Characterization and functional analysis of the Hydroxycinnamoyl-CoA: shikimate hydroxycinnamoyl transferase (HCT) gene family in poplar

Nan Chao 1,2,3, Qi Qi 2,4, Shuang Li 2, Brent Ruan 5, Xiang-Ning Jiang 2,6, Ying Gai Correspond. 2, 6

1 School of Life Science, Tsinghua University, Beijing, China
2 College of Biological Sciences and Biotechnology, Beijing Forestry University, Beijing, China
3 School of Biotechnology, Jiangsu University of Science and Technology, Zhenjiang, China
4 College of Horticulture, China Agricultural University, Beijing, China
5 Department of Agricultural and Biological Engineering, University of Illinois at Urbana Champaign, Urbana Champaign, USA
6 National Engineering Laboratory for Tree Breeding, the Tree and Ornamental Plant Breeding and Biotechnology Laboratory of Chinese Forestry Administration, Beijing, China

Corresponding Author: Ying Gai
Email address: gaiying@bjfu.edu.cn

Hydroxycinnamoyl-CoA: shikimate hydroxycinnamoyl transferase (HCT) divides the mass flux to H, G and S units in monolignol biosynthesis and affects lignin content. Ten HCT homologs were identified in the Populus trichocarpa (Torr. & Gray) genome. Both genome duplication and tandem duplication resulted in the expansion of HCT orthologs in Populus. Comprehensive analysis including motif analysis, phylogenetic analysis, expression profiles and co-expression analysis revealed the divergence and putative function of these candidate PoptrHCTs. PoptrHCT1 and 2 were identified as likely involved in lignin biosynthesis. PoptrHCT9 and 10- are likely to be involved in plant development and the response to cold stress. Similar functional divergence was also identified in Populus tomentosa Carr. Enzymatic assay of PtoHCT1 showed that PtoHCT1 was able to synthesize caffeoyl shikimate using caffeoyl-CoA and shikimic acid as substrates.
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Nan Chao¹,²,⁴, Qi Qi²,⁵, Li Shuang², Brent Ruan⁶, Xiang-Ning Jiang²,³ and Ying Gai²,³†

¹School of Life Science, Tsinghua University,
²College of Biological Sciences and Biotechnology, Beijing Forestry University,
³National Engineering Laboratory for Tree Breeding, the Tree and Ornamental Plant Breeding and Biotechnology Laboratory of Chinese Forestry Administration, Beijing 100083, China,
⁴School of Biotechnology, Jiangsu University of Science and Technology,
⁵College of Horticulture, China Agricultural University,
⁶Department of Agricultural and Biological Engineering, University of Illinois at Urbana Champaign

†To whom correspondence should be addressed: Tel (Fax): +86 10 6233 8063;
E-mail: gaiying@bjfu.edu.cn, Beijing Forestry University.

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Abstract: Hydroxycinnamoyl-CoA: shikimate hydroxycinnamoyl transferase (HCT) divides the mass flux to H, G and S units in monolignol biosynthesis and affects lignin content. Ten HCT homologs were identified in the *Populus trichocarpa* (Torr. & Gray) genome. Both genome duplication and tandem duplication resulted in the expansion of HCT orthologs in *Populus*. Comprehensive analysis including motif analysis, phylogenetic analysis, expression profiles and co-expression analysis revealed the divergence and putative function of these candidate *PoptrHCTs*. *PoptrHCT1* and 2 were identified as likely involved in lignin biosynthesis. *PoptrHCT9* and 10- are likely to be involved in plant development and the response to cold stress. Similar functional divergence was also identified in *Populus tomentosa* Carr. Enzymatic assay of *PtoHCT1* showed that *PtoHCT1* was able to synthesize caffeoyl shikimate using caffeoyl-CoA and shikimic acid as substrates.

