

Identification of tomato F-box proteins functioning in phenylpropanoid metabolism

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Abstract

Phenylpropanoids, a class of specialized metabolites, play crucial roles in plant growth and stress adaptation and include diverse phenolic compounds such as flavonoids. Phenylalanine ammonia-lyase (PAL) and chalcone synthase (CHS) are essential enzymes functioning at the entry points of general phenylpropanoid biosynthesis and flavonoid biosynthesis, respectively. In Arabidopsis, PAL and CHS are turned over through ubiquitination-dependent proteasomal degradation. Specific kelch domain-containing F-Box (KFB) proteins as components of ubiquitin E3 ligase directly interact with PAL or CHS, leading to polyubiquitinated PAL and CHS, which in turn influences phenylpropanoid and flavonoid production. Although phenylpropanoids are vital for tomato nutritional value and stress responses, the post-translational regulation of PAL and CHS in tomato remains unknown. We identified 31 putative KFB-encoding genes in the tomato genome. Our homology analysis and phylogenetic study predicted four PAL-interacting SIKFBs, while SIKFB18 was identified as the sole candidate for the CHS-interacting KFB. Consistent with their homolog function, the predicted four PAL-interacting SIKFBs function in PAL degradation. Surprisingly, SIKFB18 did not interact with tomato CHS and the overexpression or knocking out of SIKFB18 did not affect phenylpropanoid contents in tomato transgenic lines, suggesting its irreverence with flavonoid metabolism. Our study successfully discovered the post-translational regulatory machinery of PALs in tomato while highlighting the limitation of relying solely on a homology-based approach to predict interacting partners of F-box proteins.

Key message

Despite its highest sequence homology with Arabidopsis CHS-interacting KFB among 31 tomato KFBs, SIKFB18 does not function in CHS degradation, while predicted PAL-interacting SIKFBs function in PAL degradation in tomato.

Keywords Phenylpropanoids · PAL · CHS · F-box · KFB · Solanum lycopersicum

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Introduction

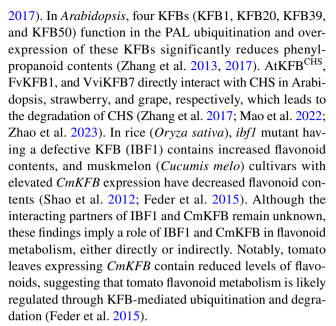
Phenylpropanoids are a group of specialized metabolites, encompassing flavonoids, condensed tannins, hydroxycinnamoyl compounds, volatile phenylpropanoids, and monolignols (Deng and Lu 2017; Dong and Lin 2021; Garibay-Hernández et al. 2021). They are ubiquitously present in the plant kingdom (Liu et al. 2015; Garibay-Hernández et al. 2021) and play vital roles in plant survival. Monolignols, for instance, serve as building blocks of lignin, providing rigidity and hydrophobicity to vascular bundles (Muro-Villanueva et al. 2019). Flavonoids protect plants from various stresses (Agati et al. 2020; Shomali et al. 2022) and regulate plant growth and development by modulating auxin transport and scavenging reactive oxygen species (Brown et al. 2001; Yin et al. 2014; Muhlemann et al. 2018; Tan et al. 2019;



Chapman and Muday 2021; Teale et al. 2021). Kaempferol, a flavonol aglycone, serves as a precursor for ubiquinone, an essential respiratory cofactor. (Soubeyrand et al. 2018, 2021; Fernández-Del-Río et al. 2020; Berger et al. 2022). Moreover, numerous phenylpropanoids, especially flavonoids, exhibit beneficial properties for human health, such as anti-cancer, anti-diabetes, and antioxidant activities (Wedick et al. 2012; Tu et al. 2017; Bondonno et al. 2019; Wen et al. 2021; Prasanna and Upadhyay 2021; Micek et al. 2021; Xian et al. 2021; Slika et al. 2022). Therefore, understanding phenylpropanoid biosynthesis and its regulation is imperative for engineering enhanced phenylpropanoid production in crops (Sun et al. 2020; Chen et al. 2021).

Phenylpropanoid biosynthesis starts with the deamination of phenylalanine to produce cinnamic acid by phenylalanine ammonia-lyase (PAL) (Zhang and Liu 2015). Subsequent hydroxylation and ligation reactions produce *p*-coumaroyl-CoA, a precursor for hydroxycinnamoyl compounds like monolignols and flavonoids (Vogt 2010) (Fig. 1a). The first enzyme directing flux from the production of hydroxycinnamoyl compounds to flavonoid biosynthesis is chalcone synthase (CHS) that produces naringenin chalcone using *p*-coumaroyl-CoA and malonyl-CoA (Fig. 1a) (Grotewold 2006; Saito et al. 2013). Then, sequential reactions of isomerases, hydroxylases, and reductases generate basic structures of flavonoid skeletons such as flavonol aglycones and anthocyanidins (Wen et al. 2020).

The phenylpropanoid pathway is regulated through intricate mechanisms, including feed-forward, feed-back, transcriptional, post-transcriptional, and post-translational regulations (Yin et al. 2012; Xu et al. 2015; Shin et al. 2015; Verweij et al. 2016; Zhang et al. 2017; Ohno et al. 2018; Wang et al. 2020). Recent studies have revealed that PAL and CHS activities are regulated post-translationally through ubiquitin-dependent proteasomal degradation (Zhang et al. 2013, 2017; Gu et al. 2019; Mao et al. 2022; Zhao et al. 2023). Ubiquitination is a protein modification that adds the small regulatory protein called ubiquitin (Ub) to lysine residues of substrate proteins and poly-ubiquitinated proteins are subsequently degraded by the 26S proteasome (Hristova et al. 2020). Ubiquitination requires ubiquitinactivating enzyme (E1), ubiquitin conjugating enzyme (E2), and ubiquitin ligase (E3). Ubiquitin is activated by the E1 enzyme, and then transferred to E2. The E3 complex then adds ubiquitin from E2 to target proteins. The E3 ubiquitin ligase includes E2 binding protein, scaffold protein, adaptor protein, and substrate binding protein such as F-box proteins (Gray and Estelle 2000). Several Kelch domain-containing F-box proteins (KFBs) were identified as subunits of E3 ligase functioning in the ubiquitination of PAL and CHS, the two vital enzymes functioning at the entry points of the general phenylpropanoid pathway and flavonoid pathway, respectively (Fig. 1a) (Zhang and Liu 2015; Zhang et al.



Tomato is one of the most widely consumed vegetables globally, serving as an excellent source of beneficial phytonutrients, including phenylpropanoids (Chandra et al. 2012; Anwar et al. 2019). Tomato accumulates various phenylpropanoids including flavonoids, caffeic acid derivatives, stilbenes, coumarins, monolignols, aurones (Zhang et al. 2015b). Despite advances in our understanding of the phenylpropanoid metabolism in tomato (Zhang et al. 2015b; Tohge et al. 2017; Rosa-Martínez et al. 2023), the post-translational regulation of phenylpropanoid metabolism in tomato remains unknown. In this study, we aimed to identify tomato KFBs (SlKFBs) involved in phenylpropanoid metabolism in tomato (Solanum lycopersicum). Our homology study identified 31 genes encoding putative KFB in the tomato genome, and we investigated their functions in phenylpropanoid metabolism.

Materials and methods

Genetic material and plant growth conditions

Micro-Tom was obtained from the tomato genetics resource center located at the University of California, Davis, led by C.M. Rick. To conduct the BiFC analysis, we used *Nicotiana benthamiana*. Tomato and tobacco plants were grown under controlled conditions of 22 °C \pm 1 °C with a 16 h day and 8 h photoperiod.

