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# Identification and Characterization of the *CCoAOMT* Gene Family in Apple, Chinese White Pear, and Peach

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**ABSTRACT.** Lignin is one of the main components of plant cell walls, which provides mechanical support for plants and also contributes to resisting against plant pathogenic fungi. In the fruit industry, the lignin content can affect the quality of fruit. The biosynthesis of lignin involves a variety of enzymes, of which *caffeoyl-CoA 3-O-methyltransferase (CCoAOMT)* is a class of methyltransferases that plays an essential role in lignin biosynthesis. Studies have been conducted on the *CCoAOMT* gene family in several species, including arabidopsis (*Arabidopsis thaliana*), black poplar (*Populus nigra*), and cotton (*Gossypium hirsutum*). Still, there is relatively little research on this gene family in the Rosaceae. In this study, we used bioinformatics to identify and characterize the *CCoAOMT* gene family in apple (*Malus domestica*), chinese white pear (*Pyrus bretschneideri*), and peach (*Prunus persica*). In total, 35 *CCoAOMT* genes were identified in the three Rosaceae species: 8 from chinese white pear, 12 from apple, and 15 from peach. By using structure analysis and collinearity analysis, we found 12 conserved motifs and 12 pairs of *CCoAOMT* genes with collinearity. In the phylogenetic tree, the gene family was mainly divided into two groups. The genes had different expression patterns during the growth and development stage of fruit, a finding that is consistent with the pattern of lignin accumulation. This study will be beneficial for further study of *CCoAOMT* genes.

Lignin is one of the main components of plant cell walls. It is a phenol polymer composed of coniferyl alcohol, sinapyl alcohol, and p-coumaryl alcohol subunits. Lignin can enhance the strength of cell walls to maintain a plant's structure and also act as a barrier against pathogen infection. When plants are infected by pathogens, the enzymes related to lignin biosynthesis become active, and the lignin content accumulates to resist the pathogen (Huang, 2001). At a global level, lignin is one of the most abundant renewable biomass resources on earth and is widely associated with a range of industrial and agricultural production systems. However, in some contexts, the existence of lignin can cause some problems. For example, the lignin in animal feed has a deleterious effect on the efficiency of digestion and absorption (Srivastava et al., 2012).

In the fruit industry, attention needs to be paid to the effect of lignin on fruit quality. On one hand, as mentioned, lignin

can resist plant pathogenic fungi and thereby help avoid any decrease in fruit quality. For example, gray mold caused by *Botrytis cinerea* is a common fungal disease in postharvest apple [*Malus domestica* (Tang et al., 2012; Yuan et al., 2014)]. When an apple is infected with this pathogen, it will rot and with it comes a decline in fruit quality, even under cold storage conditions. Although there are some agricultural methods to resist this disease, enhancing apple's inherent disease resistance is the most effective method. The induction of lignin synthesis and other phenolic compounds may help in this approach (Valentines et al., 2005).

The content of lignin itself can affect other aspects of fruit quality, and the most representative example is the effect of lignin on the quality of chinese white pear fruit (*Pyrus bretschneideri*). Stone cells, formed by the accumulation of lignin and other substances, represent one crucial factor that affects the taste of fruit (Rogers and Campbell, 2004). The stone cells mainly exist in groups, and the size and the diameter of these groups both affect the taste of such fruit (Qiao et al., 2005). The russet pericarp characteristic is another specific factor that affects the appearance of pear (*Pyrus* sp.) fruit. This color of the pericarp is formed by the accumulation of the cork layer (Teng et al., 2005), of which lignin is one of the main components (Lopes et al., 2001; Pereira, 1988). Another defect related to lignin is found in peach (*Prunus persica*), a Rosaceae fruit tree that originated in China. The formation of the endocarp of peach

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seed is a process dependent on lignification. During the growth and development of peach fruit, a split-pit phenomenon may occur in which the suture of the endocarp does not close tightly or has an obvious gap (Yang et al., 2009); this causes the fruit to be easily infected by a disease. A study has found that the closure mechanism of the peach endocarp is similar to the development of the silique in arabidopsis [*Arabidopsis thaliana* (Tani et al., 2009)]. It is known that certain transcription factors affect the cracking of the silique by regulating the lignification in this species, and this accumulation of lignin may be related to the split-pit phenomenon of peach fruit (Hu et al., 2012). Therefore, studying the mechanism of the lignification of such fruit may help control their lignification to increase their commercial value.

Considering the process of lignification, the biosynthesis of lignin begins with a series of complex reactions that produce three major monomers: p-hydroxyphenyl (H) lignin, guaiacyl (G) lignin, and syringyl (S) lignin. These three monomers form lignin (Vanholme et al., 2010). This process involves a variety of enzymes, of which caffeoyl-CoA 3-O-methyltransferases (*CCoAOMT*), a class of S-adenosyl-L-methionine methyltransferases (Boerjan et al., 2003), plays an important role. *CCoAOMT* enzymes are encoded by specific gene families, members of which have been cloned from many species, including poplar [*Populus* (Chen et al., 2000)], arabidopsis (Do et al., 2007), cotton [*Gossypium hirsutum* (Ni et al., 2010)], *Neosinocalamus affinis* (Wu et al., 2012), and *Citrus maxima* (Xu et al., 2014), and there have been studies that have found the specific relationship between the *CCoAOMT* gene family and the lignin. For example, three members of the *CCoAOMT* gene family have been cloned from rice (*Oryza sativa*) and are found to be closely related to the lignification process in this species (Zhao et al., 2004). Similarly, a *CCoAOMT* gene was cloned from pears, and it was found that the gene expression level was similar to the trend of stone cell production in pear fruit (Wang et al., 2015). Studies on tobacco [*Nicotiana tabacum* (Zhong et al., 1998)], black poplar [*Populus nigra* (Zhong et al., 2000)], and alfalfa [*Medicago sativa* (Guo et al., 2001)] found that the deletion of *CCoAOMT* genes or the inhibition of *CCoAOMT* gene expression both reduced the lignin content. The expression of *CCoAOMT* genes was significantly decreased in transgenic maize (*Zea mays*) expressing an RNA interference construct and led to a 22.4% decrease in lignin content (Li et al., 2013). Similarly, the tobacco transformed with a full-length RNA interference fragment of *CCoAOMT* had a reduced level of gene expression and consequently a significant decrease in G-lignin content (Chen et al., 2018). Most recently, compared with the wild type, transgenic *Betula platyphylla* with antisense *CCoAOMT* genes showed a significantly lower lignin content (Yao et al., 2019).

At present, much research has been conducted on *CCoAOMT* genes, and the whole genome sequence of some Rosaceae species has been completed, like apple (Velasco et al., 2010), pear (Wu et al., 2013), and peach (Verde et al., 2013). However, due to the limited bioinformatics analysis of the *CCoAOMT* gene family in Rosaceae species, we still know little about the characterization of *CCoAOMT* genes. The present study was designed to identify and analyze characteristics of the *CCoAOMT* gene family in these representative species of the Rosaceae to explain its collinearity and determine its evolution in these species, including apple, chinese white pear, and peach. The study involved structure analysis, phylogenetic analysis, collinearity analysis,

and expression analysis, and was to provide a theoretical basis for further research on the specific function of *CCoAOMT* genes and lignin biosynthesis.

