Elucidation of the core betalain biosynthesis pathway in *Amaranthus tricolor*

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*Amaranthus tricolor* L., a vegetable *Amaranthus* species, is an economically important crop containing large amounts of betalains. Betalains are natural antioxidants and can be classified into betacyanins and betaxanthins, with red and yellow colors, respectively. *A. tricolor* cultivars with varying betalain contents, leading to striking red to green coloration, have been commercially produced. However, the molecular differences underlying betalain biosynthesis in various cultivars of *A. tricolor* remain largely unknown. In this study, *A. tricolor* cultivars with different colors were chosen for comparative transcriptome analysis. The elevated expression of *AmCYP76AD1* in a red-leaf cultivar of *A. tricolor* was proposed to play a key role in producing red betalain pigments. The functions of *AmCYP76AD1*, *AmDODAα1*, *AmDODAα2*, and *AmcDOPA5GT* were also characterized through the heterologous engineering of betalain pigments in *Nicotiana benthamiana*. Moreover, high and low L-DOPA 4,5-dioxygenase activities of *AmDODAα1* and *AmDODAα2*, respectively, were confirmed through in vitro enzymatic assays. Thus, comparative transcriptome analysis combined with functional and enzymatic studies allowed the construction of a core betalain biosynthesis pathway of *A. tricolor*. These results not only provide novel insights into betalain biosynthesis and evolution in *A. tricolor* but also provide a basal framework for examining genes related to betalain biosynthesis among different species of *Amaranthaceae*.

Betalains are classified into betacyanins and betaxanthins, which provide red-violet and yellow coloration, respectively¹. Similar to anthocyanins, betalains exhibit antioxidant activity in the form of free-radical scavenging and accumulate in response to different stresses, such as UV-B radiation, high-intensity light, salinity, heat, and drought²-⁵. In addition to their potential roles in protecting plants against abiotic stresses, betalains play a role in defense against pathogenic fungi⁶. Moreover, as water-soluble natural pigments, betalains are widely used as food additives because of their health-promoting properties and color stability over a wide range of pH values⁷,⁸.

Betalains occur only in Caryophyllales and have never been detected in anthocyanin-producing plants⁹,¹⁰. Although the molecular basis for the mutual exclusion of betalains and anthocyanins is still unclear, breakthroughs in the identification of genes involved in betalain biosynthesis have shed light on the evolution of betalain pigmentation in Caryophyllales⁹-¹². Unlike anthocyanins, which are derived from L-phenylalanine, betalains are synthesized from L-tyrosine¹³,¹⁴. Initially, L-tyrosine is hydroxylated to produce L-DOPA by tyrosinases encoded by *CYP76AD1* and its orthologs¹⁵,¹⁶. L-DOPA can be converted into betalamic acid by L-DOPA 4,5-dioxygenase encoded by *DODA*, and betalamic acid can then spontaneously condense with amino acids to form betaxanthins¹,¹²,¹⁷. Alternatively, L-DOPA can be converted into *cyclo*-DOPA through the oxidase activity of *CYP76AD1*, and *cyclo*-DOPA can then spontaneously condense with betalamic acid to form betanin⁹,¹⁸. Betanidin is further glycosylated by betanidin 5-O-glucosyl-transferase encoded by *B5GT* to form betanin, the most common betacyanin. Glycosylation can also occur on *cyclo*-DOPA, catalyzed by *cyclo*-DOPA 5-O-glucosyltransferase encoded by *cDOPA5GT*, to produce *cyclo*-DOPA-glucoside, which then spontaneously condenses with betalamic acid to form betanin¹⁰,¹⁹.

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Phylogenetic analyses revealed that the CYP76AD and DODA genes, encoding key enzymes in the core biosynthetic pathway of betalains, are highly duplicated in Caryophyllales. The CYP76AD gene lineage has undergone at least three duplication events, giving rise to three clades: CYP76Ada, CYP76Adβ, and CYP76Adγ. The CYP76Adγ clade includes the CYP76Ad1 and CYP76Ad3 genes, whose products possess both the tyrosine hydroxylase and L-DOPA 4,5-dioxygenase activities required for L-DOPA and cyclo-DOPA formation, respectively. The CYP76Adγ clade includes the CYP76Ad5, CYP76Ad6, and CYP76Ad15 genes, which possess only the tyrosine hydroxylase activity required for L-DOPA formation. However, the functions of the genes in the CYP76Adγ clade in betalain biosynthesis have not been determined. Duplication events in the DODA gene lineage also gave rise to two major clades: DODAA and DODAB. The function of DODAB is unknown, but the evolution of L-DOPA 4,5-dioxygenase activity in betalain-producing plants was proposed to be led by DODAA. Nevertheless, only one paralogous gene from the DODAA clade shows high L-DOPA 4,5-dioxygenase activity in each species, and others exhibit barely detectable activity.