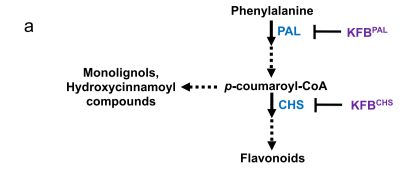
Retrieval of KFBs from tomato genome

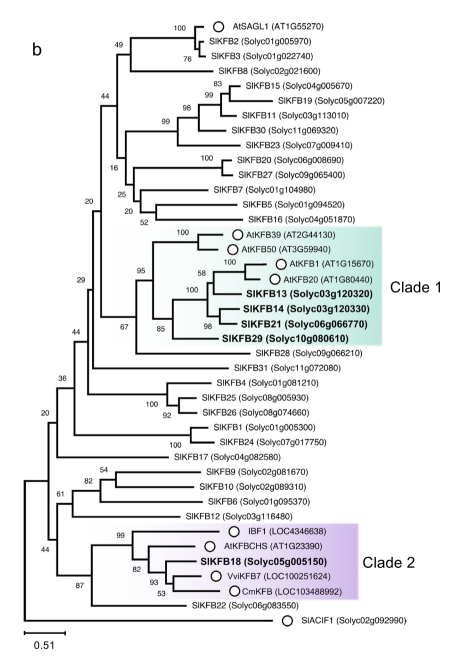
Kelch-domain containing F-box proteins (KFBs) were identified in the tomato genome (SL4.0 build; ITAG4.0 annotation) (Tomato Genome Consortium 2012) using the



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Fig. 1 Proteolytic regulation steps in the phenylpropanoid pathway in Arabidopsis and a phylogenetic tree of tomato KFBs. a Chalcone synthase (CHS) and phenylalanine ammonia-lyase (PAL) are regulated through ubiquitinmediated proteolysis in Arabidopsis. AtKFBCHS and AtKFBPAL are kelch-domain containing F-box proteins responsible for ubiquitination of CHS and PAL, respectively. **b** Phylogenetic analysis with 31 putative SlKFBs retrieved from the tomato genome and 9 characterized KFBs from other plant species identified two sub clades including AtKFBCHS and AtKFB^{PAL}. A phylogenetic tree was constructed using Maximum Likelihood method with 1000 bootstrap samples and the JTT model. The tree was constructed with the full-length protein sequences of 31 SIKFBs from the tomato reference genome (SL4.0 build; ITAG4.0 annotation) (Tomato Genome Consortium 2012) and 9 KFBs from Arabidopsis, rice, grape, and muskmelon (marked with open circle) known for their role in regulating phenylpropanoid metabolism. Branch lengths are drawn to scale, with the scale bar indicating the number of amino acid substitutions per site. The four AtKFBPALs (AtKFB1, 20, 39, and 50) are clustered in Clade 1, and the KFBs known to regulate flavonoids are in Clade 2. SIACIF1 serves as an outgroup







Jackhmmer program (version 2.41.2) within HmmerWeb (https://www.ebi.ac.uk/Tools/hmmer/) (Fernandez-Pozo et al. 2015; Potter et al. 2018). A Hidden Markov Model (HMM) profile was constructed by initially querying the sequence of Kelch domain and F-box domain in the Arabidopsis KFB^{CHS} (AT1G23390). This model was then employed to iteratively search the tomato protein database, which includes UniProtKB and SwissProt. The search was conducted separately for proteins containing either the Kelch domain or F-box domain until no additional proteins were added to the retrieved protein lists. Proteins containing both the F-box and Kelch domain were selected by cross-referencing the protein lists obtained with the HMM profile of each domain separately. In total, this method identified 31 KFBs in the tomato genome.

Gene matrix construction

To generate the amino acid sequence identity matrix, we utilized Clustal Omega (version: Clustal2.1). We used the default configurations for the analysis.

Yeast two hybrid (Y2H) assay

The coding sequences (CDS) of seven SIKFBs (Solyc01g005970, Solyc03g120320, Solyc03g120330, Solyc05g005150, Solyc06g066770, Solyc06g083550, Solyc10g080610), AtKFBCHS (At1g23390), CmKFB (XP008446188), SICHS1 (Solyc09g091510), SICHS2 (Solyc05g053550), S1CHI (Solyc05g010320), S1F3H (Solyc02g083860), SIPAL5 (M83314), and AtCHS (At5g13930) with the appropriate restriction enzyme site at the end of the CDS were synthesized from Twist Bioscience (CA, USA). Arabidopsis PAL1, PAL2, PAL3, and PAL4 were cloned from vectors purchased from ABRC (stock number: pDEST-DB004H07 for AtPAL1, pDEST-DB030E01 for AtPAL2, CIW05433 for AtPAL3, and pDEST-DB101F06 for AtPAL4) using primer numbers 32-39 (Supplementary Table S1). The CDS of KFBs were subcloned into the pGADT7 vector (catalog number: 630442, Takara Bio, Otsu, Japan), while the CDS of CHS, PAL, CHI, F3H were subcloned into the pGBKT7 vector (catalog number: 630443, Takara Bio, Otsu, Japan). The empty pGADT7 and pGBKT7 vectors were used as negative controls. The constructed vectors were then co-transformed into the yeast strain Y2HGold (catalog number: 630489, Takara Bio, Otsu, Japan) using the lithium acetate-mediated transformation method (Gietz and Woods 2002). To screen the transformed yeast strains, we used SD media (5 g of ammonium sulfate, 3.4 g of yeast nitrogen base without amino acids, 20 g of D-glucose, 20 g of agar per liter) supplemented with dropout amino acids, excluding leucine and tryptophan. We used two SD media, dropout-SD and dropout-SD with Aureobasidin A (AbA), to assess protein–protein interactions. *AUR1-C*, a mutated version of the *AUR1* reporter gene in this Y2H system, allows yeast to survive on media containing AbA. AbA inhibits the wild-type AUR1 protein, which is lethal to yeast, but AUR1-C provides resistance. Dropout-SD lacked leucine, tryptophan, histidine, and adenine, while dropout-SD with AbA was the same as dropout-SD but included AbA for strong selectivity.

Bimolecular fluorescence complementation (BiFC) assay

Two different BIFC systems were used in this study. For the BiFC with PAL, we adopted a vector system from (Han et al. 2022). PCR amplification was performed using specific primers (Supplementary Table S1). The PCR products of KFBs (full length or truncated proteins lacking F-box domain) were cloned into the pUC19/Vc-C vector (Addgene #183,158), while SIPAL5 was cloned into the pYL322d1/ Vn-C vector (Addgene #183,154). The pUC19/Vc-c constructs were then linearized with the AscI enzyme, and the pYL322d1/Vn-C constructs were linearized with AscI and SbfI enzymes. Additionally, a linearized DNA fragment containing the mCherry marker fused with a nuclear localization sequence (NLS) was obtained from pUC19/NLS-mChe (Addgene #183,162) using the AscI enzyme. The three DNA fragments were assembled into the plant binary vector, pYL1300UaUf (Addgene #183,173) using the NEBuilder® HiFi DNA Assembly Cloning Kit (catalog number: E5520S, NEB, Ipswich, MA, USA). The protocol for the BIFC with CHS was adopted from a previous report with slight modifications (Nakabayashi et al. 2015). The F-box domains of SIKFB18 and AtKFB^{CHS} were removed to avoid substrate degradation. The truncated KFBs (lacking F-box domain) and the full-length of CHSs were amplified with the attB sequence at the end of CDS by PCR with specific primers listed in Supplementary Table S1. The PCR products were subsequently inserted into the pCC1155 (Zhang et al. 2020) for gateway cloning using BP ClonaseTM II Enzyme Mix (catalog number: 11789020; Thermo Fisher Scientific, Waltham, MA, USA). The pCC1155 with KFB constructs were subcloned into the pBatTL-B-sYFPn vector, and the pCC1155 with CHS constructs were subcloned into pBatTL-B-sYFPc using LR ClonaseTM II Enzyme Mix (catalog number: 11791; Thermo Fisher Scientific, Waltham, MA, USA). Empty pBatTL-B-sYFPn and pBatTL-B-sYFPc were utilized as negative controls.

All BiFC constructs were transformed into *A. tumefaciens* strain GV3101 and co-infiltrated into 4 weeks-old *Nicotiana benthamiana* leaves. The YFP, Venus, and mCherry fluorescence signals were detected using a confocal scanning microscope (Olympus IX81-DSU) 48 h after infiltration.



