Quinolizidine alkaloid biosynthesis: recent advances and future prospects

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INTRODUCTION

Plant secondary metabolites play multiple roles in the interaction between plants and their environment (Dixon, 2001). Almost 100,000 secondary metabolites have been discovered from plant species (Verpoorte, 1998; Hostettmann and Terreaux, 2000; Afendi, 2004). Several alkaloids have beneficial properties for humans and have been used in medicine. However, the molecular mechanisms underlying the biosynthesis of these alkaloids are not well understood. In the present article, we discuss recent advances in our understanding of Lys-derived alkaloids, especially the biochemistry, molecular biology, and biotechnology of quinolizidine alkaloid (QA) biosynthesis. We have also highlighted Lys decarboxylase (LDC), the enzyme that catalyzes the first committed step of QA biosynthesis and answers a longstanding question about the molecular entity of LDC activity in plants. Further prospects using current advanced technologies, such as next-generation sequencing, in medicinal plants have also been discussed.

Keywords: Lopin, lycopodium, lysine decarboxylase, lysine-derived alkaloids, N-tigloyltransferase, quinolizidine alkaloids

Lys-derived alkaloids, including piperidine, quinolizidine, indolizidine, and lycopodium alkaloids, are widely distributed throughout the plant kingdom. Several of these alkaloids have beneficial properties for humans and have been used in medicine. However, the molecular mechanisms underlying the biosynthesis of these alkaloids are not well understood. In the present article, we discuss recent advances in our understanding of Lys-derived alkaloids, especially the biochemistry, molecular biology, and biotechnology of quinolizidine alkaloid (QA) biosynthesis. We have also highlighted Lys decarboxylase (LDC), the enzyme that catalyzes the first committed step of QA biosynthesis and answers a longstanding question about the molecular entity of LDC activity in plants. Further prospects using current advanced technologies, such as next-generation sequencing, in medicinal plants have also been discussed.
FIGURE 1 Biosynthesis of universal intermediates, skeleton structures, and some Lys-derived alkaloids found in plants. Piperine, (−)-pelletierine, and (−)-lobeline are piperidine alkaloids. (−)-Lupinine and (−)-sparteine are quinolizidine alkaloids. (−)-Swainsonine is an indolizine alkaloid. Lycopodine and (−)-huperzine A are lycopodium alkaloids. LDC, Lys decarboxylase; CuAO, copper amine oxidase.

Spenser, 1988; Saito and Murakoshi, 1995; Ma and Gang, 2004). In contrast, indolizidine alkaloids are formed from L-Lys via L-pipecolic acid which is formed via the aldehyde and Schiff base, with retention of the nitrogen of the α-amino acid group (Figure 1; Harris et al., 1988a,b; Wickwire et al., 1990; Dewick, 2001). Δ¹-Piperideine and L-pipecolic acid are universal intermediates that can be modified by condensation, hydrolysis, or methylation to produce diverse Lys-derived alkaloids (Figure 1; Evans, 2009).

The genes and enzymes involved in the biosynthesis of Lys-derived alkaloids are not well documented. The best-studied pathway at the genetic level is the one leading to the formation of QAs, which will be discussed in the following section. One other study addresses partially purified piperonyl-CoA: piperidine N-piperoyltransferase (EC 2.3.1.145) from P. nigrum which catalyzes the synthesis of piperine in the presence of piperoyl-CoA and piperidine (Geisler and Gross, 1990).

QUINOLIZIDINE ALKALOIDS
Quinolizidine alkaloids are a group of alkaloids possessing a quinolizidine ring or a piperidine ring (Saito and Murakoshi, 1995). QAs occur mainly in the family Leguminosae, especially in the genera Lupinus, Baptisia, Thermopsis, Genista, Cytisus, and Sophora (Ohmiya et al., 1995). More than 500 Lupinus species have been described and 200–300 species are distributed in North and South America (Wink et al., 1995). QAs are important as potential sources of medicine. They have a broad range of pharmacological properties, including cytotoxic, oxytocic, antipyretic, antibacterial, antiviral, and hypoglycemic activities, as determined by in vivo pharmacological screening (Saito and Murakoshi, 1995). Some QA-containing plants, for example Sophora flavescens, have been used as sources of crude drugs in Chinese–Japanese medicine (KAMPO; Tang and Eisenbrand, 1992).