Key words: Hydroxycinnamoyl- CoA: shikimate hydroxycinnamoyl transferase; enzymatic synthesis; divergence; gene family; monolignol; *Populus*

Introduction

Primary walls and secondary walls protect plant cells and define the shapes of cells, tissues, organs and ultimately the whole plant body (Zhong et al. 2019). Lignin is an important component for secondary cell walls and is one of the most abundant components of biomass in plants (Boerjan et al. 2003; Tang & Tang 2014). Therefore, lignin plays a vital role in plant physiology. Owing to the recalcitrant chemical nature and the complexity of lignin, lignin limits the conversion efficiency of lignocellulosic biomass to ethanol (Poovaiah et al. 2014; Vanholme et al. 2010). Modifying trees to have less lignin or more-degradable lignin along with normal growth, can reduce the high processing costs and carbon footprint of making paper, biofuels, and chemicals (Ralph et al. 2019; Tang & Tang 2014; Wang et al. 2019; Xu & Li 2016; Zhao 2016).

The biosynthetic pathway for lignin has been studied extensively and the phenylpropane pathway which begins with phenylalanine, is responsible for monolignol biosynthesis. (Boerjan et al. 2003; Karkonen & Koutaniemi 2010; Maeda 2016; Ralph et al. 2019; Vanholme et al. 2010; Wang et al. 2019; Xu & Li 2016). Monolignol is the general name for lignin building blocks. Our understanding of the monolignol biosynthetic pathway has continued to grow, and now 11 enzyme families and 24 metabolites are associated with it (Vanholme et al. 2019). Hydroxycinnamoyl- CoA: shikimate hydroxycinnamoyl transferase (HCT) is located at a key point in the monolignol biosynthetic pathway and is conserved across all land plants. In conjunction with
C3H (p-coumarate 3-hydroxylase), HCT catalyzes two steps to direct the mass flux from the H monolignol to G and S monolignols (Figure 1). HCT first catalyzes the coupling of p-coumaroyl-CoA with shikimate to produce p-coumaroyl shikimate (Hoffmann et al. 2004; Hoffmann et al. 2003). Caffeoyl shikimate is generated by C3H and is then transesterified by HCT to form caffeoyl-CoA. This reaction is probably reversible based on the reported in vitro activity (Lepelley et al. 2007; Wang et al. 2014). Caffeoyl shikimate esterase (CSE), a new member in monolignol biosynthesis pathway recently discovered in plants can hydrolyze caffeoyl shikimate to release caffeate (Ha et al. 2016; Saleme et al. 2017; Vanholme et al. 2013; Vargas et al. 2016). Although down-regulation of HCT expression improves forage digestibility and saccharification efficiency, it negatively affects plant growth resulting in shorter plants (Li et al. 2010; Shadle et al. 2007).

HCT (GO:0102660) belongs to the BAHD acyltransferase family and is able to utilize many non-native substrates. Some HCTs (also called HQT) can use quinate as a substrate in addition to shikimate (Eudes et al. 2016; Kim et al. 2013) for the biosynthesis of chlorogenic acid. As an acyl-CoA-dependent transferase, HCT is capable of acylating a wide variety of acceptors, with some exhibiting broad substrate flexibility (Chiang et al. 2018; Eudes et al. 2016). Crystal structures of HCTs from different plants have been determined for both the apo-form and complexed structure with diverse substrates allowing determination of active sites. For example, the apo-form and ternary complex with p-coumaroyl-CoA and shikimate of SbHCT from *Sorghum bicolor* (L.) Moench revealed the catalytic mechanism of HCT (Walker et al. 2013). Structures of AtHCT from *Arabidopsis thaliana* (L.), CbHCT from *Coleus blumei* Benth, CcHCT from *Coffea canephora* Pierre ex Froehn and SmHCT from *Selaginella moellendorffii* Hieron. have also been reported (Chiang et al. 2018; Lallemand et al. 2012; Levsh et al. 2016).

Similar to other key genes involved in monolignol biosynthesis, HCT is found as a gene family in many species in plant kingdom (Carocha et al. 2015; Ferreira et al. 2019; Ma et al. 2017; Raes et al. 2003; Zhang et al. 2018). The structural information and the proposed active sites of HCT, can help us to distinguish bona fide HCT utilizing shikimate as an acceptor and involved in monolignol biosynthesis in plants. In this study, we used genome-wide screening to identify 10 HCT homologs in *Populus trichocarpa*. Further motif and active site analysis showed the divergence of *PoptrHCTs*. Expression profiles and co-expression network analysis identified *PoptrHCT1* and 2 as the lignin-related HCTs. Finally, we cloned and characterized the catalytic
activity of PtoHCT1 from *Populus tomentosa* in vitro, which generated caffeoyl shikimate. PtoHCT1 could be used as the target gene for genetic modification to alter lignin content and composition.