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Generation of transgenic lines

To generate Arabidopsis transgenic lines overexpressing SIKFB13 and SIKFB14, coding sequences of SIKFB13 and SIKFB14 were synthesized by Twist Bioscience (San Francisco, CA, USA). These sequences were subsequently cloned into the pCC1155 entry vector using the BP ClonaseTM II enzyme mix (catalog no. 11789020; Thermo Fisher Scientific, Waltham, MA, USA), and then were recombined into the pCC0995 destination vector via the LR ClonaseTM II enzyme mix (catalog no. 11791; Thermo Fisher Scientific, Waltham, MA, USA). These constructs were then introduced into wild-type Arabidopsis plants (Col-0) using the Agrobacterium tumefaciens-mediated floral dip method, as described by (Zhang et al. 2006). The resulting T1 generation seedlings were initially selected on soil with 0.2% (w/v) BASTA (glufosinate ammonium), and surviving plants were subsequently transplanted to fresh soil for further assays. To determine the expression levels of SlKFBs in the transgenic Arabidopsis lines, RT-PCR was performed using specific primers No. 24 and 25 for SIKFB13, No. 26 and 27 for SIKFB14, and No. 28 and 29 for AtTUB3 as an internal control.

To generate tomato transgenic lines overexpressing *SlKFB18* and CRISPR lines,

the gRNA-containing (5'-GCTTCAACAAGCCGAAGC CG-3') pHSE401 CRISPR vectors to target the upstream region of the SIKFB18 gene and the pGWB502 overexpression vectors harboring the SIKFB18 CDS were introduced into the A. tumefaciens GV3101 using the heat shock method. The tissue culture-mediated tomato transformation was conducted using the previously described method with slight modifications (Gupta and Van Eck 2016). Cotyledons from 10 days-old seedlings were excised and incubated on filter paper on preculture media plates containing 4.3 g of Murashige and Skoog (MS) salt, 100 mg of myoinositol, 1 ml of modified Nitsch vitamins stock(composed of 10 mg of glycine, 50 mg of nicotinic acid, 2.5 mg of pyridoxine HCl, 2.5 mg of thiamine HCl, 2.5 g of folic acid, and 0.2 mg of d-biotin in 10 ml), 20 g sucrose, 5.2 g of TC gel, and 2 ml of 1 mg/ml trans-zeatin stock per litter, with the pH adjusted to 6. The cotyledons were incubated under a 16 h photoperiod for 24 h. Agrobacterium cells carrying the vector were incubated overnight and subsequently harvested via centrifugation. The pellet of Agrobacterium cells was re-suspended in a buffer containing 4.3 g of MS salts, 100 mg of myoinositol, 0.4 mg of thiamine HCl, and 20 g of sucrose per liter, with the pH adjusted to 5.8. The cotyledons were then immersed in the Agrobacterium suspension for 5 min and placed back onto preculture media for co-culture in the dark at 22 °C for 48 h. The cotyledons were then transferred to callus induction media, which contained 3 ml of 100 mg/ml timentin stock and 3 mg/l hygromycin per liter and were incubated for 2 weeks. Once small calli had formed, the cotyledons with the calli were transferred to a shoot induction media. The concentration of trans-zeatin in shoot induction media was reduced to 50% of the amount in the callus induction media, while all other ingredients remained the same. The calli were transferred to fresh shoot induction media every 2 weeks until shoot formation was observed. Once the shoots reached a minimum length of 2 cm with at least one node, they were excised and planted in a root regeneration media composed of 4.3 g of MS salt, 1 ml of modified nitch vitamins stock, 30 g of sucrose, 8 g of Difco Bacto Agar, 2 ml of 100 mg/ml timentin stock, 3 mg of hygromycin, and 1 mg of indole acetic acid per liter with the pH adjusted to 6. When the roots were sufficiently formed, the plantlets were moved to soil and grown until seed maturation. The CRISPR knock-out lines were isolated via sequencing of the target genes. To check the genotype of modified region of SIKFB18, the primer set, No. 30 and 31, was used (Supplementary Table S1). Transgenic plants overexpressing SIKFB18 were isolated based on their resistance to hygromycin. The expression of SIKFB18 was then confirmed by qRT-PCR using the primer set of No. 22 and 23. SIACTIN2 expression was used as an internal control (Supplementary Table S1).

PAL activity

The PAL activity was measured by a modified method of the procedure described in (Kim et al. 2015). Total proteins were extracted from the leaves of 3 weeks-old Arabidopsis, which were pulverized in a Benchmark BeadBlaster 24 homogenizer (Benchmark Scientific, NJ). The powdered tissue was then incubated in extraction buffer comprising 0.1 M Tris-HCl (pH 8.3), 10% glycerol, and 5 mM dithiothreitol (DTT) for one hour and then crude proteins were collected after centrifugation. Protein concentrations in the extracts were quantified via the Bradford assay, employing the Bradford Reagent (Sigma-Aldrich, St. Louis, MO, USA) according to the manufacturer's protocol. The enzyme reaction of PAL was started by adding 150 µl of protein extract with 400 µl of a reaction buffer that contained 5 mM L-phenylalanine, and the mixture was incubated at 37 °C for 90 min. The reaction was terminated by adding 40 µl of 30% (v/v) acetic acid. The product of the enzyme reaction was then extracted with ethyl acetate, the volume of which was 600 µl, and subsequently concentrated using an Eppendorf Vacufuge Plus (Eppendorf, Hamburg, Germany). The dried extract was then redissolved in 100 µl of 50% methanol and 10 µl of extract was analyzed using HPLC with a solvent B (100% acetonitrile) gradient in solvent A (0.1% formic acid in water). The gradient starting from 12 to 30% of solvent B over 2.6 min, increasing from 30 to 95% in the next 4 min, and holding at 95% for an additional 3 min. The



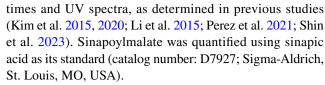
flow rate was set at 0.7 ml/min, and the column temperature was maintained at 40 °C. The PAL reaction product, trans-cinnamic acid, was quantified by measuring the peak area at 270 nm and comparing it to a calibrated curve of a standard *trans*-cinnamic acid solution (Sigma-Aldrich, St. Louis, MO, USA).

RNA extraction and gene expression analysis

Leaf samples from 4 weeks old tomato plants were collected and immediately frozen in liquid nitrogen. The samples were then homogenized using a Benchmark BeadBlaster 24 homogenizer (Benchmark Scientific, Edison, NJ, USA) with 500 µl of 1.25 mm Zirconia oxide beads (A Norstone Company, Bridgeport, PA, USA) to obtain a complete ground sample. Total RNA was extracted from the samples using Trizol reagent (Life Technologies Inc., Gaithersburg, MD, USA) according to the manufacturer's instructions. For cDNA synthesis, 2 µg of total RNA and a reverse transcription kit (catalog number: 4368814; Thermo Fisher Scientific, Waltham, MA, USA) were used. Quantitative real-time Reverse Transcription PCR (qRT-PCR) and RT-PCR were performed using PCR kits (catalog number: FERK1071; Thermo Fisher Scientific, Waltham, MA, USA, and catalog number: K1081; Thermo Fisher Scientific, Waltham, MA, USA) with 1 µl of cDNA. AtTUB3 (AT5G62700) and SlAC-TIN2 (Solyc11g005330) were used as internal controls for gene expression analysis in Arabidopsis and tomato, respectively. Specific forward and reverse primers, as listed in Supplementary Table S1, were used for PCR.

