genes of primary metabolism is an important basis for gene recruitment in secondary metabolism (Nakajima et al., 1993; Hashimoto et al., 1998; Ober and Kaltenegee, 2009; Teuber et al., 2007; de Kraker and Gershenzon, 2011). A good example would be the identification of the enzyme homoserpinimide synthase, which is involved in pyrrolizidine alkaloid biosynthesis, as a modified version of deoxypusine synthase, an enzyme that is involved in the primary metabolism of transalational regulation (Ober and Hartmann, 1999). The results of this study (i.e., no indication of the presence of an additional paralogous gene of L/ODC in the genome of L. angustifolius) indicate that the novel L/ODCs have presumably been derived from common plant ODCs with a slight modification as an adaptation aimed at broadening substrate specificity under a certain environmental stress as discussed in other organisms (Ganfornina and Sanchez, 1999; Harlin-Cognato et al., 2006). The second possible scenario is that gene duplication took place from an ancestor ODC, one of the duplicated genes having original ODC activity was deleted, and the other gene evolved to L/ODC by expanding substrate acceptability, as often reported in the evolution of plant genes (Simillion et al., 2002; Blanc and Wolfe, 2004). ODC is not an enzyme absolutely necessary in plants as Arabidopsis does not have ODC (Hanfrey et al., 2001). Further research is required to address the evolutionary consequence of how specialized secondary metabolism has evolved to expand plants’ chemical diversity.

In this context, it is worthwhile noting that the changes in the possible first amino acid (from His-344 in common ODC to Tyr-344 in soybean ODC) and the second amino acid (from Tyr-344 to Phe-344 in QA-producing LDC) were a result of a single nucleotide replacement (a C-to-T transition in the former and an A-to-T transversion in the latter). Such a critical mutation might take place just at two subsequent minimal nucleotide replacements during the evolution. It is less likely but possible that the changes from His to Tyr and His to Phe independently evolved during evolution. The change from His to Phe needs two nucleotide replacements at the same time, but only one nucleotide replacement is needed for the change from Tyr to Phe. A similar change in substrate specificity attributable to a single amino acid difference was reported in the O-methyltransferases involved in isoquinoline alkaloid and phenylpropanoid biosynthesis (Frick and Kutchan, 1999; Gang et al., 2002).

In this study, we demonstrated that the isolated LDC genes are involved in the decarboxylation of L-Lys, which is the first step of QA biosynthesis in QA-producing plants. In addition, we also demonstrated a point mutation responsible for substrate promiscuity that is supposed to have occurred via adaptive evolution of LDC within the QA-producing plants. These findings can serve as the basis for future research on not only QA biosynthesis but also on the control of plant alkaloid production. Since cadaverine-derived alkaloids have potential applications in the
development of new drugs and in pest control in plants, a further trial for metabolic engineering by combinatorial biochemistry (Oksman-Caldentey and Inzé, 2004) should use the LDC genes.

**METHODS**

**Plant Materials**

Seeds of *Lupinus angustifolius* cv Fest (QA-producing bitter cultivar) and cv Uniharvest (sweet cultivar that does not produce QAs) were from Colin G. Smith (The Australian Lupin Collection Crop Improvement Institute, Australia). Seeds of soybean (*Glycine max*; B01151) and *Lotus japonicus* (Gifu) were from National BioResource Project (Japan) and Hiroshi Sudo (Hoshi University, Japan), respectively. *Sophora flavescent*, *Echinopsphora koreensis*, *Thermopsis chinensis*, and *Baptisia australis* were obtained from the Medicinal Plant Gardens of the Graduate School of Pharmaceutical Sciences at Chiba University, Japan.

**Cloning of L/ODC cDNA from QA-Producing Plants**

The bitter cultivar-specific cDNA fragments were profiled by PCR-select cDNA subtraction (Clontech; PCR-Select cDNA Subtraction Kit) between bitter and sweet cultivars in *L. angustifolius*. Among 71 bitter-specific cDNA fragments, three fragments showed homology to ODC in deduced amino acid sequences. The cDNA encoding La-ODC was isolated by primers that were designed from the nucleotide sequence of these three fragments (see Supplemental Table 1 online for the primers used in this study). Sf-L/ODC, Ek-L/ODC, Tc-L/ODC, and Ba-L/ODC cDNAs were isolated by PCR using degenerate primers designed based on the La-L/ODC and other plants ODC sequences. Each cDNA fragment was obtained with the first-strand cDNA prepared from young leaves as the template and the full-length cDNAs were obtained using 5'- and 3'-rapid amplification of cDNA ends (TakaRa).