## Material and Methods

**IDENTIFICATION OF *CCoAOMT* GENES IN APPLE, CHINESE WHITE PEAR, AND PEACH.** The entire nucleotide sequences, amino acid sequences, and gene annotation files of pear were downloaded from GigaScience database [GigaDB (Wu et al., 2013)]. The complete nucleotide sequences, amino acid sequences, and gene annotation files of apple were downloaded from the Apple Genome and Epigenome [AGE version 1.1 (Daccord et al., 2017)]. The whole nucleotide sequences, amino acid sequences, and gene annotation files of peach were downloaded from the Genome Database for Rosaceae [GDR (Jung et al., 2019)]. Then we constructed an amino acid sequence database of apple, chinese white pear, and peach to perform protein basic local alignment search tool (BLASTP). The query sequence used in the BLASTP was the *CCoAOMT* amino acid sequence of arabidopsis (NP\_564916.2), which was downloaded from the National Center for Biotechnology Information [NCBI, Bethesda, MD (Theologis et al., 2000)]. The resulting sequences with E value  $\leq 10^{-10}$  were treated as candidate *CCoAOMT* sequences. We used NCBI's CD-Search tool (Lu et al., 2020) to check whether the candidate sequence has the characteristic domain of *CCoAOMT*. If the candidate proteins contained domain of Methyltransf\_3 (PFAM: PF01596), it would be considered as the members of *CCoAOMT* gene families (Pospiech et al., 1996). Basic data about the *CCoAOMT* proteins were calculated, including protein length, molecular weight, isoelectric point, and transmembrane helices.

**PHYLOGENETIC ANALYSIS OF *CCoAOMT* GENES IN APPLE, CHINESE WHITE PEAR, AND PEACH.** In this study, we used the neighbor joining method of MEGA 5.0 (Tamura et al., 2011) to construct the phylogenetic tree, and the website phyML 3.0 [1 Mar. 2020 (Guindon et al., 2010)] was used to construct the maximum likelihood tree as verification. We chose *CCoAOMT* genes in arabidopsis as outgroup. The *CCoAOMT* potential protein sequences of arabidopsis were also identified using BLASTP with E value  $\leq 10^{-10}$  among the whole genome. The data of arabidopsis whole protein sequence were downloaded from the Arabidopsis Information Resource [TAIR (Philippe et al., 2012)]. Multiple sequence alignment and neighbor joining tree construction of *CCoAOMT* protein sequences in apple, chinese white pear, peach, and arabidopsis were performed using MEGA 5.0 with a selection of Poisson model and pairwise deletion of vacant/missing data. The phylogenetic tree was constructed with a bootstrap value of 1000.

**MOTIF ANALYSIS OF *CCoAOMT* GENES IN APPLE, CHINESE WHITE PEAR, AND PEACH.** Motif analysis was performed using the Web site multiple em for motif elicitation [MEME (Bailey and Elkan, 1994)]. Based on the MEME motif, a phylogenetic tree was constructed to combine the information of motifs and phylogenetic analysis.

**POSITIVE SELECTION ANALYSIS OF *CCoAOMT* GENES IN APPLE, CHINESE WHITE PEAR, AND PEACH.** The positive selection analysis was based on the preceding motif analysis, and the genes were divided into different groups to be analyzed. First, a multiple sequence alignment of *CCoAOMT* amino acid sequences was performed using CLUSTALW (Larkin et al., 2007). The alignment

results and the corresponding nucleotide sequences were input into the pal2nal (Suyama et al., 2006) to generate a file in the format of phylogenetic analysis by maximum likelihood [PAML (Xu and Yang, 2013)]. Then a phylogenetic tree was needed as well. The whole positive selection analysis was performed using the X version of PAML with a model of codons. The likelihood ratio test (LRT) was also required in this analysis, using the  $\chi^2$  test to calculate whether there was a significant difference. Four of the different models of PAML were used in this study: M0 (Goldman and Yang, 1994; Yang and Nielsen, 1998), M3, M7, and M8 (Yang, 2000a, 2000b). The LRT statistic between M0 and M3 was used to evaluate whether there was a significant difference in the selection pressure between the sites. The LRT statistic between M8 and M7 was used to evaluate whether there was a positive selection between the sites. If the LRT statistic between M8 and M7 had a significant difference and the  $\omega$  value of M8 was more than 1, then the Bayesian method was used to evaluate the positively selected sites.

In general, if the results of a group met these three conditions, the group could be considered to have a positive selection: 1) LRT statistic between M8 and M7 had a significant difference; 2) the  $\omega$  value of M8 was more than 1; and 3) there existed positively selected sites after evaluation by the Bayesian method.

**COLLINEARITY ANALYSIS OF *CCoAOMT* GENES IN APPLE, CHINESE WHITE PEAR, AND PEACH.** Multiple Collinearity Scan X [MCS-X (Tang et al., 2008)] was used to conduct the collinearity analysis of each pair of species using a basic local alignment search tool (BLAST) file, which was obtained by blasting the whole genomes and annotated genes.

**EXPRESSION ANALYSIS OF *CCoAOMT* GENES IN APPLE, CHINESE WHITE PEAR, AND PEACH.** To examine the expression of *CCoAOMT* genes among chinese white pear genes, chinese white pear fruit samples at 15, 36, 80, 110, 145, and 167 d after flowering (DAF) were used (Wu et al., 2013), which was calculated in reads per kilobase million (RPKM). To examine the expression of *CCoAOMT* genes of peach, the fruit samples of peach at 41, 54, 69, 83, 111, and 125 DAF were used [GEO accession GSE71561 (Zaffolon et al., 2017)]. To examine the expression of *CCoAOMT* genes in apple fruit, the apple fruit samples (hypanthium) at young fruit stage, expanding stage and maturity stage were collected in Zhengzhou Fruit Research Institute (Zhengzhou, China), Chinese Academy of Agricultural Sciences. The apple samples included the wild accession Xifuhaitang and the cultivar Golden Delicious. The flesh tissues of 0.2 to 0.3 g apple were ground with liquid nitrogen, and RNA extracted by improved CTAB method 61. According to the Illumina TruSeq (San Diego, CA) RNA sample preparation process, the complementary DNA library was constructed. The size and purity of the complementary DNA library were determined by Agilent DNA1000 kit and Agilent 2100 bio-assay (Agilent Technologies, Santa Clara, CA). Raw reads of the experiment are submitted to NCBI SRA database (Bethesda, MD), the accession number of the study is SRR9291270 and SRR9291271. Transcripts per million (TPM) was calculated after sequencing by Illumina technique.