Phenylpropanoid quantification

The first three true leaves from 3 weeks old tomato plants and 3rd and 4th rosette leaves of 3 weeks old Arabidopsis were used for metabolite analysis using 50% (v/v) methanol at 65 °C for 1 h. The extracts were then subjected to analysis using an UltiMate 3000 HPLC system (ThermoFisher Scientific, Waltham, MA, USA) equipped with an AcclaimTM 120 C18 column (75 mm \times 3 mm; 2.2 μ m) coupled with a C18 guard column (10 mm × 3 mm; 5 µm) (ThermoFisher Scientific, Waltham, MA, USA). Metabolites from Arabidopsis samples were separated using a mobile phase composed of solvent A (0.1% formic acid (v/v) in water) and solvent B (100% acetonitrile), with a gradient of 5% to 14% for 2.2 min, followed by 14% to 18% (v/v) solvent B for 9 min, and finally 18% to 95% solvent B for 3.5 min. Three kaempferol glycosides were identified by comparing the HPLC profiles of wild type, ugt78d1, and ugt78d2 mutants, following previous studies (Yin et al. 2012, 2014). The levels of kaempferol glycosides were compared based on their HPLC peak areas. Sinapate esters, including sinapoylmalate and sinapoylglucose, were identified based on their retention



Metabolites from the tomato samples were separated with a mobile phase consisting of solvent A (0.1% formic acid (v/v) in water) and solvent B (100% acetonitrile) with a gradient of 3% to 18% for 17 min, followed by 18% to 50% (v/v) solvent B for 6 min. The contents of rutin and chlorogenic acid were quantified based on peak area and a standard curve of rutin (catalog number: R5143; Sigma-Aldrich, St. Louis, MO, USA) and chlorogenic acid (catalog number: C0181; TCI America, Portland, OR, USA).

Results

putative SIKFBs were identified in the tomato genome

From the tomato genome (SL4.0 Assembly), we identified a total of 31 genes encoding proteins with both F-box and Kelch domains, which we considered to be putative KFB homologs in tomato. We designated them as SIKFB1 to SIKFB31 based on their positions within the tomato chromosomes (Supplementary Table S2). To identify SIKFB candidates targeting CHS and PAL, we conducted a comparative analysis with the amino acid sequences of tomato KFBs and previously characterized KFBs from Arabidopsis, rice, grape, and muskmelon that were reported to regulate phenylpropanoid production (Shao et al. 2012; Zhang et al. 2013, 2015a, 2017; Feder et al. 2015; Zhao et al. 2023) (Fig. 1b). In the phylogenetic tree, four PALinteracting KFBs from Arabidopsis (AtKFB01, AtKFB20, AtKFB39, and AtKFB50) were clustered in clade 1, while clade 2 included KFBs functioning in flavonoid metabolism, including Arabidopsis CHS-interacting KFB (AtKFBCHS) (Fig. 1b).

Four tomato KFBs, S1KFB13 (Solyc03g120320), S1KFB14 (Solyc03g120330), S1KFB21 (Solyc06g066770), and S1KFB29 (Solyc10g080610), were clustered with PALinteracting KFBs in clade 1 (Zhang et al. 2013, 2015a) (Fig. 1b). S1KFB13, S1KFB14, and S1KFB21 showed 45 to 50% amino acid sequence identities with AtKFB1 and AtKFB20, while S1KFB29 showed 36% and 37% amino acid sequence identities with AtKFB1 and AtKFB20, respectively (Supplementary Fig. S1). All four S1KFBs displayed sequence identities ranging from 30 to 36% with AtKFB39 and AtKFB50. S1KFB28 (Solyc09g066210), the closest KFB to those in clade 1, showed only 24% to 27% sequence identities with AtKFB1, 20, 39, and 50 (Fig. 1b, Supplementary Fig. S1). Thus, S1KFB13, S1KFB14, S1KFB21, and



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SIKFB29 in clade 1 were selected as PAL-interacting SIKFB candidates.

SIKFB18 was the only SIKFB in clade 2 having AtKFB^{CHS}, IBF1, VviKFB7, and CmKFB, the KFBs functioning in flavonoid metabolism (Fig. 1b). SIKFB22 (Solyc06g083550) was SIKFB close to clade 2, but SIKFB22 showed 20% to 24% sequence identities with AtKFB^{CHS}, IBF1, CmKFB, and VviKFB7 while SIKFB18 exhibited over 40% sequence identity with them (Supplementary Fig. S1). Thus, SIKFB18 was selected as a CHS-targeting KFB candidate.

SIKFB13, SIKFB14, SIKFB21, and SIKFB29 regulate PAL stability

In Arabidopsis, four Phenylalanine Ammonia-Lyase (PAL) enzymes, AtPAL1, AtPAL2, AtPAL3, and AtPAL4, function redundantly (Rohde et al. 2004; Huang et al. 2010) and four KFBs, AtKFB1, AtKFB20, AtKFB39, and AtKFB50, participate redundantly in the ubiquitination of all four AtPALs (Zhang et al. 2013, 2015a). We identified six PAL homologs (Solyc05g056170, Solyc10g086180, Solyc09g007890, Solyc09g007910, Solyc09g007900, and Solyc09g007920) in the tomato genome (SL4.0 build; ITAG4.0 annotation) (Tomato Genome Consortium 2012) that showed over 70% sequence identities with Arabidopsis PALs (Supplementary Fig. S2). Solyc10g086180, Solyc09g007890, Solyc09g007910, Solyc09g007900, and Solyc09g007920

were shown to be upregulated in *AtMYB12* overexpression tomato, along with other flavonoid biosynthesis enzymes (Zhang et al. 2015b). Additionally, SIPAL5 (M83314.1) has been reported as a tomato PAL in previous studies (Guo and Wang 2009; Løvdal et al. 2010). Interestingly, all seven SIPALs are more closely related to AtPAL1 and AtPAL2 than to AtPAL3 and AtPAL4 in the phylogenetic tree (Supplementary Fig. S2). As SIPAL5 has been previously characterized (Guo and Wang 2009; Løvdal et al. 2010), we decided to use SIPAL5 to identify PAL-interacting SIKFBs in tomato.

In our Y2H analysis, we included four SIKFBs, SIKFB13 (Solyc03g120320), SIKFB14 (Solyc03g120330), SIKFB21 (Solyc06g066770), and SIKFB29 (Solyc10g080610) from clade 1 (Fig. 1b). Among them, SIKFB14, SIKFB21 and SIKFB29 interacted with SIPAL5 (Fig. 2). Notably, SIKFB13 (Solyc03g120320) did not interact with SIPAL5 in our Y2H assay, despite its high sequence similarity with SIKFBs in clade 1 (Fig. 1b, 2; Supplementary Fig. S1).

To test if the interaction of SIKFB14/21/29 with SIPAL5 affects protein stability, we conducted BiFC experiments using intact SIKFBs and truncated SIKFBs (SIKFB (Δ)) with the F-box domain removed (Fig. 3). In this test, we included a nuclear-localized mCherry cassette to evaluate proper transformation (Fig. 3a). All tested samples showed mCherry signals in nucleus, indicating properly expressed transgenes. The detection of the Venus signal was evident in leaves that were infiltrated with SIKFB14, SIKFB21, and

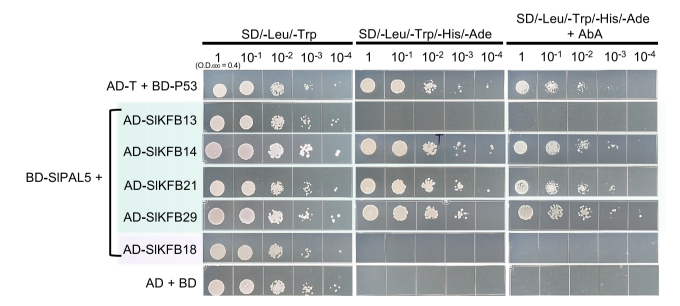


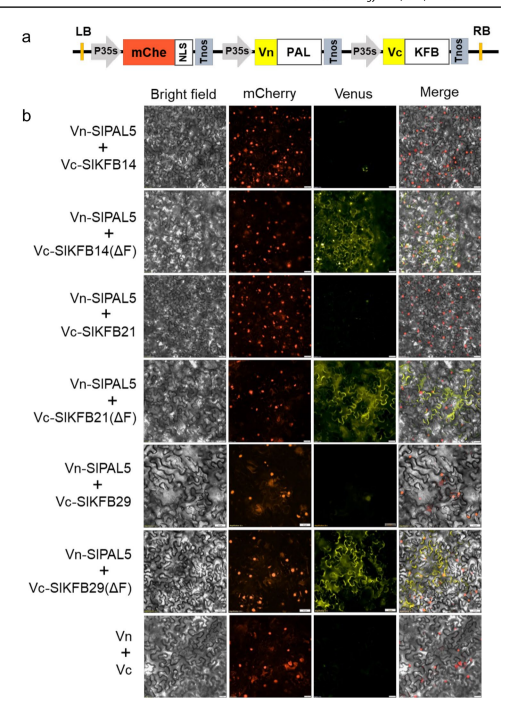
Fig. 2 Three SIKFBs interacted with SIPAL5 in the Y2H assay SIKFBs were fused with the activation domain (AD), and SIPAL5 was fused with the binding domain (BD). SD media excluding leu and trp were used to check the introduction of AD and BD vectors in the yeast. SD media excluding leu, trp, his, and ade were used to assess protein interactions. SD media excluding leu, trp, his, and ade

along with Aureobasidin A (AbA) were used to increase stringency of interactions. SIKFB14, 21, and 29 in clade 1 showed interaction with SIPAL5, while SIKFB13 in the same clade and SIKFB18 in clade 2 did not. The pair of AD-T and DB-P53 was used as a positive control, and the pair of AD and BD vector without insert DNA was used as a negative control