Quinolizidine alkaloids are synthesized through the cyclization of the cadaverine unit (Golebiewski and Spenser, 1988), which is produced through the action of LDC (Figure 2; Saito and Murakoshi, 1995). Various QA skeletons are further modified by tailoring reactions (e.g., dehydrogenation, oxygenation, or esterification) to yield several hundred structurally related alkaloids (Ohmiya et al., 1995). Lupinus accumulates two types of QA esters, the derivatives of lupinine and lupanine. QA
FIGURE 2 | Biosynthetic pathway of quinolizidine alkaloids. Quinolizidine alkaloids are synthesized by the decarboxylation of Lys, yielding cadaverine, which is further modified by various reactions as shown. The metabolites derived from the pathway are shown. Enzymes involved in the synthesis of these alkaloids are indicated: LDC, Lys decarboxylase; HMT/HLT, tigloyl-CoA: (-)13α-hydroxymultiflorine/(+)-13α-hydroxylupanine O-tigloyltransferase; ECT/EFT-LCT/LFT, p-coumaroyl-CoA/feruloyl-CoA: (+)-epilupinine/(-)-lupinine O-coumaroyl/feruloyltransferase.

esters are assumed to be end products of biosynthesis and storage forms (Ohmiya et al., 1995). Two acyltransferases (ATs) involved in QA ester biosynthesis have been identified and characterized: (-)-13α-hydroxymultiflorine/(+)-13α-hydroxylupanine O-tigloyltransferase (HMT/HLT; EC 2.3.1.93), which catalyzes the transfer of the tigloyl group from tigloyl-CoA to (-)-13α-hydroxymultiflorine or (+)-13α-hydroxylupanine, and p-coumaroyl-CoA/feruloyl-CoA: (-)-epilupinine/(+)-lupinine O-coumaroyl/feruloyltransferase (ICT/EFT-LCT/LFT), which catalyzes the transfer of p-coumaroyl-CoA or feruloyl-CoA to the hydroxyl moiety of (+)-epilupinine or (-)-lupinine (Figure 2; Saito et al., 1992, 1993; Suzuki et al., 1994). However, only HMT/HLT has been cloned and characterized at the molecular level (Okada et al., 2005). HMT/HLT belongs to the BAHD superfamily, a plant acyl-CoA-dependent AT gene family that transfers acyl groups from their CoA esters to various plant metabolites (D’Antu, 2006).

An investigation of an alkaloid-rich “bitter” cultivar and an alkaloid-poor “sweet” cultivar of L. angustifolius using random amplified polymorphic DNA (RAPD) showed that total DNA polymorphisms did not differ substantially between the two cultivars (Hirai et al., 2000). A cDNA-amplified fragment length polymorphism (cDNA-AFLP) was then used to isolate the genes specifically expressed in the bitter plant, however, no bitter-specific gene was isolated (Hirai et al., 2000). The PCR-select subtraction technique was then used to profile differentially expressed genes in the two cultivars. Among 71 bitter-specific fragments, three genes exhibited homologies to Orn decarboxylase (ODC, EC 4.1.1.17), CuAO, and AT, representing potential candidates for structural genes encoding QA biosynthetic enzymes (Bunsupa et al., 2011). Further study showed that the AT might be involved either in the formation of QA esters or N-acylated polyamine conjugates (Bunsupa et al., 2011) and ODC was identified as Lys/Orn decarboxylase (LDC) (see below) (Bunsupa et al., 2011). In addition, partial sequences encoding candidate transcription factors that might regulate the expression of genes in QA biosynthesis were obtained; these included DNA binding protein EREBP3, leucine zipper protein, and coronatine-insensitive 1, which provide clues about the regulation of the QA biosynthetic pathway (Bunsupa et al., 2011). Genomic PCR of L. angustifolius AT and LDC indicated the presence of at least one copy of both genes in the genomes of both the bitter and sweet cultivars. However, transcripts of both genes were expressed only in the bitter cultivar. This finding is interesting in terms of the regulation of the entire QA biosynthetic pathway (Bunsupa et al., 2011, 2012).