## Results

***CCoAOMT* GENES IN APPLE, CHINESE WHITE PEAR, AND PEACH.** In total, 35 *CCoAOMT* predicted proteins were identified in the three Rosaceae species: eight from chinese white pear, 12 from apple, and 15 from peach by BLASTP (Table 1). The longest of

the 35 sequences is *MD02G1073400*, which has 525 amino acids and a molecular weight of 58272.8. The shortest sequence is *MD02G1073300*, which has only 74 amino acids and a molecular weight of only 8441.1. The length of most *CCoAOMT* amino acid sequences is between 100 and 300 amino acids and their molecular weights range from 10,000 to 30,000. The isoelectric points of *CCoAOMT* sequences distributed between 4 and 10. Twelve *CCoAOMT* amino acid sequences were found to have a transmembrane segment: *Pbr025246.1*, *MD16G1119300*, *MD02G1230900*, *MD05G1209400*, *Prupe.1G227100.2*, *Prup.2G107200.1*, *Prup.2G107300.1*, *Prup.4G148100.3*, *Prup.4G148100.4*, *Prupe.4G148100.5*, *Prupe.4G148100.1*, and *Prupe.7G131200.1*. Their transmembrane segments were similar in length, all between 25 and 30 amino acids. Based on the location of several of the predicted *CCoAOMT* proteins, a number of them were derived from a single gene, representing splice variations, which is shown in Table 1. Therefore, there are a total of 22 *CCoAOMT* genes in our study considering the splice variations. However, splice variations are a great source for divergence in function-duplicated genes.

**PHYLOGENETIC ANALYSIS OF *CCoAOMT* GENES IN APPLE, CHINESE WHITE PEAR, AND PEACH.** Thirteen *CCoAOMT* protein sequences were identified in arabidopsis. A phylogenetic tree of the *CCoAOMT* gene family in apple, chinese white pear, peach, and arabidopsis was constructed (Fig. 1). The phylogenetic tree was divided into two groups: A and B. The A group consisted of 39 sequences and the B group consisted of nine sequences. The A group was further divided into four subgroups: A1, A2, A3, and A4. The A1 subgroup included only the *CCoAOMT* genes from apple, chinese white pear, and peach, indicating that this group was a unique group of Rosaceae. The A2 subgroup included only *CCoAOMT* genes in arabidopsis. The A3 subgroup did not have the *CCoAOMT* genes of chinese white pear, and there was only one *CCoAOMT* gene from arabidopsis (*AT4G26220.1*) in this subgroup. These results indicated that the differentiation of A1 and A2 was after the species divergence. The A4 subgroup and B group all contained *CCoAOMT* genes from the four species. The sequence distribution of the A4 subgroup and B group was similar: *CCoAOMT* genes of apple and chinese white pear were in the closest position of the phylogenetic tree, and *CCoAOMT* genes of peach and arabidopsis each had a separate branch. Among the 35 *CCoAOMT* proteins identified in the Rosaceae species, the *CCoAOMT* genes in apples and chinese white pear were always the most similar. The distribution of *CCoAOMT* genes on the phylogenetic tree was consistent with their species taxonomy (Fig. 2).

**MOTIF ANALYSIS IN *CCoAOMT* GENE FAMILY OF APPLE, CHINESE WHITE PEAR, AND PEACH.** Twelve conserved motifs were predicted in this study. The conserved motifs, together with a phylogenetic tree, were integrated into the same figure (Fig. 3). This figure was divided into four blocks named 1, 2, 3, and 4. None of the 12 motifs existed in all sequences, and motifs 1, 2, 3, 4, 5, 7, and 8 frequently appeared in the sequences. Motifs 6 and 11 were only present in the six sequences of block 4: *Pbr025246.1*, *MD05G1209400*, *Prupe.4G148100.3*, *Prupe.4G148100.4*, *Prupe.4G148100.5*, and *Prup.4G148100.1*, which were on the same branch of the phylogenetic tree. Except for *Prupe.8G128100.3*, seven genes among the eight sequences in block 3 all had motif 12; these seven genes were *Prupe.7G131200.1*, *Prupe.8G128100.1*, *Prupe.8G128100.4*, *Pbr038709.1*, *MD05G1209400*, *Pbr034039.1*, and *MD00G1088100*. *MD02G1073300* had only one motif, because this sequence only had 74 amino acids. *MD02G1073400* had the most motifs, of which motifs 1, 2, 3, 4, 5, 7, and 9 all appeared twice.

Table 1. Basic properties of *caffeoyl-CoA 3-O-methyltransferase (CCoAOMT)* protein genes in apple, chinese white pear, and peach.

Species	No.	Gene	Start	End	Protein length (amino acids)	Molecular wt (Da)	pI <sup>2</sup>	Transmembrane
Pear	PbCCoAOMT01	<i>Pbr015171.1</i>	6095377	6096533	161	17590.3	8.04	None
	PbCCoAOMT02	<i>Pbr015180.1</i>	6147661	6151135	235	26323.8	4.81	None
	PbCCoAOMT03	<i>Pbr019305.1</i>	21283935	21288384	235	26697.3	5.39	None
	PbCCoAOMT04	<i>Pbr025246.1</i>	19507699	19511795	307	34112.3	8.84	# 1:233–261 29aa
	PbCCoAOMT05	<i>Pbr028157.1</i>	254343	255523	149	16337.1	8.04	None
	PbCCoAOMT06	<i>Pbr028166.1</i>	313064	316536	235	26308.8	4.9	None
	PbCCoAOMT07	<i>Pbr034039.1</i>	5204856	5206214	247	27850.1	5.39	None
	PbCCoAOMT08	<i>Pbr038709.1</i>	10033	11412	247	27814.1	5.18	None
	MDCCoAOMT01	<i>MD00G1088100</i>	17871472	17873066	243	27330.6	5.97	None
	Apple	MDCCoAOMT02.1	<i>MD02G1073100</i>	5899257	5901470	240	26717.9	5.17
MDCCoAOMT02.2		<i>MD02G1073300</i>	5902929	5903673	74	8441.1	4.52	None
MDCCoAOMT02.3		<i>MD02G1073400</i>	5904355	5908568	525	58272.8	5.41	None
MDCCoAOMT03.1		<i>MD02G1230800</i>	27660687	27660983	98	11216.9	5.18	None
MDCCoAOMT03.2		<i>MD02G1230900</i>	27666886	27669257	118	13667.8	8.19	# 1:44–72 29aa
MDCCoAOMT04		<i>MD05G1209400</i>	34097139	34100658	307	34051.2	8.79	# 1:233–261 29aa
MDCCoAOMT05		<i>MD05G1083900</i>	17486739	17488329	247	27774	5.39	None
MDCCoAOMT06		<i>MD13G1117900</i>	8626272	8629143	237	26678.1	4.83	None
MDCCoAOMT07.1		<i>MD16G1118200</i>	8401568	8407573	251	28336.7	6.4	None
MDCCoAOMT07.2		<i>MD16G1119200</i>	8519352	8521735	235	26207.7	4.64	None
MDCCoAOMT07.3	<i>MD16G1119300</i>	8523351	8524410	175	19463.9	4.48	# 1:8–36 29aa	
Peach	PpCCoAOMT01.1	<i>Prupe.1G227100.1</i>	24032115	24034470	234	26384.9	5.17	None
	PpCCoAOMT01.2	<i>Prupe.1G227100.2</i>	24032115	24034470	166	18885.4	5.01	# 1:0–24 25aa
	PpCCoAOMT01.3	<i>Prupe.1G227100.3</i>	24032115	24033691	222	25148.5	4.85	None
	PpCCoAOMT02.1	<i>Prupe.2G107200.1</i>	16475915	16478143	237	26847.1	5.06	# 1:46–71 26aa
	PpCCoAOMT02.2	<i>Prupe.2G107300.1</i>	16478452	16480777	235	26751.3	5.72	# 1:158–185 28aa
	PpCCoAOMT03.1	<i>Prupe.4G148100.3</i>	8460021	8464276	299	33087	9.18	# 1:219–247 29aa
	PpCCoAOMT03.1	<i>Prupe.4G148100.4</i>	8460023	8464276	294	32569.5	9.18	# 1:219–247 29aa
	PpCCoAOMT03.1	<i>Prupe.4G148100.5</i>	8460023	8464276	308	34064	9.02	# 1:233–261 29aa
	PpCCoAOMT03.1	<i>Prupe.4G148100.1</i>	8460012	8464276	313	34581.5	9.02	# 1:233–261 29aa
	PpCCoAOMT04	<i>Prupe.7G131200.1</i>	15043867	15045183	186	21009.2	5.23	# 1:41–65 25aa
	PpCCoAOMT05	<i>Prupe.7G214300.1</i>	19456381	19457966	238	26819.3	6.53	None
	PpCCoAOMT06	<i>Prupe.7G214400.1</i>	19458449	19460610	240	26878	4.81	None
	PpCCoAOMT07.1	<i>Prupe.8G128100.1</i>	15185748	15187399	247	27831.9	5.37	None
	PpCCoAOMT07.2	<i>Prupe.8G128100.3</i>	15185606	15187399	187	20955.5	5.06	None
PpCCoAOMT07.3	<i>Prupe.8G128100.4</i>	15185748	15187399	246	27702.8	5.65	None	