An increased understanding of the betalain biosynthesis pathway has facilitated the metabolic engineering of betalains, providing new sources for basic research studies and commercial applications. For example, the fluorescent betalain pigments produced by the expression of MjDODAA1 in yeast were used as chemical biosensors to reveal the tyrosine hydroxylase activity of BvCYP76AD1. The production of semisynthetic betaxanthins by spontaneous condensation between fluorescent betalamic acid and the amino groups of proteins provides an alternative method for labeling proteins for multiple biological applications. Fine-turning the content ratio of betacyanins and betaxanthins produced by the expression of BvCYP76AD1, BvDODAα1, and BvDODAβ1 in tobacco enhances the resistance of transgenic plants to Botrytis cinerea infection. In addition, betanin rice generated through the coexpression of BvCYP76AD1, BvDODAA1, and BvDODAα1 in rice endosperm shows higher antioxidant activity and can be provided as a functional food.

Amaranthus species are economically important crops containing large amounts of betalains. They can be classified into three categories: vegetable Amaranthus, grain Amaranthus, and weed Amaranthus species. Amaranthus tricolor L., a vegetable Amaranthus species, is widely distributed in warm and tropical regions and is cultivated as a leafy vegetable. As rich in natural antioxidants, A. tricolor is able to tolerate abiotic stresses and has been used as a traditional Chinese medicinal herb for the treatment of eruptive fever, pain, sore throat, dysentery, anemia, bronchitis, colic, etc. A. tricolor cultivars with various contents of betacyanins and betaxanthins, resulting in striking colors ranging from red to green, have been commercially produced. However, the molecular basis underlying betalain biosynthesis in A. tricolor remains largely unknown. Recently, virus-induced gene silencing was applied to elucidate the function of CYP76AD1 in producing betalain pigments in A. tricolor. In addition, a comparative analysis of a transcriptome database constructed from different leaf samples of A. tricolor cv. Dahong was performed to construct a putative metabolic pathway of betalains in A. tricolor. Candidate genes encoding enzymes catalyzing the formation of L-DOPA, cyclo-DOPA, betalamic acid, cyclo-DOPA-glucoside, and betanin were obtained from a transcriptome database and showed higher expression levels in red areas of A. tricolor leaves than in green areas. However, functional characterization and enzyme activity analyses are still needed to elucidate the roles of these candidate genes in the betalain biosynthesis pathway of A. tricolor.

In this study, A. tricolor cultivars with different colors were chosen for comparative transcriptome analysis. The key gene showing elevated expression in a red-leaf cultivar of A. tricolor was identified, and the results indicated that the dual activities of tyrosine hydroxylase and L-DOPA oxidase are important for producing red betalain pigments in A. tricolor. The core betalain biosynthesis pathway of A. tricolor was further constructed based on the functional characterization of betalain biosynthesis genes through the heterologous engineering of betalain pigments in Nicotiana benthamiana and in vitro enzymatic assays of L-DOPA 4,5-dioxygenase activities. These results provide novel insights into betalain biosynthesis and evolution in A. tricolor.