### Materials and Methods

#### Materials

Leaves of six--year-old *Populus tomentosa* 741 were collected from Hebei, China (Hu et al. 2019; Tian et al. 2013). Samples were immediately frozen in liquid nitrogen and then stored at -80 °C until use.

#### Genome-wide identification of HCT gene family members

To identify the HCT sequences in *Populus*, we first built a hidden Markov model (HMM) using reported HCT and HQT sequences. HMMsearch using Hmmer 3.0 software against the proteome data of *Populus trichocarpa* was performed based on the HMM model (Eddy 2010). The cutoff for PoptrHCT homolog screening was an E-value (<E–100) of both the domain and full sequence and scores of full sequences (>400) (Supplementary Table S1). The stable gene ID and symbols for HCTs reported in a previous study were also marked in Supplementary Table S1 Table S2. The sequences used for building the HMM model are shown in Supplementary Table S2.

#### Distribution of HCT genes and HCT orthologs on *Populus* chromosomes

Ten candidate HCT and HCT orthologs were located on chromosomes in specific duplicated blocks which were determined based on the *Populus* genome and the WGDotplot in the PLAZA platform (Proost et al. 2009).

#### HCT sequence alignment and phylogenetic analysis

Alignment of PoptrHCTs and SbHCT, AtHCT were performed using DNAman 8.0 (Lynnon BioSoft) with default parameters. A phylogenetic tree was obtained using Mega 7.0 with the maximum-likelihood method (Kumar et al. 2016; Tamura et al. 2011). The phylogenetic tree was assessed by bootstrapping using 1000 bootstrap replicates and marked above nodes only if greater than 50. The JTT substitution model and G+I rates among sites model were selected as parameters for building the tree. The putative HCT sequences are listed in Supplementary Table S2.

#### HCT expression profiles in *P. trichocarpa* and *P. tomentosa*

We obtained gene expression profiles for various tissues in *P. trichocarpa* using the GEO database with the accession number GSE30507. In addition, RNA-seq dataset GSE78953 including the transcriptome of various
monolignol biosynthesis related mutants in *P. trichocarpa*, was used for co-expression analysis to explore the functions of the *PoptrHCT* orthologs. We also examined the expression profiles of *PtoHCT* orthologs in *P. tomentosa* in different seasons (Spring, Summer, Fall and Winter) and organs or tissues (roots, buds, phloem and xylem) using our microarray dataset (accession number: GSE56023) (Chao et al. 2014b). The corresponding *PtoHCTs* were identified using *PoptrHCTs* as queries by local blastn against the probe sequences database (Christiam et al. 2009) TBtools v0.6652 and Cytoscape 3.4 were used to visualize the HCT expression profile or co-expression network (Chen et al. 2018; Shannon et al. 2003) (Supplementary Table S3).

**Cloning and purification of recombinant HCT from *P. tomentosa***

Isolation of RNA and cDNA synthesis have been described in a previous study (Chao et al. 2014a). We cloned the homologous *HCT1* from *P. tomentosa* based on the sequence information from *P. trichocarpa* (GenBank accession number: KT021003). Primer pair used for PCR amplification of *PtoHCT1* is as follow: forward, 5’-CGATAAATAGAGCATTAGCACGGGG-3’; and reverse, 5’-ATAGCTCGGCTCATTCTTT-3’. PCR products were purified and cloned into the pMD18-T vector (Takara Dalian), propagated in *Escherichia coli* DH5α and inserts were confirmed by sequencing. *PtoHCT1* was constructed with pET28a (Novagen) through a digestion-ligation way using restriction enzymes BamHI, HindIII and T4 ligation (Takara, Dalian). pET28a-PtoHCT1 was then transformed into *E.coli* BL21(DE3). To induce expression, Isopropy-β-D-thiogalactoside (IPTG) was added to a final concentration of 0.8mM and incubation was continued at 28°C for four hours. Cells were collected by centrifugation at 4000g and 4°C for 15min. The pellets were resuspended in lysis buffer (50 mM NaH₂PO₄, 300 mM NaCl with 10 mM imidazole, pH 8.0) and then disrupted by sonication. After centrifugation at 12,000 g and 4°C for 30 min, the lysates were mixed with pretreated 1 ml Ni-NTA agarose (Qiagen Shanghai, China) After washing using lysis buffer supplemented with 20mM imidazole, the His-tagged *PtoHCT1* was eluted with 100mM imidazole in lysis buffer.