**Phylogenetic Analysis**

Phylogeny and amino acid alignments and phylogenetic data were performed using MEGA version 4 (Tamura et al., 2007). A phylogenetic tree was constructed using the neighbor-joining method. Bootstrap values were statistically calculated with the default sets of the MEGA program at 1000 replicates and seed = 44,251. The accession numbers of the L/ODC amino acid sequences used for comparison are listed in Supplemental Table 2 online.

**Gene Expression Analysis**

**Expression Analysis of L/ODC in *L. angustifolius***

Total RNA was prepared from mature leaves, young leaves, cotyledons, hypocotyls, and roots in both bitter and sweet forms of *L. angustifolius*. The cDNA encoding La-ODC was amplified by PCR using the RNeasy plant mini kit (Qiagen). To avoid the contamination of genomic DNA, DNase treatment of total RNA solution was performed. Each total RNA was reverse transcribed as described before. PCR was performed using primers LaDC-full-F and LaDC-full-R (see Supplemental Table 1 online). The constructs were then introduced into *Escherichia coli* BL21 competent cells. The *E. coli* cells harboring designated gene constructs were precultured in Luria-Bertani broth containing 100 μg/mL ampicillin at 37°C overnight. This culture was diluted 1/100 into new Luria-Bertani medium (containing 100 μg/mL ampicillin) and cultured at 37°C until the OD600 reached 0.4 to 0.6. Then, isopropyl-β-D-1-thiogalactopyranoside was added to a final concentration 0.1 mM and cells were cultured at 20°C for 6 h. The cells were collected and suspended in 1 × PBS (140 mM NaCl, 2.7 mM KCl, 10 mM Na2HPO4, and 1.8 mM KH2PO4, pH 7.3). Phenylmethylsulfonyl fluoride and pepstatin A were added to the cell suspension, which was then disrupted by sonication. The disrupted proteins were centrifuged and the supernatants were used as a crude protein.

**Gene Expression Analysis of La-L/ODC-Overexpressing Lines**

Total RNA were extracted (RNeasy kit; Qiagen) and reverse transcribed to cDNA as described previously from shoots of each transgenic line. PCR was conducted with KOD polymerase (Toyobo) using primers listed in Supplemental Table 1 online. The ORFs of La-L/ODC were amplified by PCR using Ex Taq DNA polymerase (TakaRa) with specific primers LaDC-full-F and LaDC-full-R (see Supplemental Table 1 online). PCR was performed with an initial denaturation at 95°C for 5 min, then 25 cycles each at 95°C for 30 s, 55°C for 30 s, and 72°C for 1.5 min. PCR products were separated by electrophoresis using 1.5% gel at 100 V, and gel was further stained by SYBR Green I nucleic acid gel stain (Invitrogen).

**Genomic PCR**

Genomic PCR of La-L/ODC was performed using Ex Taq DNA polymerase (TakaRa) with specific primers LaDC-full-F and LaDC-full-R (see Supplemental Table 1 online). PCR was performed with an initial denaturation at 95°C for 5 min, then 25 cycles each at 95°C for 30 s, 55°C for 30 s, and 72°C for 1.5 min.

**DNA Gel Blot Analysis**

Genomic DNA was extracted by the standard cetyltrimethyl ammonium bromide method (Sambrook and David, 2001). Aliquots of 15 μg of genomic DNA were digested with EcoRI, EcoRV, and HindIII and separated under denaturing conditions on a 0.8% agarose gel. After transfer to a Hybond N+ membrane (Amersham), hybridization was performed with the 32P-labeled probe prepared from the ORF of La-L/ODC cDNA using the Random Primer DNA labeling kit, version 2.0 (TakaRa). Hybridization signals were detected with a Storm 860 image analyzer (Amersham).