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Fig. 3 BiFC assay for the interactions of SIKFBs with SIPAL5. a The illustration of the BiFC vector used in this assay. P35s (35S promoter), mChe (mCherry CDS), NLS (Nuclear localization signal sequence), Tnos (Nos terminator), Vn (Venus N-terminal fragment), Vc (Venus C-terminal fragment), PAL (Phenylalanine ammonialyase CDS), and KFB (Full length or partial CDS of Kelch domain-containing F-box). b SIKFB14/21/29 interacting with SIPAL5 exhibited fluorescence when their F-box domain was removed, but intact SIKFBs coinfiltrated with SIPAL5 did not show fluorescence. The BiFC was conducted using intact SIKFBs, and truncated SIKFBs that lack the F-box domains (SIKFB14 (\Delta F), SIKFB21 (ΔF) , and SIKFB29 (ΔF) that first 46, 51 and 49 amino acids were removed, respectively). Images captured under identical exposure conditions depict bright field, mCherry (captured using Wide-band Green-Red excitation light with a center wavelength of 593 nm, and a TRITC filter), Venus (captured using blue excitation light with a center wavelength of 470 nm and a FITC filter) and merged two-channel views. More than five infiltrated leaves were examined, and similar images were obtained



SIKFB29 lacking their F-box domains, along with SIPAL5 (Fig. 3b). However, this signal was not observed with intact SIKFBs (Fig. 3b). This result suggests that SIPAL5, upon interacting with SIKFBs, undergoes degradation through a mechanism requiring the F-box domain, likely involving ubiquitination-mediated degradation.

In Arabidopsis, all four KFBs redundantly interact with four PALs (AtPAL1 ~ AtPAL4) (Zhang et al. 2013, 2015a). As there are at least six additional PAL-encoding genes in the tomato genome, SIKFB14/21/29 may interact with other

SIPALs in addition to SIPAL5. Similarly, SIKFB13 may interact with other PAL proteins, although it did not interact with SIPAL5. To test if SIKFB13 functions in PAL degradation, we took advantage of Arabidopsis system. Arabidopsis rosette leaves accumulate sinapoylmalate, a blue fluorescent phenylpropanoid (Ruegger and Chapple 2001), causing Arabidopsis leaves to emit a bluish color under UV light. Conversely, Arabidopsis plants with reduced sinapoylmalate would appear red under UV light due to chlorophyll auto-fluorescence (Ruegger and Chapple 2001). As PAL



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activity is necessary for the production of phenylpropanoids, including sinapovlmalate, accelerated PAL degradation can be detected with leaf UV fluorescence. Given that tomato PALs showed over 80% sequence identities with Arabidopsis PAL1/2/4, which is higher than 73% sequence identities of AtPAL3 when compared with AtPAL1 and 2 (Supplementary Fig. S2a), it is possible that SIKFBs may interact with AtPALs. Thus, we overexpressed *SlKFB13* in Arabidopsis to test its impact on phenylpropanoid production. We also overexpressed SlKFB14, which interacts with SlPAL5 from Y2H and BiFC assays (Fig. 2, 3). As shown in Fig. 4, some Arabidopsis transgenic lines overexpressing SlKFB13 or SlKFB14 exhibited a red color under UV light, while others displayed a bluish color. Consistently, the lines showing a red color under UV light accumulated lower levels of phenylpropanoids, including three kaempferol glycosides, sinapolymalate, and sinapoylglucose, compared to the lines exhibiting a blue color (Fig. 4, Supplementary Fig. S3a). The reduced phenylpropanoid contents correlated with PAL activity and the expression levels of SIKFB13 and SIKFB14 (Fig. 4), suggesting the repressive roles of SIKFB13 and SIKFB14 on PAL activity. Notably, several Arabidopsis transgenic lines with strong expression of SIKFB13 and SlKFB14 displayed alteration in growth and development, such as stunted inflorescence growth (Supplementary Fig. S3b), similar to those observed in the Arabidopsis pal mutants (Huang et al. 2010).

To determine whether any interactions of SIKFBs with AtPALs contributed to the reduced PAL activity and phenylpropanoid contents, we used Y2H assays to examine the interactions between SIKFBs (SIKFB13 and SIKFB14) and Arabidopsis PALs (AtPAL1, AtPAL2, AtPAL3 and AtPAL4) (Rohde et al. 2004; Huang et al. 2010). SIKFB14 indeed interacted with AtPAL1, AtPAL2, and AtPAL4, while SIKFB13 interacted with AtPAL1 and AtPAL4, suggesting a possible role of SIKFBs in PAL stability (Supplementary Fig. S4). Interestingly, both SIKFBs did not interact with AtPAL3.

SIKFB18 does not function in flavonoid metabolism in tomato

In Arabidopsis, AtCHS (At5g13930) is the only CHS functioning in flavonoid biosynthesis as its loss-of-function mutant failed to make any flavonoids (Schmelzer et al. 1988; Shirley et al. 1995). We identified four CHS homologs in the tomato genome (SL4.0 Assembly), which showed over 60% sequence identities with the Arabidopsis CHS (AtCHS) (Supplementary Fig. S5). Among the four identified tomato CHS homologs, SICHS1 (Solyc09g091510) and SICHS2 (Solyc05g053550) have been previously characterized (Schijlen et al. 2007; España et al. 2014; Kong et al. 2020). Silencing SICHS1 resulted in a notable reduction in

flavonoid content (Schijlen et al. 2007; España et al. 2014; Kong et al. 2020). The other two SICHS homologs were designated as SICHS-like1 (Solyc12g098090) and SICHS-like2 (Solyc05g053170) (Supplementary Fig. S5a). SICHS1 and SICHS2 showed approximately 85% sequence identities with AtCHS, while SICHS-like1 and SICHS-like2 exhibited sequence identities of 65% and 66% with AtCHS, respectively (Supplementary Fig. S5b). According to the public database (Ruprecht et al. 2017), SICHS1 and SICHS2 are expressed in most organs, while the expression of SICHS-like1 and SICHS-like2 are barely detected (Supplementary Fig. S5c). Thus, we used SICHS1 and SICHS2 to identify SICHS-interacting SIKFBs.

To assess protein-protein interactions between SIKFB candidates and SICHS1/2, we employed the Y2H system. Given that AtKFB^{CHS} interacts with AtCHS (Zhang et al. 2017), we included them as a positive control for our Y2H assay. Previous studies have indicated that overexpression of CmKFB from muskmelon reduces flavonoid content in tomato, yet its target protein(s) remains unknown (Feder et al. 2015). It is possible that CmKFB functions in CHS degradation. Therefore, we also included CmKFB in our Y2H assay. As expected, AtKFB^{CHS} interacted with AtCHS (Fig. 5). Interestingly, AtKFBCHS and CmKFB interacted with both SICHS1 and SICHS2 (Fig. 5). The interaction of AtKFB^{CHS} and CmKFB with tomato CHS suggests that SICHS1 and SICHS2 may have the binding site for KFBs (Fig. 5). Our phylogenetic study identified only one SIKFB, SIKFB18, in clade 2, where CHS-targeting KFBs or flavonoid-related KFBs clustered (Fig. 1b). Despite having the highest sequence identity with known CHS-targeting KFBs (Supplementary Fig. S1), SIKFB18 did not interact with either SICHS1 or SICHS2. Similarly, SIKFB22, the KFB closest to SIKFB18, also did not interact with SICHS1 and SICHS2 (Figs. 1b and 5). We further investigated whether SIKFB18 interacts with two other flavonoid biosynthesis enzymes downstream of CHS in the flavonoid biosynthesis pathway, namely SICHI (Solyc05g010320) and SIF3H (Solyc02g083860). However, SIKFB18 did not interact with either in our Y2H assay (Supplementary Fig. S6).