Results
AmCYP76AD1 is highly expressed in a red-leaf cultivar of A. tricolor. A. tricolor cultivars are important leafy vegetables that display leaf colors ranging from red to green depending on the betalain content. To elucidate the genetic factors that influence betalain pigment contents in red- and green-leaf cultivars of A. tri-
AmDODA exhibits a marginal level of L-DOPA 4,5-dioxygenase activity. Although candidate transcripts related to betalain biosynthesis were identified previously in *A. tricolor*, their functional and enzymatic activities have not yet been characterized. To functionally characterize the enzyme activities of AmCYP76AD1, AmDODA, and AmDOPAG5T in the core pathway of betalain biosynthesis (Fig. 1e), 35S promoter-driven cDNAs encoding C-terminal YFP- or FLAG (SF)-tagged AmCYP76AD1, AmDODA, and AmDOPAG5T were transiently coexpressed in *N. benthamiana* leaves by agroinfiltration. Upon expression, only a small amount of betalain pigment was produced in *N. benthamiana* leaves, which was barely detectable (Fig. 3a). In contrast, as a positive control, high production of betalain pigments with red-violet color was observed when the *Beta vulgaris* tyrosinase gene (*BvDODAa1*), and the *Mirabilis jalapa* cyclo-DOPA 5-O-glucosyltransferase gene (*MjcDOPA5GT*), were coexpressed in *N. benthamiana* leaves (Fig. 3a). To elucidate the *A. tricolor* genes responsible for the negligible activity of betalain synthesis in transient analysis, a series of coinfiltration assays were carried out by replacing the positive control genes individually with *AmCYP76AD1, AmDODA, and AmcDOPA5GT* in the core pathway of betalain biosynthesis (Fig. 1d). The highly differential expression pattern of *AmCYP76AD1* between AMR and AMG was also observed in 4-week-old *A. tricolor* (Supplementary Fig. S1c). Moreover, as a key element in the initiation of the betalain biosynthesis pathway, *AmCYP76AD1* displayed higher transcript levels in the upper leaves of AMR, which contained higher content of betalains than those in the lower leaves of AMR (Fig. 2a,b, Supplementary Fig. S2). Further phylogenetic reconstruction and LOGO analysis revealed that *AmCYP76AD1* belongs to the CYP76Adα clade (Fig. 2c,d), whose members possess both the tyrosine hydroxylase and L-DOPA oxidase activities required for L-DOPA and cyclo-DOPA formation, respectively (Fig. 1e). These results suggest that the elevated expression of *AmCYP76AD1* is necessary for betalain pigment accumulation, which leads to an obvious red-violet color in the leaves and stems of AMR, but not in those of AMG.
AmDODAα1, but not AmDODAα2, exhibits a high level of L-DOPA 4,5-dioxygenase activity. As a key step in betalain biosynthesis, L-DOPA 4,5-dioxygenase can convert L-DOPA into betalamic acid, the basic structural unit of all betalains1,32. To functionally characterize the L-DOPA 4,5-dioxygenase activity of AmDODAα1, AmDODAα1 was coexpressed with BvCYP76AD1 and MjcDOPA5GT by agroinfiltration. As a result, high production of betalain pigments and betanin was observed when comparable amounts of proteins were expressed in N. benthamiana leaves (Fig. 3b–d, Supplementary Fig. S3). These results indicate that AmDODAα1, but not AmDODAα2, exhibits a high level of L-DOPA 4,5-dioxygenase activity, similar to that of BvDODAα1.

To verify enzyme activity in vitro, AmDODAα1 and AmDODAα2 were expressed as SUMO-fused recombinant proteins in an Escherichia coli expression system (Fig. 5a). Enzymatic reactions were conducted following the method described by Sasaki et al32, in which crude extracts prepared from E. coli were used. After incubation for 5 min at 30 °C, a bright yellow color derived from betalamic acid was observed in the reaction mixture containing L-DOPA, ascorbic acid, and a crude extract prepared from E. coli harboring AmDODAα1 or BvDODAα1, but not AmDODAα2 (Fig. 5b). However, only a very weak yellow color was observed when the reaction mixture contained twofold crude extract prepared from E. coli harboring AmDODAα2 (Fig. 5b). As a control, a reaction mixture containing the crude extract was prepared from E. coli harboring only the vector, and no color was
observed (Fig. 5b). The reaction products were then subjected to LC–MS/MS analysis and revealed that the clear peak at a retention time of 7.5 min was betalamic acid (Fig. 5c). These results confirm that AmDODAα2 exhibits marginal levels of L-DOPA 4,5-dioxygenase activity.

Reconstruction of the core betalain biosynthesis pathway of *A. tricolor* in *N. benthamiana*. In this study, we also attempted to use TRV-based virus-induced gene silencing (VIGS) to examine the functional activities of genes involved in betalain biosynthesis in *A. tricolor*. However, the transient silencing of AmCYP76AD1 in *A. tricolor* was particularly challenging and failed in our hands. In addition, the attempted overexpression of AmCYP76AD1 to complement the betalain pigments in the leaves of AMG was unsuccessful using an agroinfiltration system. These differences might have resulted from the different varieties and low transformation efficiency of *A. tricolor*.