A NEW LYS DECARBOXYLASE FROM QUINOLIZIDINE ALKALOIDS PRODUCING PLANTS

Lys decarboxylase and ODC are key enzymes involved in the formation of cadaverine and putrescine by the decarboxylation of Lys and Orn, respectively. Cadaverine, putrescine, and other aliphatic polyamines, such as spermidine and spermine, are involved in a
number of growth and developmental processes (Bagini and Tassoni, 2003). Unlike plant ODCs, plant LDCs have not been well studied at the molecular level. There are several studies on crude protein extracts or partially purified protein extracts from Glycine max (Kim et al., 1996; Ohe et al., 2009) and L. polyphyllus (Hartmann et al., 1980). There was speculation that a member of group IV of the PLP-dependent amino acid decarboxylases might be responsible for the production of cadaverine from Lys (Sandri-Messer et al., 1994) and that ODC may also have LDC activity in plants that make cadaverine-derived alkaloids (Sindelar et al., 2004).

Recently, we isolated a cDNA of L/ODC from L. angustifolius (La-L/ODC) using differential transcript screening in QA-producing and non-producing cultivars (Bunsupa et al., 2011, 2012). We also obtained L/ODC cDNAs from S. flavescens and Eichinopsphora koreensis, which also produce QAs. Recombinant L/ODCs from QA-producing plants almost equally catalyzed the decarboxylation of L-Lys and L-Orn unlike the known ODCs that only accept Orn (Breker and Fischer, 1983; Bunsupa et al., 2012). In vivo experiments of La-L/ODC expression in Arabidopsis thaliana and Nicotiana tabacum provided evidence that this enzyme is involved in the first step of QA biosynthesis (Bunsupa et al., 2012). This is the first plant LDC identified at the molecular level by cDNA cloning and the second QA biosynthesis enzyme for which the cDNA has been identified.

Identification of La-L/ODC answers a longstanding question about the molecular entity of LDC activity in plants. We found LDCs in plant species that produce QAs. A point mutation from His-344 to Phe-344 (La-L/ODC numbering) is key for La-L/ODC to exhibit substrate promiscuity for Lys and Orn. This amino acid substitution has presumably occurred via adaptive evolution of LDC within the QAs producing plants to expand substrate accessibility to Lys (Bunsupa et al., 2012). 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 may be found not only in QA-producing Leguminosae but also in plants that produce cadaverine-derived alkaloids. This finding is a breakthrough in the understanding of both QA biosynthesis and LDC occurrence and function in plants.

A major safety issue of lupine-based foods is the presence of QAs, which are toxic to human and mammals. Genetic mapping of L. angustifolius and L. albus has been completed, which enables the mapping of the genes controlling key domestication traits, including alkaloid production, water permeability, flowering time, pod fertility, increased protein content, and related traits (Molvig et al., 1997). However, transformation systems for Lupinus sp. have been developed; however, only a few have been successful. For example, shoot regeneration using Agrobacterium-mediated gene transfer to proorganzed meristematic tissue combined with auxillary regeneration has been successful in L. angustifolius (Pigeaire et al., 1997). L. mutabilis (Li et al., 2000), and L. mutabilis (Babaoglu et al., 2000). However, transformation efficiency is poor and transformation in some Lupinus species and even in different cultivars of the same species is difficult. In L. mutabilis, the transformed roots synthesize isoflavones, but not QAs (Babaoglu et al., 2004). Genetic engineering of L. angustifolius by over expressed sun flower seed albumin gene has succeeded to enhance methionine levels and increase nutritive value of seeds of transgenic Lupinus plants (Molvig et al., 1997).

**ENGINEERING IN QA-PRODUCING PLANTS**

The concentration of QAs in cell suspension cultures of Lupinus and related plants is approximately 100–1000 times lower than that of concentration in leaves of differentiated plants (Hartmann et al., 1980; Hernandez et al., 2011). In cell suspension cultures, the storage tissue is probably repressed; thus, the QAs cannot be stored in large quantities and are degraded rapidly either inside the cells or in the cell culture medium (Wink, 1987). In callus, the concentration of QAs is correlated to the amount of chlorophyll in the cells (Saito et al., 1989a,b).

Lupinus spp. are the major grain legumes in Australia and have been used as feed supplements. In the past decade, they have gained attention as ingredients for human food because of their high percentage of protein (34–43% of dry matter) and acceptable content of essential amino acids (Sujak et al., 2006). To genetically engineer plants successfully, a routine transformation system is needed. Thus, it is necessary to develop a transformation system for Lupinus lines to confer desired traits, such as herbicide resistance, stress tolerance, resistance to certain pathogens, increased pod and seed set, and improved seed quality (Pigeaire et al., 1997). Many transformation systems for Lupinus sp. have been developed; however, only a few have been successful. For example, shoot regeneration using Agrobacterium-mediated gene transfer to proorganzed meristematic tissue combined with auxillary regeneration has been successful in L. angustifolius (Pigeaire et al., 1997), L. hirsutum (Li et al., 2000), and L. mutabilis (Babaoglu et al., 2000). However, transformation efficiency is poor and transformation in some Lupinus species and even in different cultivars of the same species is difficult. In L. mutabilis, the transformed roots synthesize isoflavones, but not QAs (Babaoglu et al., 2004). Genetic engineering of L. angustifolius by over expressed sun flower seed albumin gene has succeeded to enhance methionine levels and increase nutritive value of seeds of transgenic Lupinus plants (Molvig et al., 1997).