<sup>2</sup>Isoelectric point.

**POSITIVE SELECTION ANALYSIS OF *CCoAOMT* GENE FAMILY IN APPLE, CHINESE WHITE PEAR, AND PEACH.** A positive selection analysis was performed separately according to the groups A1, A2, A3, A4, and B of Fig. 1. As shown in Table 2, the  $\omega$  value of each group was less than 1, indicating that the purification selection effect occurred in all five groups. The probability values of the LRT statistic between M3 and M0 of A1, A3, and B groups were less than 0.05, indicating that the three groups had a significant difference in the selection pressure. In contrast, the other A2 and A4 groups did not have a significant difference in the selection pressure. The LRT statistic between M8 and M7 of each group did not have a significant difference, indicating that there was no positive selection in these five groups. The program evaluated no reliable positively selected site. In conclusion, none of the groups met the three conditions at the same time, so the *CCoAOMT* genes were considered to have not undergone any positive selection.

**COLLINEARITY ANALYSIS OF *CCoAOMT* GENE FAMILY IN APPLE, CHINESE WHITE PEAR, AND PEACH.** Through collinearity analysis by MCSanX, 55,158 pairs of collinear genes were identified in apple

and chinese white pear, 18,677 pairs in apple and peach, and 15,331 pairs in peach and chinese white pear. There were 12 pairs of *CCoAOMT* genes with collinearity, eight pairs were in apple and chinese white pear, two pairs were in apple and peach, and two pairs were in chinese white pear and peach (Table 3, Fig. 4). Six *CCoAOMT* genes in chinese white pear had collinearity with *CCoAOMT* genes in other species: *Pbr015171.1*, *Pbr019305.1*, *Pbr025246.1*, *Pbr015180.1*, *Pbr028166.1*, *Pbr028157.1*. These six genes were distributed on Chr5, Chr9, Chr16, and scaffold466.0. Chinese white pear genes *Pbr015171.1*, *Pbr019305.1*, *Pbr025246.1*, *Pbr028157.1*, and *Pbr028157.1* had collinearity with a *CCoAOMT* gene in apple. *Pbr015180.1* and *Pbr028166.1* both had collinearity with two *CCoAOMT* genes in apple and one *CCoAOMT* gene in peach. Five *CCoAOMT* genes in apple had collinearity with *CCoAOMT* genes in other species: *MD02G1230800*, *MD05G1209400*, *MD13G1117900*, *MD16G1118200*, and *MD16G1119200*. These five genes were distributed on Chr02, Chr05, Chr13, and Chr16. Only one *CCoAOMT* gene *Prupe.1G227100.2* in peach had collinearity with *CCoAOMT* genes in other species;