To reconstruct the core betalain biosynthesis pathway of *A. tricolor* in *N. benthamiana*, AmCYP76AD1, AmDODAα1, and AmcDOPA5GT were transiently overexpressed in *N. benthamiana* leaves by agroinfiltration for the heterologous engineering of betalain pigments. Similar to the vector-only control, the heterologous expression of single AmCYP76AD1, AmDODAα1, or AmcDOPA5GT was not sufficient to produce any betalain pigment in *N. benthamiana* (Fig. 6a). However, low production of betalain pigments was observed when AmCYP76AD1 and AmDODAα1 were coexpressed in *N. benthamiana* (Fig. 6a). In contrast, no betalain pigment was observed when AmCYP76AD1 and AmcDOPA5GT or AmDODAα1 and AmcDOPA5GT were coexpressed in *N. benthamiana* (Fig. 6a). Only the coexpression of AmCYP76AD1, AmDODAα1, and AmcDOPA5GT together was sufficient to produce high amounts of betalain pigments in *N. benthamiana*, which resulted in a strong red-violet color (Fig. 6a). The strong red-violet color was similar to that in the positive control in which BvCYP76AD1, BvDODAα1, and MjcDOPA5GT were coexpressed in *N. benthamiana* (Fig. 6a). As expected, the coexpression of AmCYP76AD1, AmDODAα2, and AmcDOPA5GT only produced marginal levels of betalain pigments, which were
barely detectable (Fig. 6a). Consistently, high production of betanin was observed only when AmCYP76AD1, AmDODAα1, and AmDOPA5GT were coexpressed in N. benthamiana leaves (Fig. 6b). Together with the comparable amount of proteins detected by western blotting (Fig. 6c, Supplementary Fig. S5), our results suggest that the enzyme activities of AmCYP76AD1, AmDODAα1, and AmDOPA5GT are sufficient to construct the core betalain biosynthesis pathway of A. tricolor.

Discussion
Molecular genetics have shed light on the betalain biosynthesis pathway and its evolutionary significance in Caryophyllales. Based on phylogenetic analysis, CYP76AD homologues can be classified into α, β, and γ clades. To date, only the functions of CYP76ADα and CYP76ADβ clade homologues, such as CYP76AD1 and CYP76AD6, have been reported. For example, the cosilencing of CYP76AD1 and CYP76AD6 represses the production of betacyanins and betaxanthins in B. vulgaris, causing a green leaf phenotype. In this study, a CYP76AD6-like (AmCYP76AD6) gene, belonging to the CYP76ADβ clade according to phylogenetic construction and LOGO analysis (Fig. 2c,d), was also identified in A. tricolor through transcriptome analysis (Supplementary Fig. S6). However, the expression of AmCYP76AD6 was extremely low and was difficult to detect in AMR and AMG. As a result, it is difficult to functionally connect AmCYP76AD6 with the production of betalains in A. tricolor. In addition, although PPO, a polyphenol oxidase gene, and CATPO, a catalase-phenol oxidase gene, were previously proposed to be involved in betalain biosynthesis via monophenolase activity, their transcripts did not show highly differential expression patterns between AMR and AMG (Fig. 1d, Supplementary Fig. S1c). As a result, we propose that the elevated expression of AmCYP76AD1 is necessary for the occurrence of a red-violet color phenotype in A. tricolor; in contrast, the loss of AmCYP76AD1 expression results in a green color phenotype in A. tricolor (Fig. 1a–d). The existence of the AmCYP76AD1 gene in AMG examined by PCR using genomic DNA as a template confirmed the loss of AmCYP76AD1 expression in AMG (Supplementary Fig. S7). Together with the functional characterization of the enzymatic activity of AmCYP76AD1 through the heterologous engineering of betalain pigments in N. benthamiana (Figs. 3b, 6a), we conclude that AmCYP76AD1, a CYP76ADα homologue required for the initiation of the betalain biosynthesis pathway, plays a key role in betalain pigment accumulation in A. tricolor. Accordingly, AmCYP76AD1 displayed higher transcript levels in the upper leaves of AMR, which contained higher content of betalains than those in the lower leaves of AMR (Fig. 2a,b, Supplementary Fig. S2).
biosynthesis in Caryophyllales \textsuperscript{16,20,30,36,37}. However, numerous duplication events have led to difficulty in elucidating the functional activities of key enzymes in betalain-pigmented species through annotation \textsuperscript{10}. For example, duplication events gave rise to two major clades of DODA homologues, DODA\textalpha{} and DODA\textbeta{}, but only one gene paralog in the DODA\textalpha{} clade of each species exhibits high levels of L-DOPA 4,5-dioxygenase activity \textsuperscript{12,21}. Thus, it is necessary to examine the possible involvement of annotated genes in betalain biosynthesis on the basis of experimental evidence. In this study, the AmDODA\alpha{}\textsubscript{1} and AmDODA\alpha{}\textsubscript{2} genes, which belong to the DODA\alpha{} clade according to phylogenetic construction and LOGO analysis (Fig. 4b,c), were identified in \textit{A. tricolor} through transcriptome analysis (Supplementary Fig. S4, Table S4). Based on the heterologous engineering of betalain pigments in \textit{N. benthamiana} and in vitro biochemical studies (Figs. 3b, 5b), we report that AmDODA\alpha{}\textsubscript{1} displayed a high level of L-DOPA 4,5-dioxygenase activity to produce betalamic acid, but such activity was barely detectable for AmDODA\alpha{}\textsubscript{2}. These results indicate that at least one duplication event has occurred in the DODA\alpha{} lineage of \textit{A. tricolor}, and the primary function of AmDODA\alpha{}\textsubscript{2} remains to be further studied.