**Heterologous Expression of Recombinant L/ODCs**

The ORFs of La-L/ODC, Sf-L/ODC, and Ek-L/ODC were amplified by PCR using KOD polymerase (Toyobo) with gene-specific primers overhung with a restriction site (see Supplemental Table 1 online). The La-L/ODC mutants were prepared by PCR-based mutagenesis (Hipuchi et al., 1998). PCR was conducted with KOD polymerase (Toyobo) using primers listed in Supplemental Table 1 online. The amplified fragments were inserted in frame into the same restriction site of the expression vector pGEX-6P-2 (Amersham) for Sf-L/ODC, Ek-L/ODC, and La-L/ODC mutants, and pGEX-6P-3 (Amersham) for La-L/ODC, which gives a recombinant gene product with an N-terminal GST protein tag. Complete constructs were sequenced to confirm correct orientation.

The constructs were then introduced into *Escherichia coli* BL21 competent cells. The *E. coli* cells harboring designated gene constructs were precultured in Luria-Bertani broth containing 100 μg/mL ampicillin at 37°C overnight. This culture was diluted 1/100 into new Luria-Bertani medium (containing 100 μg/mL ampicillin) and cultured at 37°C until the OD600 reached 0.4 to 0.6. Then, isopropyl-β-D-1-thiogalactopyranoside was added to a final concentration 0.1 mM and cells were cultured at 20°C for 6 h. The cells were collected and suspended in 1 × PBS (140 mM NaCl, 2.7 mM KCl, 10 mM Na2HPO4, and 1.8 mM KH2PO4, pH 7.3). Phenylmethylsulfonyl fluoride and pepstatin A were added to the cell suspension, which was then disrupted by sonication. The disrupted proteins were centrifuged and the supernatants were used as a crude protein.
Plasmid Construction and Plant Transformation

The full-length ORF of La-L/ODC was transferred to the binary expression vector pGWB2 (Nakagawa et al., 2007) via Gateway technology (Invitrogen) to provide pGWB2-La-L/ODC (3SSpro:La-L/ODC). Subsequently, an electroporation method was used to introduce pGWB2-La-L/ODC into Agrobacterium tumefaciens strain LBA4404 for the generation of transgenic tobacco BY-2 suspension cells (Nicotiana tabacum cv Bright Yellow-2) and Arabidopsis plants, and into Agrobacterium rhizogenes strain 15834 for the generation of tobacco hairy root.

Tobacco BY-2 cells suspensions were maintained as described (Nakagawa et al., 2007). Transgenic tobacco BY-2 cells harboring pGWB2-La-L/ODC were generated by A. tumefaciens–mediated transformation according to the protocol (Häkkinen et al., 2007). Transformed colonies were picked and transferred to fresh plates, and their transgenic nature was confirmed by PCR. The transformed calli were subsequently suspended in liquid medium containing 50 ppm of kanamycin and hygromycin to keep selection pressure. Multiple independent transgenic lines were selected for metabolite analysis. These cells were elicited by methyl jasmonate for alkaloid production as described (Häkkinen et al., 2007).

The transgenic tobacco (N. tabacum cv Petit Havana line SR1) hairy roots were generated as described (Shoji and Hashimoto, 2008). The transformed tobacco hairy roots were subcultured in Gamborg B5 medium with 2% Suc every 2 weeks. For metabolite analysis, hairy root were inoculated in 20 mL medium and cultivate in rotary shaker (55 rpm, 25°C) in liquid Gamborg B5 medium with 2% Suc for 2 weeks. Transgenic Arabidopsis harboring pGWB2-La-L/ODC were generated via A. tumefaciens–mediated transformation by floral dip method (Clough and Bent, 1998). The seeds were collected from the dipped plants and selected in Murashige and Skoog medium (Murashige and Skoog, 1962) containing 50 ppm of kanamycin and hygromycin. T2 seeds were grown on Murashige and Skoog medium containing 1% (w/v) Suc, 50 ppm of kanamycin, and hygromycin. Plates were incubated at 22°C in a growth chamber with 16/8-h light/dark cycle. After 14 d, all shoots from each transgenic line and wild-type line were pooled separately, frozen in liquid nitrogen, and stored at −70°C to perform analysis of gene expression and metabolites.

