An acyltransferase gene that putatively functions in anthocyanin modification was horizontally transferred from Fabaceae into the genus Cuscuta

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\textbf{A B S T R A C T}

Horizontal gene transfer (HGT) refers to the flow of genetic materials to non-offspring, and occasionally HGT in plants can improve the adaptation of organisms in new niches due to expanded metabolic capability. Anthocyanins are an important group of water-soluble red, purple, or blue secondary metabolites, whose diversity results from modification after the main skeleton biosynthesis. Cuscuta is a stem holoparasitic genus, whose members form direct connection with hosts to withdraw water, nutrients, and macromolecules. Such intimate association is thought to increase the frequency of HGT. By transcriptome screening for foreign genes in Cuscuta australis, we discovered that one gene encoding a putative anthocyanin acyltransferase gene of the BAHD family, which is likely to be involved in anthocyanin modification, was acquired by C. australis from Fabaceae through HGT. The anthocyanin acyltransferase-like (AT-like) gene was confirmed to be present in the genome assembly of C. australis and the transcriptomes of Cuscuta pentagona. The higher transcriptional level in old stems is consistent with its putative function in secondary metabolism by stabilizing anthocyanin at neutral pH and thus HGT of this AT-like gene may have improved biotic and abiotic resistance of Cuscuta.

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1. Introduction

Plants synthesize numerous secondary metabolites that are beneficial for adaptation to biotic or abiotic stresses and thus contribute to the fitness of species (Pichersky and Lewinsohn, 2011). Flavonoids are an important class of plant secondary metabolites. In the model plant Arabidopsis thaliana, at least 54 flavonoid molecules (35 flavonols, 11 anthocyanins and 8 proanthocyanidins) have been identified (Saito et al., 2013). Genes that play a role in flavonoid modification, including glycosylation, methylation and acylation (Grotewold, 2006; Lepiniec et al., 2006), constitute large families and thus greatly increase the structure and genetic diversity of flavonoids in plants. Anthocyanins are mostly red, purple, or blue water-soluble pigments derived from a branch of the flavonoid pathway. As antioxidants and pigments, anthocyanins have applications in agriculture, food products, and human health. Biosynthesis of anthocyanins requires acylation which is carried out by a group of flavonoid-specific acyl CoA-dependent acyltransferases in the BAHD family, named according to the first letter of the family (BEAT, AHCT, HCBT, DAT) (St-Pierre and De Luca, 2000). Over 60 BAHD acyltransferases have been assigned functions in plants based on the genetic mutants and/or biochemical assays (Tuominen et al., 2011). Aromatic acylation and aliphatic acylation are two major types of acylation: the former usually involves the addition of cinnamoyl, coumaroyl, caffeoyl groups and the latter is normally malonylation (Winkel-
Shirley, 2001) (Fig. 1). The acyltransferases that have anthocyanins as acceptor substrates constitute a superclade of phylogenetic tree but functional predictions on the catalytic versatility is difficult to infer from the primary sequences alone (Luo et al., 2007).

Horizontal gene transfer (HGT) refers to the transfer of genetic material to non-offspring genomes (Bock, 2010; Zhang et al., 2014b). In plants, most HGTs involve mitochondrial DNA transferred to nuclear or mitochondrial genomes (Davis and Xi, 2015; Rice et al., 2013; Xi et al., 2013). Such transfer events have been considered neutral or to confer no advantage to the recipients because the majority of transferred genes evolve into pseudogenes (Bock, 2010). Gene transfer between plant nuclear genomes is important because the newly introduced genes may endow the recipients with adaptive advantages. Nevertheless, such transfers are rare in plants. Furthermore, nuclear transfers that have been reported mostly involve parasitic plants with a narrow host range (Xi et al., 2012). Very few nuclear genes were found to have been transferred between parasitic plants with a broad host range and their hosts. Two notable exceptions include albumin 1 and strictosidine synthase-like (SSL) genes from Fabaceae and Brassicaceae hosts to both Phelipanche aegyptiaca and Cuscuta australis, respectively (Zhang et al., 2013, 2014a).

Plants in the genus Cuscuta (Convolvulaceae) are world-wide obligate stem holoparasites. There are about 200 species of this genus and most have broad host ranges (Garcia et al., 2014), including Brassicaceae, Leguminosae, Solanaceae, Apiaceae, Asteraceae, and Cucurbitaceae. By establishing intimate connections with their hosts through haustoria, Cuscuta can take up water, nutrients, and macromolecules, including proteins and mRNAs (Jiang et al., 2013; Kim and Westwood, 2015). Relatively high frequency genetic material exchanges enhance the occurrence of horizontal gene transfer. Surprisingly, except for the foreign albumin 1 and SSL gene, no nuclear gene transfer events have been reported in Cuscuta to date. Utilizing the next-generation sequencing technology, we generated eight transcriptomes from different tissues and developmental stages of C. australis. Multiple Cuscuta pentagona transcriptomes are publicly accessible, offering reliable data sources for screening for foreign genes at the transcriptional level. In this study, we show that one putative anthocyanin acyltransferase gene (AT-like) has been transferred from Fabaceae to Cuscuta, although the possibility of convergent evolution of these genes can not be absolutely ruled out.