Betalains are composed of betacyanins and betaxanthins. In contrast to betaxanthins, which are derived from betalamic acid via spontaneous condensation with amino acids or other amines, a large number of betacyanins are composed of betanidin conjugated with glycosyl moieties \textsuperscript{9,10}. We characterized the function of AmcDOP-A5GT, a cyclo-DOPA 5-O-glucosyltransferase gene, through the heterologous engineering of betalain pigments in \textit{N. benthamiana}. The coexpression of AmCYP76AD1, AmDODA\alpha{}\textsubscript{1}, and AmcDOPA5GT enabled the production of high levels of betalain pigments with a dark red color (Fig. 6a). In contrast, low production of betalain pigments was observed when AmCYP76AD1 and AmDODA\alpha{}\textsubscript{1} were coexpressed (Fig. 6a). Our results suggest the importance of AmcDOPA5GT in the glycosylation reaction during betalain biosynthesis in \textit{A. tricolor}. In fact, the metabolic pathway of betalain biosynthesis is very complex due to multiple glycosylation steps, and different betacyanins have been identified \textsuperscript{39,40}. For example, betanin, the most common betacyanin, is not only produced by cyclo-DOPA 5-O-glucosyltransferase but is also produced by betanidin 5-O-glucosyl-transferase through the glycosylation of betanin \textsuperscript{39,40}. In this study, AmB5GT, a betanidin 5-O-glucosyl-transferase gene, was also identified through comparative transcriptome analyses (Supplementary Table S4). Although AmcDOPA5GT showed higher expression levels than AmB5GT in both AMR and AMG (Supplementary Table S4), it remains to be determined which of the two glycosylation routes is more important for the formation of betanin in \textit{A. tricolor}.

Recently, betalain biosynthesis in different pitaya species, such as \textit{Hylocereus polyrhizus}, \textit{Hylocereus costaricensis}, \textit{Hylocereus undatus}, and \textit{Hylocereus megalanthus}, has been intensively studied through comparative transcriptome analysis \textsuperscript{36,37,41,42}. However, further studies remain to be conducted to provide experimental evidence and strengthen the understanding of the roles of candidate genes in betalain biosynthesis. Here, complementation assays conducted through the heterologous engineering of betalain pigments in nonbetalain-producing plants...
provided a solution for the easy and rapid comparison of the functional activities of genes involved in the core betalain biosynthesis pathway between betalain-pigmented species of Caryophyllales. Using the coexpression of BvCYP76AD1, BvDODAα1, and MjcDOPA5GT in N. benthamiana as a positive control, the functional activities of A. tricolor genes responsible for betalain synthesis could be compared through a series of complementation assays (Fig. 3b–d). We showed that comparable amounts of betalain pigments were observed when the functional activities of positive genes were individually replaced with AmCYP76AD1, AmDODAα1, and AmcDOPA5GT in transient coexpression assays (Fig. 3b–d). Our results indicate that AmCYP76AD1, AmDODAα1, and