Measurement of Alkaloids and Amines

Alkaloids of the transformed tobacco BY-2 cells were extracted as described (Häkkinen et al., 2007). For transgenic tobacco hairy roots, fresh samples were homogenized and 25 mg of ground sample was dissolved in extraction solution (methanol containing 250 ng/mL of 2,4′-dipyridyl as internal standard). After sonication in an ultrasonic bath for 2 h, the sample was centrifuged (8000 rpm, 5 min). Supernatant was collected and filtered through a 0.22-μm polyvinylidene difluoride filter. The samples were stored at −20°C until analysis. Quantitative analysis was performed based on the internal standard method by the HPLC/diode array detector (DAD) system (binary pump, L-7100; DAD, L-7455; autosampler, L-7200; Hitachi) using the same column and chromatographic conditions as described (Häkkinen et al., 2007). Alkaloids were also identified by liquid chromatography–photodiode array detector–electrospray ionization/MS consisting of an Agilent 6120 single quadrupole liquid chromatography/MS and an Agilent HPLC 1100 series (Agilent technologies) using the same column and chromatographic condition as HPLC/DAD. Nitrogen gas was used as a sheath gas for positive ion electrospray ionization/MS performed at a...
capillary temperature and voltage of 250°C and 4 kV, respectively. The tube lens offset was set at 2.0 V. Full-scan mass spectra were acquired from 100 to 700 mass-to-charge ratio at one scan s⁻¹. Identification of alkaloids was made by comparison with the authentic compounds, such as nicotine (Nacalai Tesque), (±) anabasine (Alfa Aesar), and (±) anatamine (Toronto Research Chemicals). Amines and amino acids of the transformed tobacco hairy root and Arabidopsis plants were extracted and analyzed by CE-MS as described (Ohkama-Ohtsu et al., 2008).

Accession Numbers
Sequence data from this article can be found in the DNA Data Bank of Japan (DDBJ) data library under the following accession numbers: La-L/ODC, L. angustifolius (DDBJ, AB566664); Si-L/ODC, S. flavescens (DDBJ, AB561138); Ek-L/ODC, E. koreensis (DDBJ, AB561139); Tc-L/ODC, T. chinensis (DDBJ, AB647178); and Ba-L/ODC, B. australis (DDBJ, AB647177).

Supplemental Data
The following materials are available in the online version of this article.

Supplemental Figure 1. Alignment of La-L/ODC, Si-L/ODC, and Eki-L/ODC Amino Sequences with Other Eukaryotic ODCs.

Supplemental Figure 2. SDS-PAGE of the Recombinant La-L/ODC, Si-L/ODC, Eki-L/ODC, and La-L/ODC Mutant Proteins in E. coli.

Supplemental Figure 3. Inhibition Study of La-L/ODC by L-Orn and α-DFMO.

Supplemental Figure 4. Comparative Protein Modeling of La-L/ODC and La-L/ODC Mutants.

Supplemental Figure 5. The Contents of Orn, Lys, Putrescine, and Cadaverine in Transgenic Tobacco Hairy Root Expressing GUS or La-L/ODC.

Supplemental Figure 6. Relative Expression between La-L/ODC Expression and Polyamines in Wild-Type, GUS-, and La-L/ODC-overexpressing Lines in Arabidopsis thaliana.

Supplemental Figure 7. Metabolite Profiles of La-L/ODC–Overexpressing Arabidopsis by CE-MS.

Supplemental Table 1. List of Primers Used in This Study.

Supplemental Table 2. Accession Numbers of Genes Used for Phylogenetic Analysis.

Supplemental Data Set 1. Text File of Alignment Used to Generate Phylogenetic Tree in Figure 2.

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AUTHOR CONTRIBUTIONS
K.S. and M.Y. designed the research. S.B., K.K., E.I., K.T., and A.O. performed the experiments. S.B., A.O., and M.Y. analyzed the data. S.B. and K.S. wrote the article.

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Lysine Decarboxylase Catalyzes the First Step of Quinolizidine Alkaloid Biosynthesis and Coevolved with Alkaloid Production in Leguminosae

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