**FUTURE PROSPECTS**

In the past few years, progress has been made in the elucidation of alkaloid biosynthetic pathways and their regulation. However, only two genes, HMT/HET and L/ODC, have been isolated and characterized so far. Rapid progress in the development of integrative approaches, such as genomics, transcriptomics, proteomics, and metabolomics, has provided essential information for the understanding of many complex biological processes at different levels (Fukushima et al., 2009; Yonekura-Sakakibara and Saito, 2009).
Comparative and integrative analyses of such "omics" data will facilitate the determination of gene function and pinpoint rate limiting steps and factors in the biosynthetic pathway of secondary metabolites. Transcriptomic and metabolomic approaches, particularly transcriptome coexpression network analysis, have been used to identify genes involved in specific pathways not only in Arabidopsis but also in crops and medicinal plants. Such strategies have proven to be useful in fundamental plant biology and applied biotechnology (Ziegler et al., 2006; Hirai et al., 2007; Saito et al., 2008; Seki et al., 2008; Yonekura-Sakuma et al., 2008, 2012; Luscombe et al., 2009; Okazaki et al., 2009; Saito and Matsuda, 2010; Okazaki and Saito, 2012).

Advances in sequencing technology, next-generation sequencing, and computer data processing (Ehrhardt and Frommer, 2012) are opening a new era in plant science. It is estimated that the cost of DNA sequencing drops at a faster rate than the cost of computer data processing (Ehrhardt and Frommer, 2012). Therefore, the development of sequencing techniques with high-throughput will likely be applied to alkaloid-producing plants and provide the basis for many sequence-based approaches that are now limited to model plants such as Arabidopsis and rice (Ziegler and Facchini, 2008).

Today, many researchers are making progress in plant genome and transcriptome sequencing. For instance, the 1KP project aims to generate large-scale gene sequence information for 1000 different plants, including a wide variety of species from algae to land and aquatic plants, to understand the vast biodiversity that has barely been touched by genomics to date. In addition, the medicinal plant consortium project focuses on providing transcriptomic and metabolic resources for 14 key medicinal plants, for instance Atropa belladonna and Capsicum annuum, to the worldwide research community for the advancement of drug production and development.

Recently, we started the deep transcriptome sequencing of Japanese medicinal plants and both cultivars of L. angustifolius to obtain more information about the enzymes and transcription factors involved in the production of plant secondary metabolites. Such publicly accessible databases will provide novel information for the research community and ultimately accelerate the progress of the study of plant science.

The predicted function or in vitro characterization of alkaloid biosynthetic enzymes and other components must be confirmed in vivo (Ziegler and Facchini, 2008). However, there is no reliable and highly efficient transformation protocol for Lupinus and related plants. Virus induced gene silencing (VIGS), a technology that exploits an RNA-mediated antiviral defense mechanism, is a fast method for the generation of transient transformed plants (Lu et al., 2003). VIGS has been used widely in plants for analysis of gene function and has been adapted for high-throughput functional genomics (Burch-Smith et al., 2004). Thus, VIGS might be a powerful tool for assessing and characterizing the function of candidate genes in the QA biosynthetic pathway. Further investigations on inter- and intracellular transport of pathway intermediates and products and on cellular and subcellular localization of enzymes are also important because they are crucial factors in controlling the accumulation of target metabolites in plants (Facchini et al., 2014).

CONCLUSION
Knowledge of the biosynthetic pathway of Lys-derived alkaloids is limited. The best-studied pathway at the genetic level is QA biosynthesis; however, only two genes have been identified. Application of current technologies, such as next-generation sequencing, high-end proteomics, and metabolomics approaches, is needed to study secondary metabolic pathways for a better understanding of metabolic networks. Such information will aid in the development of successful strategies in genetic engineering of secondary metabolites in plants.

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