Figure 6. Reconstruction of the core betalain biosynthesis pathway of A. tricolor in N. benthamiana by agroinfiltration for the heterologous engineering of betalain pigments. (a) N. benthamiana leaves coinfiltrated with Agrobacterium harboring plasmids for the expression of BvCYP76AD1-YFP (BvAD1), BvDODAα1-SFP (BvDAα1), MjcDOPA5GT-SFP (Mj5GT), AmCYP76AD1-YFP (AmAD1), AmDODAα1-SFP (AmDAα1), AmDODAα2-SFP (AmDAα2), and AmcDOPA5GT-SFP (Am5GT). Both the adaxial (left) and abaxial (right) sides of leaves are presented in each panel. Bottom right corner indicated the betalain pigments produced in N. benthamiana. (b) Extracted betalain pigments were examined for betanin content by LC–MS/MS analysis. Shown are XICs of masses corresponding to betanin (m/z = 551). Time, retention time (min). (c) Western blotting assays were conducted to examine the expression levels of YFP-tagged CYP76AD1 (upper panel), SFP-tagged DODA (middle panel), and SFP-tagged cDOPA5GT (middle panel) using antibodies against the YFP- or FLAG (SFP)-tag. As a loading control, the large subunit of Rubisco visualized with Coomassie brilliant blue staining is indicated by the arrowhead (lower panel).
AmCYP76AD1 exhibit high tyrosinase, L-DOPA 4,5-dioxygenase, and cyclo-DOPA 5-O-glucosyltransferase activities, respectively, which are similar to those in *B. vulgaris* and *M. jalapa*. Accordingly, in vitro biochemical studies demonstrated that AmDODAα1 displayed comparable L-DOPA 4,5-dioxygenase activity to BvDODAα1 in producing betalamic acid (Fig. 3b). These results provide novel insights into betalain biosynthesis and evolution in *A. tricolor*.

In conclusion, a comparative transcriptome analysis combined with functional and enzymatic studies were performed to reveal the core betalain biosynthesis pathway of *A. tricolor*. The heterologous engineering of betalain pigments through the coexpression of AmCYP76AD1, AmDODAα1, and AmCYP76AD5GT in *N. benthamiana* enabled the production of high amounts of betalain pigments with a red-violet color similar to those in the red-leaf cultivar of *A. tricolor*. Although the metabolic pathway of betalain biosynthesis is very complex, the core betalain biosynthesis pathway of *A. tricolor* constructed here not only provides a basal framework for examining genes related to betalain biosynthesis within the species of *Amaranthaceae* but also sheds light on the evolution of the betalain biosynthesis pathway in Caryophyllales.

**Methods**

**Plant materials and growth conditions.** *A. tricolor*, *B. vulgaris*, *M. jalapa*, and *N. benthamiana* plants were grown at 26 °C in a semicontrolled walk-in chamber under a 16:8-h light:dark photoperiod. Soil (Jiffy) mixed with vermiculite and pearlstone was used. Seeds of *A. tricolor* cv. Hung Hsien (red-leaf cultivar) and *A. tricolor* cv. Pai Hsien (green-leaf cultivar) were purchased from KNOWN-YOU SEED CO., LTD.

**Betalain pigment extraction and measurement.** For betalain pigment measurement, betalain contents were determined as described previously with some modification. Briefly, leaves of seedlings were collected and ground into powder in liquid nitrogen. Betalain pigments were extracted with extraction solution (methanol:chloroform:H₂O [1:2:1]). After centrifugation, the upper (hydrophilic) layer was collected to measure the absorbance at 538 nm and 476 nm for betacyanins and betaxanthins, respectively. The relative betalain content was calculated with the following equation: \((A_{538} + A_{476})/\text{gram}\).