**Catalytic activity of recombinant PtoHCT1***

Caffeoyl-CoA was chemically synthesized as reported (Chao et al. 2017). We determined the activity of recombinant PtoHCT1 by synthesis of caffeoyl shikimate using caffeoyl-CoA and shikimic acid as substrates. The reaction was performed according to Cesarino *et al.* (2013) and Luis *et al.* (2014). Briefly, total 40μl standard reaction mix contained 100 mM Tris-HCl pH 7, 1mM DTT, 100 μM caffeoyl-CoA, 100μM shikimic acid and
10 μg purified recombinant HCT protein. The reaction was initiated by adding the HCT proteins or the same amount of boiled protein as negative control. After incubating at 30 °C for 30 min, the reaction was terminated by boiling the samples for 5 min. Flow for HPLC analysis was 0.1 mL/min in solvent A (acetonitrile) and solvent B (0.01% formic acid in water). The gradient was 0% A to 35% in B for 0 to 24 min, 35% A in B to 100% B for 24 to 27 min, 100% B for 27 to 32 min, 100% A to 100% B for 32 to 35 min, and 100% B for 35 to 55 min. The parameters used for MS analysis was sheath gas (nitrogen) flow rate, 40 arb; aux/sweep gas (nitrogen) flow rate, 10 arb; spray voltage, 4.5 kV; capillary temperature, 320 °C. Optimized detailed parameters for dissociation of parent ions into product ions for each compound were provided in Supplementary Table S4.

Structure modeling of PtoHCT1

The crystal structure of AtHCT (accession number 5KJT) (Levsh et al. 2016) was obtained from the Protein Data Bank to build a homolog model for PtoHCT (https://www.rcsb.org). Molecular docking was performed using CDOCKER assembled in Discovery Studio 4.5. Visualization of the active sites and 3-D structures were generated by Discovery Studio 4.5.

Results

Genome-wide identification and distribution of HCT orthologs in Populus

Ten PoptrHCT homologs were found based on HMMsearch against the Populus genome (Supplementary Table S1). These HCT candidate genes are located on six different chromosomes. Among the 10 PoptrHCT orthologs, PoptrHCT3, 4 and 5 were located on chromosome V, and PoptrHCT7, 8, 9, and 10, were on chromosome XVIII, representing two clusters respectively (Figure 2A). Tandem duplication is likely to be responsible for the formation of HCT homolog clusters. Ks (substitution per synonymous site) value distributions can be used for revealing whole genome duplication (WGD) events (Jiao et al. 2011; Tang et al. 2010). PoptrHCT1 and PoptrHCT2 formed a homolog duplicate pair with Ks value 0.2174 and were located at corresponding homologous duplicated blocks, as the result of whole genome duplication. The organization of the PoptrHCT orthologs indicates that both genome duplication and tandem duplication played roles in the formation of the HCT family.

Alignment and phylogenetic analysis of HCT orthologs

Putative protein sequences of PoptrHCT orthologs and crystal structures of two shikimate-specific HCTs
(AtHCT and SbHCT) and LeHQT (*Lycopersicon esculentum* Mill.) were aligned. Characteristic of the BADH superfamily, two motifs HXXXD(G) and DFGWG were conserved in AtHCT, SbHCT and all PoptrHCT orthologs (except PoptrHCT6) (D’Auria 2006). Based on previous studies of the structure of HCTs including site-directed mutagenesis, molecular docking and crystallographic analyses we summarized the active sites of HCTs (*Table 1*) and marked these active sites in Figure 3. Active sites for the carbonyl group of the *p*-coumaroyl moiety binding and the catalysis related sites of LeHQT correspond with HCTs (red full circles) while divergence is obvious in terms of active sites for shikimate binding (red full stars). PoptrHCT1 and PoptrHCT2 showed conservation at these active sites and kept correspondence with AtHCT and SbHCT, while PoptrHCT3-10 showed poor conservation at these key sites. Thus while the ten candidate PoptrHCTs mostly belong to the BADH superfamily, only PoptrHCT1 and PoptrHCT2 appear to be associated with monolignol biosynthesis. Phylogenetic analysis showed PoptrHCT1 and PoptrHCT2 grouped with Group I HCTs, which transfer hydroxycinnamates to shikimate and have been implicated in monolignol biosynthesis, strongly suggesting this role for PoptrHCT1 and 2 as well (Figure 2B). Other PoptrHCTs (3-10) clustered with HQTs and other HCT-like as Group II and especially, PoptrHCT6 without DFGWG seem unlikely to be shikimate-specific transferases involved in monolignol biosynthesis. Thus these Group II. PoptrHCTs might have different catalytic activity (e.g., utilize acceptors other than shikimic acid) and are likely to play different roles in plants.