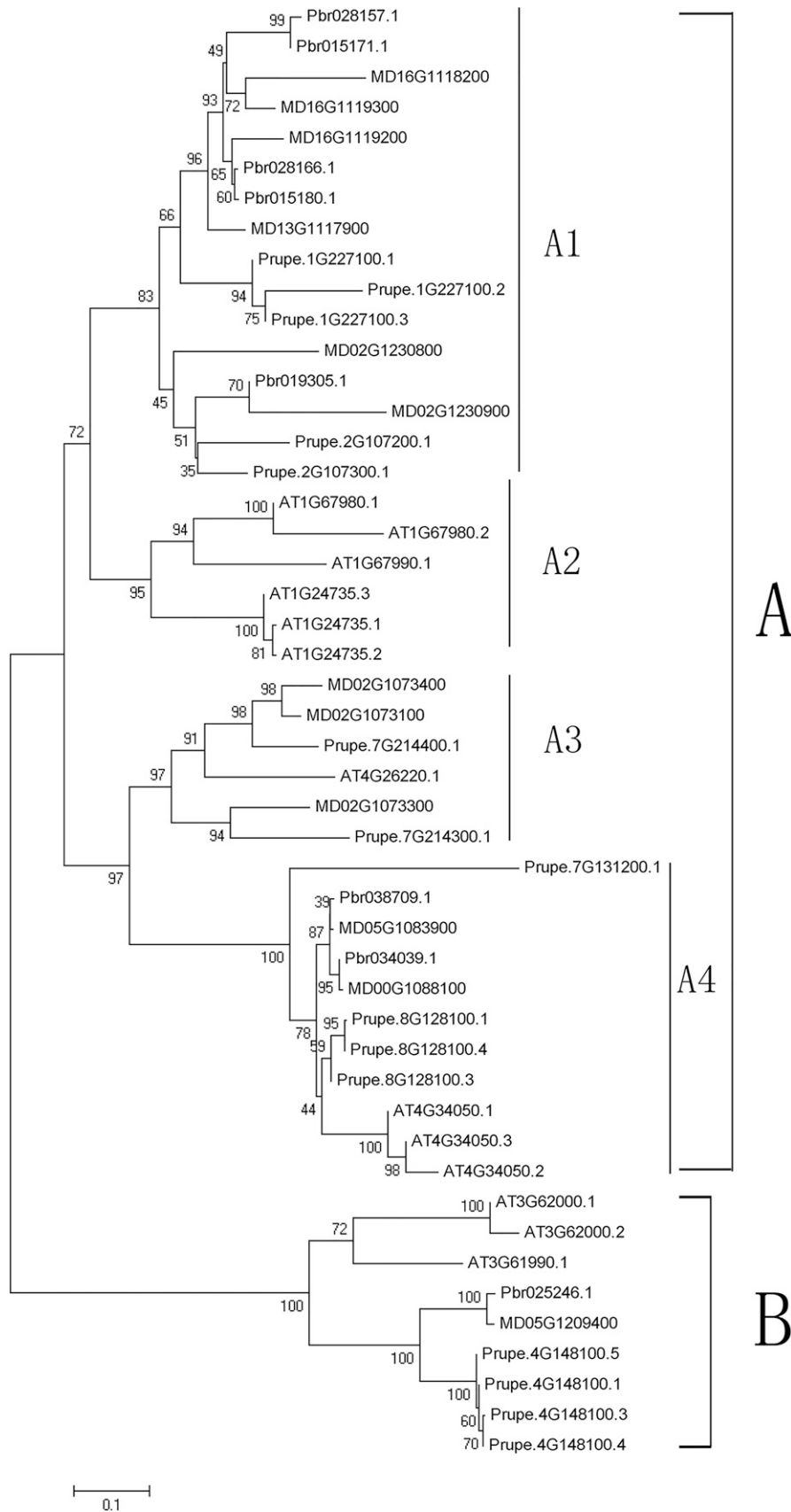


Fig. 1. Phylogenetic tree of *caffeoyl-CoA 3-O-methyltransferase (CCoAOMT)* gene family in apple, Chinese white pear, peach, and Arabidopsis. Neighbor joining method of MEGA 5.0 (Tamura et al., 2011) was used to construct the phylogenetic tree with a bootstrap value of 1000. A (A1, A2, A3, A4) and B are different groups of *CCoAOMT* genes.

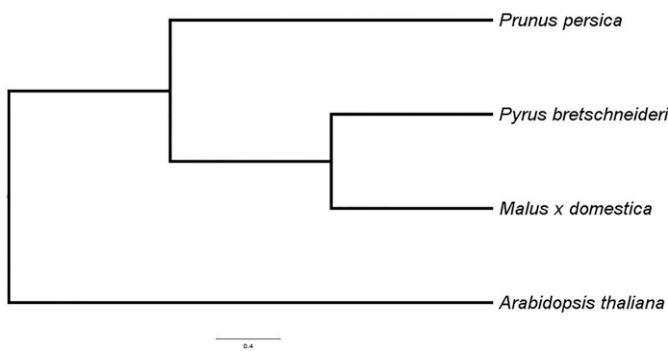


Fig. 2. Species tree of apple, chinese white pear, peach, and arabidopsis.

this gene was located in Chr01. This gene had collinearity with two *CCoAOMT* genes in apple and two *CCoAOMT* genes in chinese white pear.

**EXPRESSION OF *CCoAOMT* GENES IN APPLE, CHINESE WHITE PEAR, AND PEACH.** To explore the role of *CCoAOMT* genes in the growth and development of fruit, we examined transcriptome data in chinese white pear, apple, and peach (Figs. 5 and 6). Five genes were expressed differently in different growth and development stages of chinese white pear fruit. *Pbr028166.1* was expressed only at 15 DAF (Table 4). *Pbr015180.1* was expressed at 15 and 80 DAF, and the expression level at 80 DAF was slightly higher than that at 15 d. *Pbr019305.1*, *Pbr025246.1*, and *Pbr034039.1* expressed in six periods. Both *Pbr019305.1* and *Pbr034039.1* had a relatively high expression level at 15 DAF, and then the expression level of *Pbr019305.1* was reduced, whereas the expression level of *Pbr034039.1* decreased sharply after the peak at 36 DAF. These two genes were expressed

relatively highly at young fruit stage and early expansion stage of chinese white pear fruit (15, 36, 80, and 110 DAF), suggesting that the two genes mainly played a role at young fruit stage to expansion stage. The expression level of *Pbr025246.1* was stable during these six stages, with a little higher expression level at 110 and 167 DAF.

In apple fruit, four apple *CCoAOMT* genes were expressed in the transcriptome data (Table 5) with their expression differing at different growth and development stages. *MD02G1230800* was only expressed at the young fruit stage of wild apple (*Malus* sp.) fruit. *MD05G1083900* (*MD00G1088100*) and *MD02G1073400* were expressed in all growth and development stages of apple fruit in both wild and cultivated types. The expression level of *MD05G1083900* in wild apple was slightly higher than in cultivated apple. Their overall expression levels were lower in cultivated apple and decreased rapidly from young fruit stage to expansion stage. In contrast, their expression levels first increased and then decreased in wild apple. Generally, *CCoAOMT* genes were expressed at a higher level in wild than in cultivated lines.

The growth and development process of peach fruit can be divided into three stages, and at the second stage, the pericarp is lignified (Chalmers and van den Ende, 1975; Lilien-Kipnis and Lavee, 1971). In peach fruit, the expression levels of *Prupe.1G227100.1*, *Prupe.8G128100.1*, *Prupe.2G107300.1*, and *Prupe.7G214400.1* of six stages (41, 54, 69, 83, 111, and 125 DAF) were used (Table 6). It showed the expression levels of all these four genes showed a trend of first rising and then falling, with reaching the highest level in the middle period. This was consistent with the changes of endocarp lignification during the second stage. The expression level of *Prupe.1G227100.1* was always low. By contrast, the expression levels of the other three genes were relatively high, which indicated that these three genes played

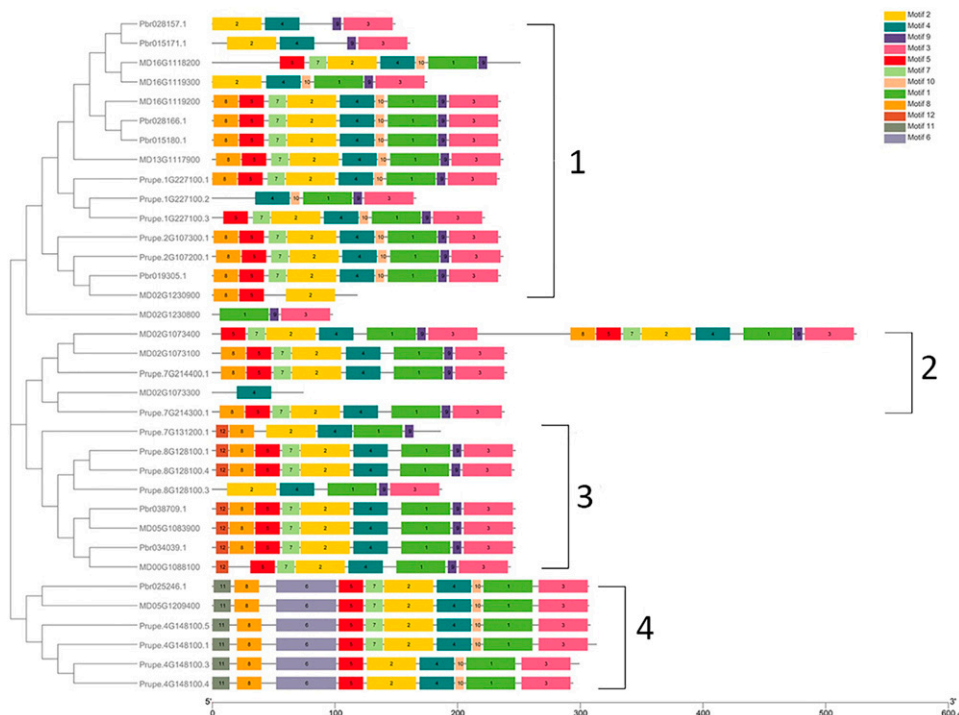


Fig. 3. Motif analysis of *caffeoyl-CoA 3-O-methyltransferase (CCoAOMT)* gene family in apple, chinese white pear, and peach. Motif analysis was performed using the Web site MEME (Bailey and Elkan, 1994). Motifs are shown in different colors.