**Plasmid construction.** All plasmid constructs were generated using standard restriction site reconstruction methods and confirmed by DNA sequencing. AmCYP76AD1, AmDODAα1, AmDODAα2, AmCYP76AD5GT, BvCYP76AD1, BvDODAα1, and MjcDOPA5GT were amplified from *A. tricolor*, *B. vulgaris*, or *M. jalapa* cDNA libraries using AccuPrime pfX DNA polymerase (Invitrogen). For the transient expression of C-terminal YFP-or FLAG (SFP)-tagged proteins in *N. benthamiana*, PCR products encoding AmCYP76AD1, AmDODAα1, AmDODAα2, AmCYP76AD5GT, BvCYP76AD1, BvDODAα1, and MjcDOPA5GT were subcloned into pBA-C-SFP or pBA-C-YFP vectors under the control of a Cauliflower mosaic virus (CaMV) 35S promoter. To produce N-terminal SUMO-tagged recombinant proteins, PCR products encoding AmDODAα1, AmDODAα2, and BvDODAα1 were subcloned into the pET-SUMO (Invitrogen) vector. For the VIGS assay, a cDNA fragment of AmCYP76AD1 was amplified and subcloned into the pTRV2 vector. The primer sequences used for plasmid construction are listed in Supplementary Table S5.

**Quantitative real-time polymerase chain reaction (qRT-PCR) and statistical analysis.** TRIzol™ (Invitrogen)-extracted total RNA was reverse transcribed using SuperScript III First-Strand Synthesis SuperMix (Invitrogen) according to the manufacturer’s instructions. Briefly, each sample was prepared from the leaves of three biologically distinct 3-week-old or 4-week-old *A. tricolor* plants. Then, cDNA was synthesized from 1 μg of total RNA using a mixture of random hexamers and oligo(dT)₂₀ under the following conditions: 25 °C for 10 min, followed by 50 °C for 40 min. The cDNA was employed as a template for qRT-PCR using the KAPA SYBR Fast qPCR Kit (Kapa Biosystems). Three technical replicates were performed on a CFX96™ Real-time System (Bio-Rad) under the following conditions: 95 °C for 3 min, followed by 40 cycles of 95 °C for 10 s and 55 °C for 30 s. The expression levels of selected genes were determined by normalization to the reference gene *Actin*. Statistically significant differences were determined using Student’s *t*-test in SPSS version 20.0. The primer sequences employed for qRT-PCR analyses are listed in Supplementary Table S1. PCR analyses using genomic DNA extracted from AMR and AMG as a template were performed to confirm the specificity of the primers (Supplementary Fig. S7).

**Transient coexpression assay and western blotting.** Plasmids for the transient expression of AmCYP76AD1-YFP, AmDODAα1-SFP, AmDODAα2-SFP, AmCYP76AD5GT-SFP, BvCYP76AD1-YFP, BvDODAα1-SFP, or MjcDOPA5GT-SFP were transformed into the *Agrobacterium tumefaciens* strain AGL1. C-terminal tagged proteins were coexpressed using a mixture of *A. tumefaciens* carrying the desired constructs in *N. benthamiana* leaves by agroinfiltration following the method described previously. After three days, the infiltrated leaves were photographed and ground into a powder in liquid nitrogen for total cell extract preparation. Briefly, 0.1 g of sample powder was added to 0.2 ml of 2.5 × SDS sample buffer (5 mM EDTA, 5% SDS, 0.3 M Tris–HCl, pH 6.8, 20% glycerol, 1% β-mercaptoethanol, and bromophenol blue), which was then heated at 95 °C for 10 min. After centrifugation at 13,000 × g for 10 min, the supernatant was obtained, and total proteins were separated by SDS-PAGE. Western blotting assays were performed to monitor protein levels using specific polyclonal and monoclonal antibodies against YFP- and FLAG-tag, respectively. Chemiluminescence signals generated by ECL reagents (PerkinElmer) were captured with an ImageQuant LAS 4000 mini imager (GE Healthcare). All experiments were repeated at least three times using biologically distinct samples prepared from two infiltrated leaves.
In vitro L-DOPA 4,5-dioxygenase activity assay and liquid chromatography-tandem mass spectrometry (LC–MS/MS) analysis. An in vitro L-DOPA 4,5-dioxygenase activity assay was performed according to the method described previously with some modifications. Briefly, plasmids for the expression of N-terminal SUMO-tagged AmDODAa1, AmDODAa2, and BvDODAa1 were transformed into Escherichia coli strain BL21 (DE3). The transformants were grown in 50 ml LB medium, and the recombinant proteins were induced with 0.2 mM IPTG at 22 °C for 16 h. Harvested cells were washed, resuspended, and disrupted by sonication in 50 mM sodium phosphate buffer (pH 7.0). The crude extract (supernatant) was used for the enzyme activity assay after centrifugation at 14,000 × g for 15 min. The amount of recombinant protein was quantified with Protein Assay Reagent (Bio-Rad) and via Coomassie blue staining SDS-PAGE with BSA as the standard. Basically, the reaction (100 μl) was performed with the crude extract containing 8 μg DODA protein, 27 mM ascorbic acid, and 6.75 mM L-DOPA at 30 °C for 5 min.