**Expression analysis of HCT homolog genes**

Based on the microarray analysis of seven different tissues and organs in *P. trichocarpa*, PoptrHCT1 and 2 showed expression preference in developing xylem (DX) and mature xylem (MX). PoptrHCT1 showed high expression levels in all detected tissues and organs. PoptrHCT9 and 10 showed expression preference in developing tissues including developing phloem, developing xylem, cambium (C) and shoots and leaf primordium (SLp) (*Figure 4A*). Co-expression network analysis shows that PoptrHCT1 and 2 have significant correlations with genes involved in lignin biosynthesis (*Figure 4B*). Monolignol biosynthesis related transcription factors also showed co-expression with PoptrHCT1 and 2. The similar expression patterns were also found in *P. tomentosa* Carr. (*Pto*). PtoHCT1 has high expression levels in almost all tissues and organs year-round while PtoHCT9 and 10 showed preference in buds and phloem especially in winter, which indicates that these two genes could be involved in development of dormancy and response to cold stress. The differential
expression of the *PoptrHCT* orthologs further supports that *HCT1* plays a major role in monolignol biosynthesis.

**Catalytic activity and structure comparison of PtoHCT1**

PtoHCT1 protein expressed in *E. coli* was purified for enzymatic assays. We monitored PtoHCT1 reactions using HPLC-MS and found that PtoHCT1 can utilize caffeoyl-CoA and shikimic acid (Figure 5). After the initiation of the reaction by adding PtoHCT1, the accumulation of caffeoyl shikimate and decrease of caffeoyl-CoA and shikimic acid were observed within 2 minutes. We built a homology model for PtoHCT1 to explore the structure of PtoHCT1 using the crystal structure of AtHCT (accession number 5KJT) as template. The main-chain root-mean-square deviation (RMSD) is 0.224 Å indicating the high structure similarity of PtoHCT1 and AtHCT (Figure 6A). According to the summarized active sites (Table 1), we found these conserved sites around the catalytic cleft, and then we successfully docked PtoHCT1 with the substrate caffeoyl-CoA, which further provided the structural evidence for PtoHCT1 catalyzing caffeoyl-CoA (Figure 6B).

**Discussion**

HCT regulates the flux at a key point in monolignol biosynthesis and has been studied in many plants (Hoffmann et al. 2004; Hoffmann et al. 2003; Shadle et al. 2007; Sun et al. 2018; Wagner et al. 2007). HCT also exists as a gene family in the land plant kingdom similar to other key genes involved in monolignol biosynthesis. We focused on the HCT gene family in this study and provided a systematic analysis of the HCT genes in poplar, and identified two lignin-related HCTs (*HCT1* and *HCT2*), which exist as a homolog pair located at a duplication block on chromosome I and III respectively (Figure 2A). Ks analysis indicated that the HCT gene pair (*PoptrHCT1* and *PoptrHCT2*) resulted from a recent genome duplication (Ks value 0.2174). Two HCT homolog clusters resulting from tandem duplication were also identified. Both genome duplication and tandem duplication provide raw genetic material for neo-function as a driving force in plant evolution and are responsible for the expansion of HCT orthologs (Zhang 2003).