We then tested these interactions in *Nicotiana benthamiana* using the bimolecular fluorescence complementation (BiFC) method. In the BiFC assay, we used truncated KFBs, AtKFB^{CHS} (AtKFB^{CHS} (Δ)) and SlKFB18 (SlKFB18 (Δ)), where the F-box domain was removed to prevent the degradation of target proteins after interaction. Consistent with Y2H results, no interaction between SlKFB18 and SlCHS1 was observed, while AtKFB^{CHS} interacted with both AtCHS and SlCHS1 (Fig. 6).

It is possible that SIKFB18 may interact with SICHS in vivo or it may interact with SICHS-likes or other flavonoid biosynthesis enzymes. To further investigate the potential involvement of SIKFB18 in flavonoid metabolism, we



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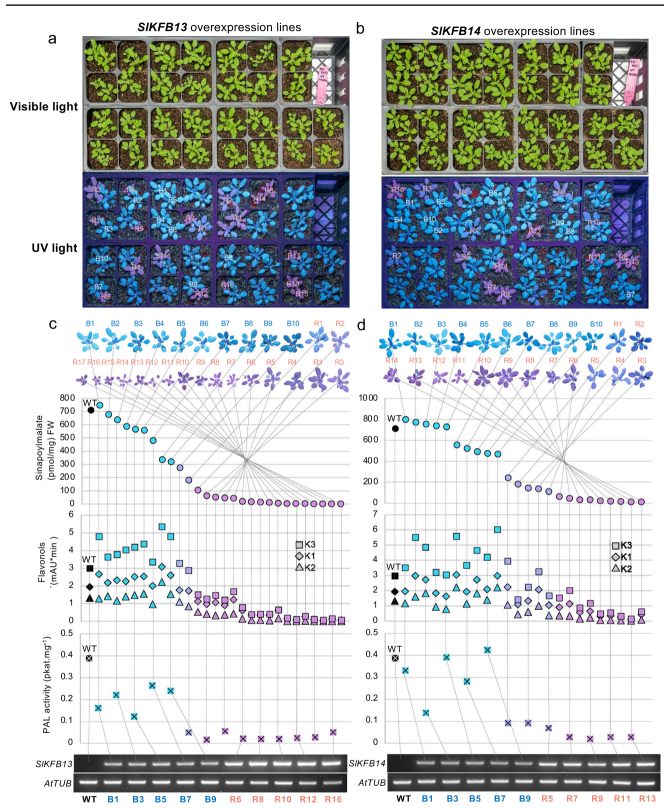


Fig. 4 PAL activity and phenylpropanoid contents in the transgenic lines overexpressing *SlKFB13* or *SlKFB14*. **a, b** 3 weeks old plants expressing *SlKFB13* (A) or *SlKFB14* (B) photographed under visible light (top) and UV light (bottom), Plants exhibiting red fluorescence under UV light indicate a low level of sinapoylmalate compared to those with blue fluorescence. **c, d** The levels of sinapoylmalate, three

kaempferol glycosides, and PAL enzyme activity in plants. The relative expression levels of *SlKFB13* and *SlKFB14* in representative high- or low-sinapoylmalate accumulation plants were shown with RT-PCR. *AtTUB3* (AT5G62700) was used for an internal control. Kaempferol-3-*O*-glu-7-*O*-rha (K2), Kaempferol-3-*O*-rha-7-*O*-rha (K3), Kaempferol-3-*O*-[rha (1->2 glu)]-7-*O*-rha (K1)



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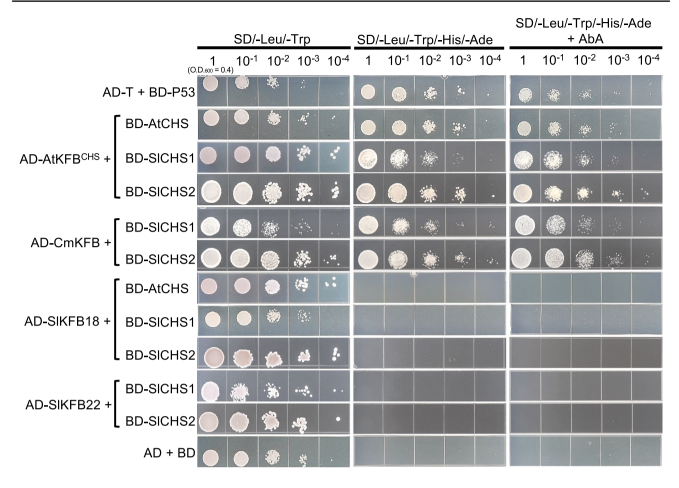


Fig. 5 Y2H for the interaction between CHS and KFB. KFBs were fused with the activation domain (AD), and CHSs were fused with the binding domain (BD). SD media excluding leu, trp, his, and ade were used to assess protein interactions. SD media excluding leu, trp, his, and ade along with Aureobasidin A (AbA) were used to increase stringency of interactions. The pairs of AD-T and DB-P53, and Arabidopsis KFB^{CHS} (AtKFB^{CHS}; AT1G23390) and AtCHS

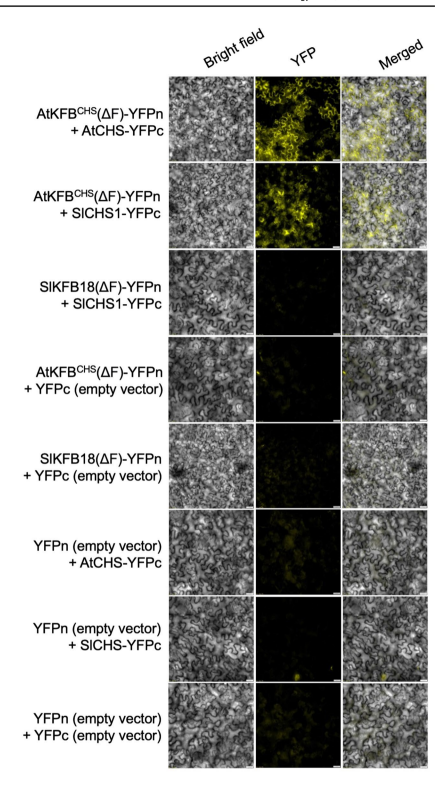
(AT5G13930) were included as positive controls. The pair of AD and BD vectors was used as a negative control. Over-expression of CmKFB (XP 008446188) increased flavonoid production in tomato, but its binding partner has not yet been discovered. Thus, CmKFB was included in the assay. SIKFB18 and SIKFB22 did not interacted with either SICHS1 or SICHS2. But, AtKFBCHS and CmKFB physically interacted with both SICHS1 and SICHS2