Table 2. Positive selection analysis of *caffeoyl-CoA 3-O-methyltransferase (CCoAOMT)* genes.

Block	Sequence no.	dN/dS ( $\omega$ ) under M0 <sup>z</sup>	2 $\Delta$ l <sup>y</sup> M3 vs. M0 <sup>x</sup>	2 $\Delta$ l M8 vs. M7 <sup>x</sup>	M8 estimates <sup>w</sup>
A1	16	0.257	10.843**	3.400*10 <sup>-4</sup>	$P = 0.717$ $q = 1.117$ $PI = 0.000$ $\omega = 1.000$
A2	6	0.266	18.687	0.014	$P = 0.532$ $q = 1.1783$ $PI = 0.000$ $\omega = 1.000$
A3	6	0.218	37.058**	0.391	$P = 0.515$ $q = 2.783$ $PI = 0.131$ $\omega = 1.092$
A4	11	0.081	13.430	0.000	$P = 0.655$ $q = 6.328$ $PI = 0.000$ $\omega = 1.000$
B	9	0.275	81.402**	0.238	$P = 0.520$ $q = 3.506$ $PI = 0.279$ $\omega = 1.000$

<sup>z</sup>dS = synonymous substitution rates, dN = nonsynonymous substitution rates.

<sup>y</sup>2 $\Delta$ l: = twice of the log likelihood difference; M0 (Goldman and Yang, 1994; Yang and Nielsen, 1998); M3, M7, and M8 (Yang, 2000a, 2000b); different models in PAML.

<sup>x</sup>Likelihood ratio test (LRT) value between M3 and M0 and LRT value between M7 and M8. If the  $\chi^2$  test result is <0.05, the data are marked with a “\*”. If the  $\chi^2$  test result is <0.01, the data are marked with “\*\*”.

<sup>w</sup>PI = percentage of positively selected sites; p and q are distribution parameters. The genes were divided into different blocks based on motif analysis.

a major role in period of fruit growth and development. And the expression level of *Prupe.8G128100.1* is the highest among these four genes.

## Discussion

Previous studies have found that *CCoAOMT* genes in different species had a certain degree of conservation. These genes in rice, tobacco, grape (*Vitis vinifera*), maize, and some other species all have unique features characteristic of this gene family.

Also, the *CCoAOMT* genes have homology in different species. One *CCoAOMT* gene in cotton has high similarity with *CCoAOMT* genes in poplar and tobacco (Ni et al., 2010). The *CCoAOMT* genes in rice are highly similar to those in maize and *Bambusa oldhamii* (Ni et al., 2010; Zhao et al., 2004). The *CCoAOMT* genes in sorghum (*Sorghum bicolor*) have high similarity with those in maize and rice (Rakoczy et al., 2018); but there is a certain degree of difference between these similarities, and these differences may be related to the specific functions of individual *CCoAOMT* genes.

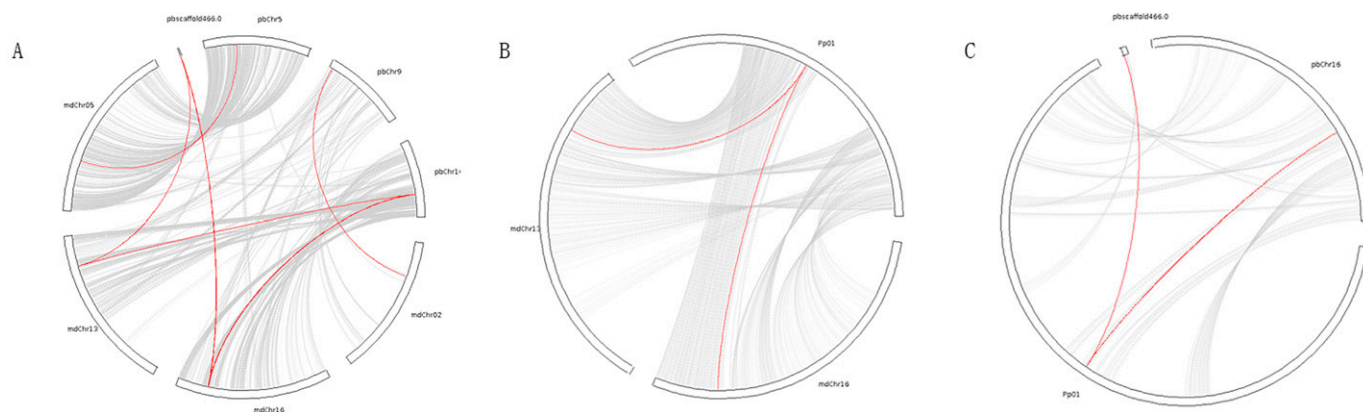


Fig. 4. Collinearity relationship of *caffeoyl-CoA 3-O-methyltransferase (CCoAOMT)* gene family in apple, chinese white pear, and peach. (A) Collinear genes on Chr5, Chr9, Chr16, and scaffold466.0 of pear with Chr02, Chr05, Chr13, and Chr16 of apple. (B) Collinear genes on Chr13 and Chr16 of apple with Chr01 of peach. (C) Collinear genes on Chr16 and scaffold 466.0 of pear with Chr01 of peach. *CCoAOMT* genes are indicated in red.

Table 3. Collinearity relationship of *caffeoyl-CoA 3-O-methyltransferase (CCoAOMT)* genes in apple, chinese white pear, and peach.

Gene	Chromosome	Gene	Chromosome
<i>Pbr019305.1</i> <sup>z</sup>	Chr9	<i>MD02G1230800</i>	Chr02
<i>Pbr025246.1</i>	Chr5	<i>MD05G1209400</i>	Chr05
<i>Pbr015180.1</i>	Chr16	<i>MD13G1117900</i>	Chr13
<i>Pbr028166.1</i>	scaffold466.0	<i>MD13G1117900</i>	Chr13
<i>Pbr015180.1</i>	Chr16	<i>MD16G1118200</i>	Chr16
<i>Pbr028166.1</i>	scaffold466.0	<i>MD16G1118200</i>	Chr16
<i>Pbr028157.1</i>	scaffold466.0	<i>MD16G1119200</i>	Chr16
<i>Pbr015171.1</i>	Chr16	<i>MD16G1119200</i>	Chr16
<i>Pbr015180.1</i>	Chr16	<i>Prupe.1G227100.2</i>	Chr01
<i>Pbr028166.1</i>	scaffold466.0	<i>Prupe.1G227100.2</i>	Chr01
<i>MD13G1117900</i>	Chr13	<i>Prupe.1G227100.2</i>	Chr01
<i>MD16G1118200</i>	Chr16	<i>Prupe.1G227100.2</i>	Chr01

<sup>z</sup>Genes in the same row have a collinearity relationship.