Based on our systematic analysis, *PoptrHCT1* and 2 are involved in monolignol biosynthesis. Both *PoptrHCT1* and 2 showed expression preference in xylem and co-expression with other monolignol related genes. *HCT1* (*PoptrHCT1* and *PtoHCT1*) had high expression levels in different tissues. PtoHCT1 showed catalytic activity for caffeoyl-CoA and shikimic acid. These results further validate *HCT1* as the dominant HCT in monolignol biosynthesis, while *HCT9* and 10 (*PoptrHCT9*, 10 and *PtoHCT9*, 10) showed expression
preference in developing tissues. *PtoHCT9* and *10* were active in winter, which suggested a function in plant
development and response to cold stress. *PtoHCT1* utilizes caffeoyl-CoA and shikimic acid to generate caffeoyl
shikimate which can be used as substrate for CSE, a new annotated enzyme involved in monolignol biosynthesis
(Ha et al. 2016; Saleme et al. 2017; Vanholme et al. 2013).

**Conclusion**

In summary, we identified ten HCT homologs and proposed the important roles of both genome duplication
and tandem duplication in the expansion of HCT orthologs in *Populous*. Two HCTs likely involved in
monolignol biosynthesis in *Populus* were identified based on phylogenetic analysis and expression profile
analysis. Enzymatic assay of *PtoHCT1* showed that *PtoHCT1* was able to synthetize caffeoyl shikimate using
caffeoyl-CoA and shikimic acid as substrates. In addition, other *PoptrHCT* orthologs showed divergence in
reported active sites and different expression pattern. *HCT9* and *10* (*PoptrHCT9, 10* and *PtoHCT9, 10*) showed
preferential expression in developing tissues and were active in winter. Further studies should help to reveal the
functions of the other HCT orthologs.

Supplementary Table S1 Genome-wide screening *PoptrHCT* genes based on HMM models;
Supplementary Table S2 Putative HCTs used for phylogenetic analysis;
Supplementary Table S3 Expression profiles and co-expression for HCT homologs.
Supplementary Table S4 Conditions used for HPLC-MS to identify caffeic acid, shikimate and caffeoyl
shikimate
Supplementary File 5: Sequences used for alignment analysis.

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**Author Contributions**

Ying Gai and Xiang-Ning Jiang guided the work and provided advice; Qi Qi, Shuang Li and Nan Chao performed
the experiments; Brent Ruan and Nan Chao read and revised the draft; Nan Chao analyzed the data and wrote
the paper.

**Conflict of Interest**
The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Figure 1

Schematic diagram of reaction catalyzed by HCT in monolignol biosynthesis pathway

R=Shikimate. 4CL, 4-coumarate-CoA ligase; C3H, p-coumarate 3-hydroxylase; HCT, Hydroxycinnamoyl-CoA: shikimate hydroxycinnamoyl transferase. Compounds in red shadow are precursors for H units and in green shadow are for G and S units.
**Figure 2**

*PoptrHCT orthologs organization and phylogenetic analysis*

(A). Organization of HCT orthologs on *Populus* chromosomes. Regions that are assumed to correspond to homologous genome blocks are shaded gray and connected by lines. The position of genes is indicated with an arrowhead. (B). Phylogenetic analysis of HCT homologs from *Populus trichocarpa* and other plant species. The PoptrHCT1 and 2 were marked with full black triangle. Two groups for HCT orthologs were shown and HCTs in Group I are likely to transfer hydroxycinnamates to shikimate and have been implicated in monolignol biosynthesis. The scale bar indicates 0.5 amino acid substitutions per site in given length. The accession numbers of sequences used are as followed: *Arabidopsis thaliana* AtHCT (AT5G48930), AtHCT-like (AT4G29250); *Amborella trichopoda* AtrHCT1 (ATR_00137G00320), AtrHCT2 (ATR_00727G00010); *Cynara cardunculus* CcaHCT (DQ104740), CcaHQT (ABK79690); *Lycopersicon esculentum* LeHQT (AJ582652); *Larix kaempferi* LkaHCT (AHA44839); *Nicotiana tabacum* NtaHCT (Q8GSM7), NtaHQT (CAE46932); *Picea lauca* PglHCT (CZ001061061); *Populus trichocarpa* PoptrHCT1 (PT01G04290), PoptrHCT2 (PT03G18390), PoptrHCT3 (PT05G02800), PoptrHCT4 (PT05G02810), PoptrHCT5 (PT05G02840), PoptrHCT6 (PT18G03270), PoptrHCT7 (PT18G10470), PoptrHCT8 (PT18G10480), PoptrHCT9 (PT18G10540), PoptrHCT10 (PT18G10550); *Physcomitrella patens* PpHCT1 (PP00022G00830); *Panicum virgatum* PviHCT1a (JX845714), PviHCT2 (KC696573); *Sorghum bicolor* SbHCT (XP_002452435.1).
Figure 3

Alignment of PoptrHCT and PoptrHCT orthologs compared to shikimate-specific HCTs from Arabidopsis and sorghum and HQT from tomato.