2. Materials and methods

2.1. Transcriptome screening for foreign genes in C. australis

To identify HGT events between host and parasitic plant, we screened C. australis transcriptomes for foreign genes according to the procedure described previously (Zhang et al., 2014a). In brief, a transcriptome assembly was obtained by combining all the RNA-seq datasets from eight tissues, seeds, germinated seeds, seedlings, pre-haustoria, stems, buds, flowers, and capsules using Trinity v 2.0.6 (Grabherr et al., 2011). The predicted coding regions with a contiguous amino acid length ≥100 were screened using AlienG (Tian et al., 2011). The alien origin of the candidate genes was predicted if the score ratio of the first non-Convolvulaceae hit to the first Convulvulaceae species hit was more than 1.2. These candidates were then manually identified as horizontally transferred genes.

2.2. Gene identification and homolog search for putative foreign genes in C. australis genome assembly and the transcriptomes of C. pentagona

We produced 73 Gb (about 170X coverage) of high quality Illumina paired-end reads from a library with ~270-bp insertion fragments. The genome assembly was performed using the SOAPdenovo package (Li et al., 2010) with a K-mer size of 41. The transcriptome contigs encoding foreign genes were used as queries to search our local genome assembly by BLASTn to obtain the corresponding DNA and homology sequences. The gene structures in the homology sequences were determined using our transcriptome data and by BLAST search (BLASTx) against the nr database. The candidate foreign genes were used as queries to search online against the RNA-seq datasets of C. pentagona deposited in the NCBI.
SRA database using BLASTn and the read pairs with more than 94% identity were extracted as the evidence of transcription.

2.3. Phylogenetic analyses of the anthocyanin acyltransferase gene in C. australis

To investigate the origin of the AT-like (anthocyanin acyltransferases-like) gene in Cuscuta, protein homologs were extracted from representative species with genome sequences (Supplementary Table 1), and our homemade genome assembly of C. australis by BLAST search using the candidate anthocyanin acyltransferase gene from C. australis as queries. To obtain insight into the putative function of the anthocyanin acyltransferase gene in Cuscuta, the putative genes or characterized proteins belonging to the clade I (Luo et al., 2007) and clade II (Tuominen et al., 2011) from two previous studies, together with the candidate BAHD acyltransferase gene and homologs in C. australis, were also collected. The sequences were aligned with ClustalX v2.1 (Thompson et al., 1997), visually inspected, and manually refined. Gaps and ambiguous sites were removed from the alignment. ModelGenerator (v.851) (Keane et al., 2006) was used to find the best-fitting model of protein substitution. The protein phylogenetic tree was inferred under maximum likelihood optimization using RAxML v 8.0.0 with the parameters "-f a,-m PROTCAMMajTTF,-# 100" (Stamatakis, 2006). Trees were viewed and edited using MEGA7 (Kumar et al., 2016).

2.4. Expression level estimation of the candidate BAHD acyltransferase gene in C. australis and C. pentagona

The expression levels of the AT-like gene in different tissues at different developmental stages of C. australis were estimated using RSEM (Li and Dewey, 2011). We mapped all clean reads to the Trinity assembly, and to obtain normalized expression levels we used the fragments per kilo base of exon per million fragments mapped (FPKM) values of the sequences. The mean FPKM values of the AT-like gene from 10 accessible public RNA-seq datasets in C. pentagona (Ranjan et al., 2014), from the tissues of seeds, seedlings, stems, pre-haustoria, haustoria, and flowers, were calculated using the RPKM formula \( \frac{10 \times C}{\text{N} \times \text{L}} \), where \( C \) is the number of read pairs mapped to a gene, \( N \) refers to the total mapped read pairs in the experiment and here is approximate to the total number of raw read pairs, \( L \) is the exon length in base-pairs for a gene. The SRA accession numbers of the 10 public RNA-seq datasets of C. pentagona were SRX345073, SRX345113, SRX345282-3, SRX345400-1, and SRX345411-4.