In our study, we also found some characteristics of the conservation and homology of *CCoAOMT* genes in three Rosaceae species. The physical and chemical properties of 35 *CCoAOMT* proteins were investigated in our study. By predicting the motifs in the sequence, it can be found that there were some similarities in the sequence structure. According to the perverse study of 2013 (Wu et al., 2013), there are nine *CCoAOMT* genes in chinese white pear and 18 in apple. The difference from our study was caused by the apple genome version, which was the 2017 version we used, and the different criteria.

Studies have shown that Rosaceae originated around the boundary between the early and late cretaceous and whole-genome duplications (WGD) occurred during the evolution of Rosaceae (Xiang et al., 2017). At least a single WGD is shared by both pear and apple (Li et al., 2019), whereas the peach has not undergone recent whole-genome duplication (Verde et al., 2013). From the phylogenetic analysis, we can find that *CCoAOMT* genes were duplicated. *MD02G1073400* had the most motifs, of which the combination of motif 5-motif 7-motif 2-motif 4-motif 1-motif 9-motif 3 was duplicated. It was probably that this sequence had gone through tandem duplication. Motifs 1 (DFIFVDADKDNY), 2 (KLINAKNTMEIGVYTGYSLLATA), 3 (GDGITLCRR), 4 (PVIQKAGVAHKIEF), and 5 (TSVYPREPEPMKELRELT) separately contained a unique tag sequence of

plant *CCoAOMT* (Joshi and Chiang, 1998), and most of the 35 sequences contained these five motifs.

The phylogenetic tree of *CCoAOMT* genes was divided into two main groups, A and B. Both groups contained *CCoAOMT* genes in apple, chinese white pear, peach, and arabidopsis, suggesting that the formation of the gene family precedes the differentiation of Rosales and Capparidales. The A group was further divided into A1, A2, A3, and A4 subgroups. The A1 subgroup only contained the genes in apple, chinese white pear, and peach, whereas the A2 subgroup only contained the *CCoAOMT* genes from arabidopsis. Also we indicated that the duplicated copies (*Pbr015180.1*–*Pbr028166.1*) of the chinese white pear genome in A1 are paralogous genes. And *Prupe.1G227100.1*, *Prupe.1G227100.2*, and *Prupe.1G227100.3* are splice variants of one gene, which are in one branch of the A1 group. We speculated that the differentiation of A1 and A2 occurred after the species divergence and that A1 and A2 are orthologous genes. The A3 subgroup lacked the *CCoAOMT* genes of chinese white pear, probably because genes in chinese white pear did not have these paralogous genes. The A4 subgroup contained two copies of apple *CCoAOMT* genes, two copies of chinese white pear genes, one copy of peach gene (considering splice variants), and one copy of arabidopsis (considering splice variants), which is as expected with the two duplicated genomes of apple and chinese white pear. The B group contained one copy of apple *CCoAOMT* gene, one copy of chinese white pear gene, one copy of peach gene (considering splice variants), and two copies of the arabidopsis gene.

Raes et al. (2003) identified seven *CCoAOMT* genes in arabidopsis. They divided plant *CCoAOMT* genes into two classes: class I contains one arabidopsis *CCoAOMT-1* (*AT4G34050*) gene with *CCoAOMT* genes from other plants, whereas class II consists of six arabidopsis *CCoAOMT* genes and a few sequences from other species. The study indicated that *CCoAOMT-5* (*AT1G67990*) and *CCoAOMT-6* (*AT1G67980*) originated through tandem duplication. In our study, *AT1G24735*, *AT1G67990*, and *AT1G67980* are located on a separate branch of phylogenetic tree, which are in the A2 group. Also, we found tandem duplications in apple, *MD02G1073100*–*MD02G1073300*–*MD02G1073400* in A3 group, *MD02G1230800*–*MD02G1230900* in A1 group, *MD16G1118200*–*MD16G1119200*–*MD16G1119300* on another branch of the A1 group. However, there is not enough information on tandem duplication of chinese white pear.

Gene collinearity shows evolutionary conservation of gene sequences on chromosomes. Gene collinearity analysis is the

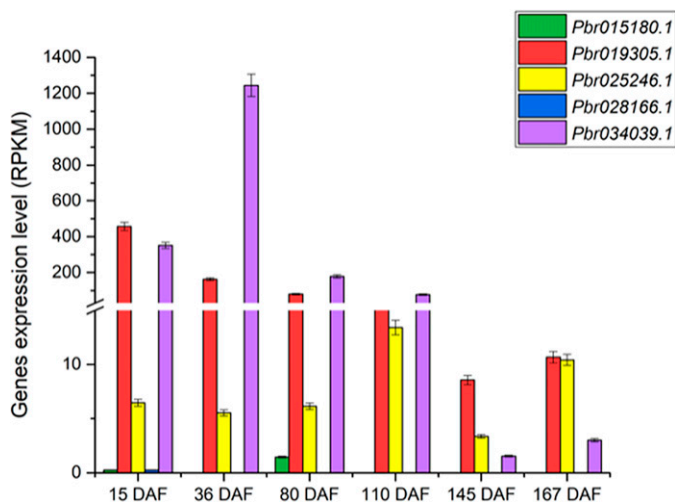


Fig. 5. Expression of *caffeoyl-CoA 3-O-methyltransferase (CCoAOMT)* genes in chinese white pear fruit at six different stages. DAF = days after flowering, RPKM = reads per kilobase million.

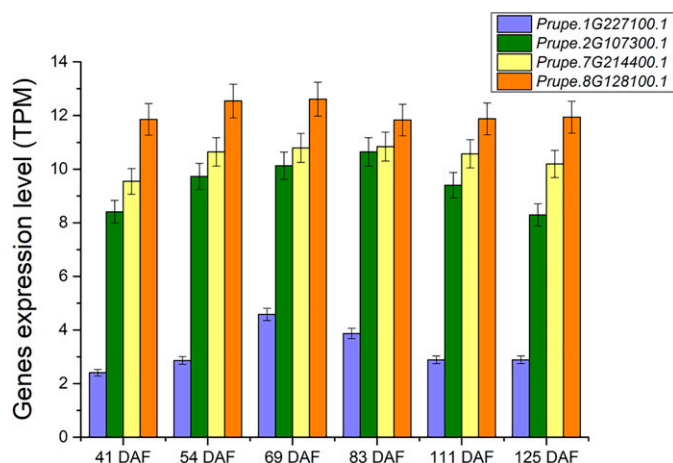


Fig. 6. Expression of *caffeoyl-CoA 3-O-methyltransferase (CCoAOMT)* genes in peach fruit at six different stages. DAF = days after flowering, TPM = transcripts per million.

basis of comparative genomics, and it can help us to understand the similarities and differences between species and their genetic relationship. It was found that the total of collinear genes between apple and chinese white pear was the most, compared with apple and peach or peach and chinese white pear. This result suggested that pear and apple had a close evolutionary distance.