generated SIKFB18 overexpression lines (SIKFB18-OX1 and SIKFB18-OX2) and CRISPR-mediated SIKFB18 knock-out lines (SIKFB18^{CR}-1 and SIKFB18^{CR}-2) in tomato Micro-Tom. SIKFB18^{CR}-1 and SIKFB18^{CR}-2 have single base pair deletion mutations at the junction of the F-box domain and the Kelch domain of SIKFB18, which result in a frameshift and premature stop codon, leading to the production of a potential nonfunctional truncated protein that lacks all Kelch domains (Fig. 7a). However, the levels of quercetin-3-O-glucoside-6"-O-rhamnoside (rutin), a major flavonol in tomato, did not alter in the CRISPR knock-out lines (Fig. 7b). Additionally, we did not observe any visible alteration of plant growth and development in the mutants compared to wild type (Fig. 7c). The coloration of the hypocotyls, indicative of anthocyanin accumulation, was observed to be the same as in wild type (Fig. 7c). Although SIKFB18 is the only SIKFB in clade 2 (Fig. 1), we cannot exclude a possibility of functional redundancy. In Arabidopsis, overexpression of AtKFB^{CHS} reduced flavonoid production (Zhang et al. 2017). Thus, we generated tomato transgenic lines overexpressing SIKFB18 driven by the 35S promoter. We isolated ten T1 transgenic lines showing resistance to hygromycin from tissue culture calli and analyzed the expression of SIKFB18, as well as the levels of rutin and chlorogenic acid (Supplementary Fig. S7). However, no statistically significant correlation was observed between the level of SIKFB18 expression and the accumulation of the two phenylpropanoids (Supplementary Fig. S7). We further analyzed T2 progeny from four overexpression lines, which exhibited over a 20 fold higher expression of SIKFB18 compared to the wild-type tomato (Fig. 7d). However, the levels of rutin and chlorogenic acid in the overexpression lines were comparable to those in the wild type and vector controls (Fig. 6e, f). Moreover, their morphology was indistinguishable from the wild



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Fig. 6 The BiFC assay confirmed the interaction between AtKFB^{CHS} and SlCHS1, but there was no interaction between SIKFB18 and SICHS1. The BiFC was conducted with intact AtKFB^{CHS} and SIKFB18 and truncated AtKFBCHS (AtKFB^{CHS} (Δ F)) and SlKFB18 (SIKFB18 (Δ F)) that lack their F-box domains. YFPn-AtKFBCHS (lacking the F-box domain, consisting of amino acids 53-395) and YFPn-SlKFB (lacking the F-box domain, consisting of amino acids 52-370) were coexpressed with YFPc-AtCHS and YFPc-SICHS1 in Nicotiana benthamiana leaves. The images were captured in bright field, YFP (captured using blue excitation light with a center wavelength of 470 nm and a FITC filter), and merged two-channel views under identical exposure conditions. More than five infiltrated leaves were examined, and similar images were obtained



type (Fig. 7g). We used Micro-Tom for the generation of transgenic lines, and the SIKFB18 sequence was retrieved from tomato reference genome (version ITAG4.0), which was generated with Heinz 1706 cultivar (Tomato Genome Consortium 2012). Notably, there was no sequence variation in *SIKFB18* between Micro-Tom (MiBASE database)

and the tomato reference genome (ITAG4.0) (Supplementary Fig. S8). Our biochemical and genetic data suggest that SIKFB18 is unlikely to be involved in flavonoid metabolism.

We tested four SIKFBs that we have shown to regulate PAL stability by using Y2H assays with SICHS1. None of the four SIKFBs interacted with SICHS1, indicating the



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specificity of these SIKFBs in targeting PAL (Supplementary Fig. S9).

Discussion

Protein-protein interactions significantly impact various cellular functions, including protein stability (Struk et al. 2019). Despite the detrimental consequence of destabilized proteins, pinpointing interacting partners is challenging, given the influence of factors such as post-translational modifications and the presence of other molecules (Liddington 2004; Keskin et al. 2008). Phylogenetic and homology analyses are commonly used to infer evolutionary relationships among proteins and identify those with similar functions. This approach proves useful, as demonstrated in our identification of PAL-interacting SIKFBs from the tomato genome (Figs. 1b, 2). However, our homology study did not yield similar results for SICHS-interacting SIKFBs. Despite its high sequence similarity with Arabidopsis KFB^{CHS} and other flavonoid-regulating KFBs from three different plant species, SIKFB18 did not interact with SICHSs (Figs. 1b, 5, 6) or the two flavonoid biosynthesis enzymes, CHI and F3H (Supplementary Fig. S6). Given that overexpression of SIKFB18 did not reduce flavonoid production in tomato, SIKFB18 unlikely functions as negative regulator in flavonoid production (Fig. 7, Supplementary Fig. S7). We also did not find any visible growth and developmental changes or alteration in fertility in either SlKFB18 overexpression lines or knock-out lines. According to expression data from a public database (Supplementary Fig. S10), SIKFB18 expresses in most organs, suggesting that SIKFB18 is not a pseudo gene and likely has functions beyond the regulation of flavonoid metabolism, which remains unknown. Given that both AtKFBCHS and CmKFB physically interacted with SICHS1 and SICHS2 (Fig. 5), SICHS1 and SICHS2 are capable of being recognized by Kelch domaincontaining proteins. SIKFB22, which is closely related to clade 2, and SIKFB13, 14, 21, and 29, which are shown to interact with PAL, did not interact with SICHS1 in our Y2H assay (Fig. 5, Supplementary Fig. S9). In a study with Paeonia, the ring-domain containing protein (PhRING-H2) is responsible for CHS ubiquitination and degradation (Gu et al. 2019). Although, we found no homolog of PhRING-H2 in the tomato genome, it is possible that, in addition to KFBs, other ubiquitination machinery could be involved in the CHS turnover mechanism in tomato.

SIKFB13, despite having the highest sequence homology with PAL-targeting AtKFBs, did not interact with SIPAL5, while SIKFB14, SIKFB21, and SIKFB29 did in both Y2H and BiFC assays (Figs. 1b, 2, 3). However, the overexpression of *SIKFB13* in Arabidopsis resulted in a significant reduction of PAL activity and phenylpropanoid

contents (Fig. 4). Under our growth condition, several Arabidopsis transgenic lines having strong expression of *SIKFB13* or *SIKFB14* exhibited dwarfism and immature siliques (Supplementary Fig. S3b), reminiscent of those observed in the Arabidopsis *pal* mutants (Huang et al. 2010). The presented biochemical and genetic data suggest that these four SIKFBs (13, 14, 21, 29) are affecting PAL stability. When targeted by these SIKFBs, SIPALs likely undergo degradation through a process that requires the F-box domain of the SIKFBs, which is similar to the Arabidopsis PAL ubiquitination and degradation process (Zhang et al. 2013, 2015a).

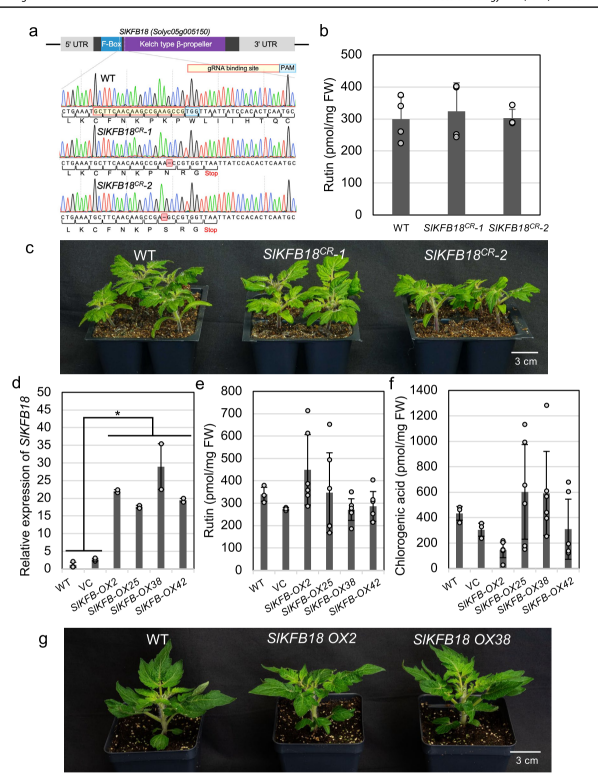
In Arabidopsis, the four Arabidopsis KFB^{PAL}s interact with all four AtPALs redundantly (Zhang et al. 2013, 2015a). Our data suggest that SIKFB13 may interact with tomato PALs, excluding SIPAL5. The four tomato SIKFB proteins could potentially interact with specific PAL enzymes, enabling a more finely tuned regulation of phenylpropanoid flux in specific organs or under particular conditions. Additionally, interactions between SIKFB13 and AtPALs were shown to be relatively weak compared to the interactions between SIKFB14 and AtPALs, although overexpression of both SIKFB13 and SIKFB14 led to the repression of phenylpropanoid biosynthesis (Supplementary Fig. S4, Fig. 4). It is possible that SIKFB13 may interact with additional targets besides PALs that require phenylpropanoid production in the Arabidopsis.