3. Results

3.1. The abnormal affiliation of one anthocyanin acyltransferase gene from Cuscuta with the homologs from Fabaceae

Using a pipeline we developed to screen transcriptomes for foreign genes, we found that a 1853-bp sequence in C. australis transcriptomes exhibited as high as 56% and 66% identities to the Fabaceae coumaroyl-CoA:anthocyanidin 3-O-glucoside-6-O-coumaroyltransferases at the amino acid and nucleotide levels, respectively. However, this sequence in Cuscuta was found to be only 24% at the amino acid level with Ipomoea and there was no significant similarity at the nucleotide level. We then searched the transcriptome datasets of C. pentagona deposited in NCBI SRA database and found 22 read pairs mapped to the C. australis transcript with high identity (Supplementary File 1). Therefore, the corresponding genes are likely present in C. pentagona as well. Using the 73 Gb high quality Illumina paired-end reads, we used SOAPdenovo to generate a genome assembly of 343.9 Mb, which covers 81.7% of the genome given that the genome size of C. australis was estimated to be 421 Mb by flow cytometer in our lab. We searched the genome assembly using the above transcriptome contig and found a 4165-bp scaffold that contained a complete open reading frame (ORF) of 1503 bp (500 amino acids) without introns. Notably, the homologous genes in Fabaceae are intron-less. Furthermore, the transcription of this gene can be detected in each of the transcriptomes from different tissues of C. australis and C. pentagona. The presence of this gene in the genome assembly of C. australis and multiple transcriptomes of C. australis and C. pentagona makes the possibility of contamination from other organisms very low.

To confirm the possible transfer of this gene from Fabaceae to Cuscuta, we constructed a phylogeny of the BAHD acyltransferase genes by collecting homologs from 37 species whose genomes have been sequenced and those with genetic and biochemical evidence (Supplementary Table 1). Multiple copies were found in most species and the homologs from monocots clustered at the base of the phylogenetic tree, which had been illustrated in other studies (Luo et al., 2007; Tuominen et al., 2011). To construct a more reliable alignment and phylogenetic tree, the homologs from monocots were excluded from further analysis and only those from dicots were used. This gene tree formed two clusters (Fig. 2). Cluster I contained homologs from asterids and Vitis, including the published anthocyanin acyltransferase and anthocyanin malonylesterase activity proteins, which corresponded to the subclade 2 of the anthocyanin superclade (Luo et al., 2007). Cluster II contained proteins from 28 of the 31 representative dicot species, which greatly expanded subclade 1 of the anthocyanin superclade (Luo et al., 2007). Both cluster I and II were included in the clade la by Tuominen et al. (2011).

The A. thaliana enzymes in cluster II have been shown to possess anthocyanin acyltransferase activity (Luo et al., 2007). Therefore, proteins in this cluster are likely to be anthocyanin acyltransferases. Five genes were found in C. australis, four in cluster I, which probably have anthocyanin acyltransferases or anthocyanin malonyltransferase activity. In addition, one gene in cluster II likely possesses AT-like activity (named CaATL in this study for convenience, accession number KX019005). As expected, CaATL formed a sister branch with the homologs from Fabaceae and the abnormal affiliation was supported by 100% bootstrap value, which suggested that horizontal gene transfer probably occurred between Fabaceae and the ancestor of Cuscuta. No other branches with obvious abnormal species affiliation were found in the gene tree of the BAHD family proteins, although some main branches contained species from multiple lineages, which may be caused by gene losses or convergent evolution as speculated by other researchers (Luo et al., 2007).

3.2. Domain structure and motif comparison of CaATL with other closely related functional BAHD members

Analysis of domain structures in the anthocyanin acyltransferases (AT), anthocyanin malonyltransferase (MAT), and CaATL by InterPro (Quevillon et al., 2005) indicated that all these proteins possess two tandemly arranged chloramphenicol acetyltransferase-like domains (IPR023213), each ~205 aa long. Both ATs and MATs can catalyze the acylation of the 6′ position of the 3-O-glucoside (3ATs and 5MATs, respectively) and the 5′ position of the 5-O-glucoside (5ATs and 5MATs, respectively) (Luo et al., 2007). Therefore, these previously characterized motifs were tested by aligning the known representatives of 3ATs, 3MATs, 5ATs and 5MATs, CaATL with their closely related putative
Fig. 2. The phylogeny of CaATL and its BAHD family homologs. The clusters I and II are labeled in the main branches. Numbers above the branches show bootstrap support values from maximum likelihood analysis. The characterized functional proteins from asterids and Arabidopsis are indicated in bold after the species names. The lineage information is indicated after the vertical bars with the branches colored, asterids in orange, fabids in blue, Vitis vinifera in purple, malvids in light green, Caryophyllales in red, the stem eudicotyledons in deep green. The horizontally transferred genes in *C. australis* from Fabaceae are indicated in brown.
Fig. 3. Multiple sequence alignment of representative ATs, MATs, CaATL and its homologs in Fabaceae. The conserved and specific motifs are indicated in green and yellow boxes. The HXXD motif in CaATL and some Fabaceae is shown in red boxes. The sequence names are displayed in the order of abbreviation names, accession numbers, genera, and species names. Dashes indicate that the sequences are incomplete or gaps introduced in the alignment. Background colors indicate the degree of conservation of the sites.
3ATs in *Medicago truncatula* (Fig. 3). The three conserved motifs of the BAHD family, consensuses of PXLKXSLSX/(T/A)/L around position 69–79, QX(T/A)XPF(N/G)XG around position 173–181, and (D/Z)FXGXG(K/R)/P/XK around position 452–460, were detected by WEBLOGO (http://weblogo.berkeley.edu/logo.cgi) (Fig. 3, Supplementary File 2). Two specific motifs, P(L/V/T)/S(F/L)/D around position 34–39 and PXXYFGNC adjacent to position 369–376, were also detected (Fig. 3, Supplementary File 2).