In this study, PAML was used to perform positive selection analysis. It was found that all four groups had significant differences in selection pressure, but no reliable positive selection sites were found. We could not determine whether the positive selection effect plays a significant role in the evolution of *CCoAOMT* genes.

The *CCoAOMT* genes had different expression patterns during fruit growth and development. The apple (*MD05G1083900*) and chinese white pear (*Pbr034039.1*) *CCoAOMT* genes were highly expressed in fruit and decreasing in mature fruit and also the highest expressed in peach (*Prupe.8G128100.1*) is in the same cluster A4 as *CCoAOMT* gene in arabidopsis. Stone cells, the formation of which involve the accumulation of lignin and other substances, are important factors affecting the quality. The genes were divided into different blocks based on motif analysis of chinese white pear fruit. Some studies found that after the flowering of pears, the number of stone cells in pear fruit first increased and then decreased during the growth of fruit. In week 7, the highest peak was reached, and the number of stone cells fell to the lowest when the fruit matured (Liu et al., 2006). Also the study showed that encoding protein of *DiCCoAOMT1*

Table 4. Expression level of *caffeoyl-CoA 3-O-methyltransferase (CCoAOMT)* genes in different fruit stages of chinese white pear.

Gene	Days after flowering					
	15	36	80	110	145	167
<i>Pbr015180.1</i>	0.26	— <sup>z</sup>	1.45	—	—	—
<i>Pbr019305.1</i>	456.53	162.38	79.95	21.65	8.55	10.68
<i>Pbr025246.1</i>	6.45	5.52	6.13	13.42	3.36	10.42
<i>Pbr028166.1</i>	0.26	—	—	—	—	—
<i>Pbr034039.1</i>	351.19	1243.92	178.45	76.86	1.54	3.00

<sup>z</sup> – indicates no data.

demonstrated a relatively high O-methyltransferase activity when using caffeic acid as a substrate in vitro (Wu et al., 2019). We found that the expression trend of *Pbr034039.1* was consistent with the findings of the preceding studies, that is, the gene reached a peak at 36 DAF. This finding suggested that *Pbr034039.1* was likely to be related to the formation of stone cells.

Cao et al. (2019) identified two true *CCoAOMT* genes: *PbCCoAOMT1 (Pbr034039.1)* and *PbCCoAOMT2 (Pbr038712.1)* in chinese white pear. In their study, *PbCCoAOMT2 (Pbr038712.1)* showed a 6-fold higher expression level compared with *PbCCoAOMT1 (Pbr034039.1)* in chinese white pear fruit. Therefore, they inferred that *PbCCoAOMT2 (Pbr038712.1)* is the most likely candidate involved in lignin biosynthesis of chinese white pear fruit. We found only *Pbr034039.1* expressed in our study, which was likely to be related to the formation of stone cells. We can find that the expression of *Pbr034039.1* gradually decreases as the fruit develops, which is of the same tendency of *Pbr034039.1* in the findings of Cao et al. (2019).

The *CCoAOMT* genes of apple *MD05G1083900* and *MD00G1088100*, which are in the same subgroup of phylogenetic tree with *Pbr034039.1* and *Pbr038712.1*, showed a downward trend during the ripening process of apple cultivars. This may infer the fact that mature cultivar apple fruit do not have a large amount of lignin content. One study has found that the lignin content was lower in the core and flesh of ‘Fuji’ apple than that in other fruits by comparing the lignin content at the time of optimal harvest in 16 cultivars including japanese pear (*Pyrus* spp.), chinese white pear, european pear (*Pyrus communis*) cultivars, quince (*Cydonia oblonga*), chinese quince (*Carica papaya*), and apple (Zhang et al., 2020). In contrast, *MD05G1083900 (MD00G1088100)* showed high expression levels in the late ripening stage of wild apple fruit, which may imply that there is a difference in lignin content between wild apple fruit and cultivated apple fruit. And because

Table 5. Expression level of *caffeoyl-CoA 3-O-methyltransferase (CCoAOMT)* genes in different fruit stages of apple.

Gene	Fruit stage <sup>z</sup>					
	Cultivated T1	Wild T1	Cultivated T2	Wild T2	Cultivated T3	Wild T3
	Gene expression (transcripts per million)					
<i>MD02G1073400</i>	20.80	23.20	23.80	21.00	24.50	31.70
<i>MD02G1230800</i>	— <sup>y</sup>	0.75	—	—	—	—
<i>MD05G1083900 (MD00G108810)</i>	9.99	16.20	1.95	3.63	1.29	15.30

<sup>z</sup>Cultivated = ‘Golden Delicious’, Wild = Xifuhaitang, T1 = young fruit stages, T2 = enlargement fruit stages, T3 = mature fruit stages.

<sup>y</sup> – indicates no data.

Table 6. Expression level of *caffeoyl-CoA 3-O-methyltransferase (CCoAOMT)* genes in different fruit stages of peach.

Gene	Days after flowering					
	41	54	69	83	111	125
<i>Prupe.1G227100.1</i>	2.41	2.86	4.58	3.87	2.89	2.89
<i>Prupe.2G107300.1</i>	8.41	9.73	10.13	10.65	9.40	8.30
<i>Prupe.7G214400.1</i>	9.55	10.64	10.79	10.84	10.57	10.20
<i>Prupe.8G128100.1</i>	11.86	12.54	12.61	11.83	11.88	11.94

<sup>z</sup>The data are the normalization data of raw data [GEO accession GSE71561 (Zaffalon et al., 2017)].

lignin has a certain resistance to stress, it may also indicate that wild apples and cultivated apples have certain disease resistance differences.

For peach fruit, the data we used in the expression analysis are collected from the mesocarp of peach fruit, and we can find that although the expression level of each *CCoAOMT* genes in peach fruit change during fruit growth and development, the changes are not particularly sharp. Moreover, previous study has found that the expression level of *CCoAOMT* genes in the endocarp was significantly higher than that in the mesocarp and exocarp, which indicates that the *CCoAOMT* genes in peaches were largely endocarp specific (Dardick et al., 2010). Therefore, our result may indicate this feature. A similar lignification process was also found in the fruit of wild roses (*Rosa multiflora*) during the period of fruit growth and development through anatomic study. The fruit of these five species were same as peach fruit, which contained endocarp, mesocarp, and exocarp (Guzicka et al., 2012). In addition, the data are from microarrays, hence not all the genes are necessarily on the array and, depending on the oligos used, they may not discriminate between the splice variants. Rosaceae species also have different fruit types, such as fruits of some achenetum, of which the fruit are dried. In general, the relationship between the structure and the lignification changes of Rosaceae fruit deserves more related studies.

Apple and chinese white pear are duplicated genomes and peach is not; however, the *CCoAOMT* gene numbers are similar in these three fruits. This would indicate that the *CCoAOMT* gene was not in the tandem duplication area of the genome. Also, from our study we can tell that although the results can give us useful information of *CCoAOMT* genes, there can be misannotation of some of the genes. More accurate genome information is needed to conduct a following study.

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