Red full stars indicate shikimate binding sites, red full circles indicate carbonyl group of p-coumaroyl moiety binding sites, purple full triangle indicate carbonyl group of shikimate moiety binding sites and the blue full circles indicate sites involved in catalysis. Accessions are as in Figure 2. Detailed references are also available in Table 1.
Figure 4

Expression profile and co-expression network of HCT orthologs in poplar.

(A). Expression profile of HCT orthologs in Populus. Tissues or specific parts of plants are indicated with the respective abbreviations: WS, whole stems; BM, Bark and mature phloem; C, cambium; DP, developing phloem; DX, developing xylem; ML, mature leaf; SLp, shoot and leaf primordium. (B) Expression profile of HCT orthologs in P. tomentosa. (C) Co-expression network of PoptrHCT orthologs with identified genes involved in lignin biosynthesis. The support information is available in Table S3. Only nodes with Pearson correlation coefficients >0.9 were shown and considered as close co-expression.
Figure 5

PtoHCT1 catalyzes enzymatic synthesis of caffeoyl shikimate.

LC separation of reactions with MS detection (selected ion signals) (A) at initiation of the reaction (B) after 80s or (C) after 120s
Figure A: Chromatogram showing the separation of Shikimate (red) and Caffeoyl-Shikimate (blue) with retention times of 32.47 minutes. The inset shows the molecular ion peaks at 111, 146, 172, and 176.

Figure B: Chromatogram with a prominent peak at 30.33 minutes for Caffeoyl-Shikimate, accompanied by a smaller peak at 119 minutes. The inset displays molecular ion peaks at 119, 135, 161, and 179.

Figure C: The chromatogram highlights a peak at 30.33 minutes for Caffeoyl-Shikimate, with another peak at 32.47 minutes. The inset includes molecular ion peaks at 119, 161, and 179.
Figure 6

The structure of PtoHCT1 and docking with caffeoyl-CoA

(A) structure alignment of AtHCT (green) and PtoHCT (purple). (B) PtoHCT docked with caffeoyl-CoA. Blue ligand is caffeoyl-CoA and active sites are labeled.
Table 1 (on next page)

Summary of active sites of HCTs
Table 1 Summary of active sites of HCTs

| Position | Amino acid | Annotation                          | Reference                      |
|----------|------------|-------------------------------------|--------------------------------|
| 31       | Val        | Shikimate binding                   | Walker et al. 2013             |
| 32       | Pro        | Shikimate binding                   | Walker et al. 2013             |
| 298      | Ala        | Shikimate binding                   | Walker et al. 2013             |
| 318      | Ile        | Shikimate binding                   | Walker et al. 2013             |
| 376      | Phe        | Shikimate binding                   | Walker et al. 2013             |
| 414      | Leu        | Shikimate binding                   | Walker et al. 2013, Lallemand et al. 2012 |
| 418      | Leu        | Shikimate binding                   | Walker et al. 2013             |
| 38       | Ser        | carbonyl group of p- coumaroyl moiety | Walker et al. 2013, Eudes et al. 2016 |
| 40       | Tyr        | carbonyl group of p- coumaroyl moiety | Walker et al. 2013, Lallemand et al. 2012, Eudes et al. 2016 |
| 384      | Trp        | carbonyl group of p- coumaroyl moiety | Walker et al. 2013             |
| 163      | His        | carbonyl group of shikimate moiety and Catalysis | Walker et al. 2013, Lallemand et al. 2012, Eudes et al. 2016 |
| 369      | Arg        | carbonyl group of shikimate moiety  | Walker et al. 2013             |
| 382      | Thr        | carbonyl group of shikimate moiety  | Walker et al. 2013, Eudes et al. 2016 |
| 36       | Thr        | Catalysis                           | Walker et al. 2013             |

Note: all positions correspond to SbHCT