Interestingly, one amino acid deletion occurred in the conserved HXXXD motif, which was changed into HXXD in the homologs in some fabaceous plants and CaATL (Fig. 3). These lines of evidence support the hypothesis that CaATL is a BAHD family member that likely uses anthocyanins as substrates, and is closely related to the homologs from Fabaceae.

### 3.3. The expression of the ATL genes in *C. australis* and *C. pentagona* implies that they still possess functional protein activity

The presence of the ATL genes in the RNA-seq data of *C. australis* indicates that they are actively transcribed. To determine how the expression level changes in different tissues at different developmental stages, we mapped all clean paired-end reads to our own transcriptome. The FPKM values of ATL varied within tissues at different developmental stages. No transcription was detected in germinated seeds or of ATL modi the gene in secondary metabolism, namely, anthocyanin transcriptional activity in old tissues is consistent with the function of old stems, which is 1.8-fold that of young stems. The high transcriptional activity in old tissues at different developmental stages, we mapped all clean paired-end reads to our own transcriptome. The FPKM values were estimated to be 0.23 using the FPKM equation (Fig. 4).

Seeds, seedlings, and capsules had similar transcriptional levels with FPKM values less than 0.1. Pre-haustoria, young stems, and buds exhibited higher expression levels with FPKM values from 0.45 to 0.68. Notably, the FPKM value reached a maximum of 1.26 in flowers, fruits, and leaves. Rich in phenolic hydroxyl groups, anthocyanins have high antioxidant properties and protect plants from reactive oxygen species produced in metabolism thus improving resistance to abiotic stresses, such as UV, drought, and low temperature, as well as biotic stresses like pathogens and herbivorous insects (Xie et al., 2013). In plants, anthocyanin acylation is catalyzed by BAHD family proteins, which show great versatility in substrate specificity (Rinaldo et al., 2015) and are able to evolve new substrate specificities rapidly (Luo et al., 2007). These traits are consistent with the phylogeny showing that cluster I is composed of 3AT, 5AT, 3MAT and 5MAT and the 3ATs are both present in cluster I and II (Fig. 2).

By searching the genome assembly, five genes belonging to the BAHD family, having anthocyanin as substrates, were identified in *C. australis*. Four of them formed a sister branch with the homologs from the relative *Ipomoea* and should be native copies. CaATL and the Fabaceae homologs constituted a cladé with 100% bootstrap support, and this gene very likely was transferred from a fabaceous plant to the common ancestor of *Cuscuta* since *C. pentagona* also possesses this gene. The reverse transfer direction is impossible since no other asterid genes were present in cluster II. The identity of the Fabaceae donor was unclear from the phylogeny given that CaATL forms a basal branch of the homologs of different Fabaceae species. The occurrence of HGT is probably mediated by direct contact via haustoria of *Cuscuta* as Fabaceae is its preferred hosts.

The acylation of anthocyanins affects its stability and light absorption in solution. In Arabidopsis, At3AT1 and At3AT2 influence the stability of anthocyanins at neutral pH and the anthocyanin absorption maxima. The higher transcription level in old compared to young stem tissues suggested this gene might still perform its functions in secondary metabolism by stabilizing anthocyanins at neutral pH and thus improving the biotic and abiotic resistance of *Cuscuta* to stresses such as UV radiation and insect herbivory. Considering the separation of cluster I with cluster II in the phylogeny, we speculate that the horizontally acquired CaATL probably endowed *Cuscuta* with certain adaptive characters, such as changes in expression pattern under stresses or varied substrate specificity. Further investigations and functional analysis are needed to understand the benefits of HGT on the evolution of this special holoparasitic plant.

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### Appendix A. Supplementary data

Supplementary data related to this article can be found at http://dx.doi.org/10.1016/j.pld.2016.04.002.

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