LC–MS/MS was performed using a Dionex UltiMate 3000 system (Thermo Fisher Scientific) linked with an amaZon speed-ion trap mass spectrometer (Bruker). Betalamic acid was detected on a Waters BEH shield RP18 column with two eluting solvent systems: (A) H2O with 0.1% formic acid, (B) 100% acetonitrile. The gradient elution program was set as follows: 0–3 min (100% A), 9 min (55% A and 45% B), 12–13 min (100% B). The flow rate was 0.3 ml min⁻¹, and the detector wavelength was 424 nm. The electrospray ionization mass parameters were set as follows: 4.5 kV capillary, 500 V end plate offset voltage, 40.0 psi nebulizer pressure, 8.0 l min⁻¹ dry gas, and 230 °C dry temperature. The measurement was operated in multiple reaction-monitoring (MRM) with the positive ion mode. The MRM was set 182 → 165 m/z to detect tyrosine, 198 → 181 m/z to detect L-DOPA, 212 → 166 m/z to detect betalamic acid, 389 → 345 m/z to detect betanidin, and 551 → 389 m/z to detect betanin.

Next-generation sequencing and MA plot. To perform next-generation sequencing, aerial tissues derived from three biologically distinct 3-week-old A. tricolor plants were collected. Total RNA was extracted using the RNaseasy Plant Mini Kit (Qiagen) according to the manufacturer’s instructions. RNA quality was examined via 1.2% (wt/vol) formaldehyde gel electrophoresis and with an Experion RNA analysis kit (Bio-Rad, Munich). Only high-quality RNA was used for next-generation sequencing performed on the Illumina HiSeq 4000 platform with 150 paired-end reads. For each dataset (AMR and AMG), 100 million reads were generated, and de novo assembly was performed with the Trinity tool. The assembled transcripts were annotated with BlastX in UniProt. Gene expression levels were normalized as FPKM values, and differentially expressed genes were identified according to an FDR < 0.05 and logFC > 2 or < −2 (Supplementary Tables S2, S3). An MA plot was generated based on the average concentration (logCPM) and fold-change (logFC) values to show the relative abundances of transcripts between AMR and AMG.

Phylogenetic tree reconstruction and LOGO analysis. Phylogenetic trees were reconstructed using MEGA-X software based on the protein sequence comparisons of CYP76AD and DODA homologues from different betalain-producing species. Multiple sequence alignments were performed using the MUSCLE program and were processed to generate a maximum likelihood phylogenetic tree via the Jones–Taylor–Thornton (JTT) model with bootstrapping to perform molecular evolutionary analysis. The numbers at the branch points are bootstrap values representing the percentages of replicate trees based on 1000 repeats. LOGO analyses were performed via WebLogo (http://weblogo.berkeley.edu/logo.cgi) based on selected conserved amino acids of CYP76AD and DODA homologues reported previously. The species, families, and accession numbers of CYP76AD and DODA homologues are available in Supplementary Table S6.

Data Availability
Sequencing data generated for this study are deposited at Short Read Archive with the accession code SRR15044103.

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**Author contributions**

Y.-C.C. carried out most of the experiments; Y.-C.C. and Y.-L.C. designed and carried out the enzymatic assays. N.-W.T. performed the LC–MS analysis. C.-M.T., Y.-H.C., and P.-C.L. performed RNA-seq and plasmid constructions. Y.-C.L., L.-C.H. and S.-Y.W. contributed to the interpretation of data. J.-Y.Y. conceived and wrote the manuscript.

**Competing interests**

The authors declare no competing interests.

**Additional information**

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