The F-box domain interacts with Skp1, a component protein of the E3 complex, while the kelch domain of KFB serves as the mediator for substrate protein interaction (Schumann et al. 2011). We found that four SlKFBs in clade 1 possessed an F-box domain and three Kelch domains, and both domains of these four SlKFBs are highly conserved when compared with four AtKFB^{PAL}s (Supplementary Fig. S11). Further study is required to understand what feature enables these SlKFBs discern different target proteins, at least for SlPAL5.

The regulation of PAL activity is precise and occurs in response to environmental stimuli and developmental cues, such as the need for the requirement of specific phenylpropanoids with unique in vivo roles or lignin in certain tissues (Edwards et al. 1985; Liang et al. 1989; Dixon and Paiva 1995; Pawlak-Sprada et al. 2011; Kim et al. 2020). Also, the process of phenylpropanoid metabolism channels a significant portion of photosynthetically derived organic carbon to downstream products, with lignin as the primary sink (Novaes et al. 2010). Thus, the stability of PAL, the gateway enzyme for the entire phenylpropanoid production, is imperative for the allocation of available energy and carbon, which is essential for plant survival under unfavorable conditions (Kim et al. 2020). The identification of the SIKFBs targeting PAL suggests the conserved regulation mechanism for phenylpropanoid metabolism in tomato.



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Given that phenylpropanoids confer health benefits to humans, as well as enhance stress tolerance and rigidity in plants, the identified PAL-interacting SIKFBs might be targets to augment phenylpropanoid content in tomato. The expression profiles of these PAL-targeting SIKFBs vary in organs, implying their possible unique roles or

regulations (Supplementary Fig. S10). However, in Arabiodpsis, AtKFB^{PAL}s function redundantly and single knockout mutants of AtKFB^{PAL}s did not affect PAL activity and phenylpropanoid production (Zhang et al. 2013). It has been reported that AtKFB^{PAL}s regulate cytokinin signaling by interacting with the type-B ARR family members,



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∢Fig. 7 Flavonoid contents and morphology of SIKFB18 knock-out mutants and SIKFB18 overexpression lines remained unaltered compared to the wild type. a Two SlKFB18 knockout lines (SlKFB18^{CR}-1 and SIKFB18^{CR}-2) were generated with the CRISPR system and SIKFB18^{CR}-1 and SIKFB18^{CR}-2 have G and A deletion mutations in its exon, respectively. The gRNA targets a region encoding the Kelch domain, as shown in the SIKFB18 gene structure (top). Single base pair deletion mutations in SIKFB18^{CR}-1 and SIKFB18^{CR}-2 lead to premature translation (bottom). **b** The level of quercetin-3-Oglucoside-6"-O-rhamnoside (rutin) in the leaves of three-week-old SIKFB18^{CR}-1 and SIKFB18^{CR}-2 compared with wild type (WT). Error bars represent standard deviation. c Representative images of 3-week-old wild type, SlKFB18^{CR}-1, and SlKFB18^{CR}-2. **d** Expression of SIKFB18 was quantified in wild-type (WT) plants, vector controls (VC), and four T2 transgenic tomato lines overexpressing SlKFB18. Expression levels were measured based on three biological replicates for WT and VC, and two siblings from each transgenic line. T-test analysis results, denoted by asterisks (*), indicate a statistically significant difference between wild type and transgenic lines with a P-value of less than 0.05. e, f Rutin and chlorogenic acid contents were determined for the plants described in (d). Analysis was conducted on metabolite extracts from three-week-old plants. Both WT and VC groups comprised three biological replicates, and for the transgenic lines, three leaves from each of two siblings were sampled, resulting in a total of six replicates for each line. Error bars represent standard deviation. g Representative images of 5 weeks old wild type and SIKFB18 overexpression lines

transcriptional regulators of the cytokinin response (Kim et al. 2013) and TCP14 (Steiner et al. 2021). Whether tomato PAL-interacting KFBs also participate in cytokinin signaling remains unknown.

In this study, we identified 31 KFB-encoding genes in the tomato genome, which is a notably modest number compared to Arabidopsis, harboring over 100 KFBs (Zhang et al 2015a, b). Majority of Arabidopsis KFBs have yet been characterized. The data presented here revealed four SIKFBs functioning in the degradation of PAL. A comprehensive study of the remaining SIKFBs would further broaden our understanding of KFB functions in plants.

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Author contributions D.S. and J.K. designed the experiments and wrote the manuscript. D.S., K.H.C., and E.T. conducted the experiments. C.Y.Y. provided the Y2H systems and guided Y2H experiments. All authors read and agreed with the manuscript.

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Data availability All data supporting the findings of this study are available within the paper and its supplementary information.

Declarations

Competing interests There is no competing interest declared.

References

- Agati G, Brunetti C, Fini A et al (2020) Are flavonoids effective antioxidants in plants? twenty years of our investigation. Antioxidants. https://doi.org/10.3390/antiox9111098
- Anwar R, Fatima T, Mattoo A (2019) Tomatoes: a model crop of solanaceous plants. Environ Sci. https://doi.org/10.1093/acrefore/9780199389414.013.223
- Berger A, Latimer S, Stutts LR et al (2022) Kaempferol as a precursor for ubiquinone (coenzyme Q) biosynthesis: an atypical node between specialized metabolism and primary metabolism. Curr Opin Plant Biol 66:102165. https://doi.org/10.1016/j.pbi.2021. 102165
- Bondonno NP, Dalgaard F, Kyrø C et al (2019) Flavonoid intake is associated with lower mortality in the danish diet cancer and health cohort. Nat Commun 10:3651. https://doi.org/10.1038/s41467-019-11622-x
- Brown DE, Rashotte AM, Murphy AS et al (2001) Flavonoids act as negative regulators of auxin transport in vivo in *Arabidopsis*. Plant Physiol 126:524–535. https://doi.org/10.1104/pp.126.2.524
- Chandra HM, Shanmugaraj BM, Srinivasan B, Ramalingam S (2012) Influence of genotypic variations on antioxidant properties in different fractions of tomato. J Food Sci 77:C1174–C1178. https:// doi.org/10.1111/j.1750-3841.2012.02962.x
- Chapman JM, Muday GK (2021) Flavonols modulate lateral root emergence by scavenging reactive oxygen species in *Arabidopsis thaliana*. J Biol Chem 296:100222. https://doi.org/10.1074/jbc. RA120.014543
- Chen W, Xiao Z, Wang Y et al (2021) Competition between anthocyanin and kaempferol glycosides biosynthesis affects pollen tube growth and seed set of *Malus*. Hortic Res 8:173. https://doi.org/10.1038/s41438-021-00609-9
- Deng Y, Lu S (2017) Biosynthesis and regulation of phenylpropanoids in plants. CRC Crit Rev Plant Sci 36:257–290. https://doi.org/10.1080/07352689.2017.1402852
- Dixon RA, Paiva NL (1995) Stress-Induced Phenylpropanoid Metabolism. Plant Cell 7:1085–1097. https://doi.org/10.1105/tpc.7.7.
- Dong N-Q, Lin H-X (2021) Contribution of phenylpropanoid metabolism to plant development and plant-environment interactions. J Integr Plant Biol 63:180–209. https://doi.org/10.1111/jipb.13054
- Edwards K, Cramer CL, Bolwell GP et al (1985) Rapid transient induction of phenylalanine ammonia-lyase mRNA in elicitor-treated bean cells. Proc Natl Acad Sci 82:6731–6735. https://doi.org/10.1073/pnas.82.20.6731
- España L, Heredia-Guerrero JA, Reina-Pinto JJ et al (2014) Transient silencing of CHALCONE SYNTHASE during fruit ripening modifies tomato epidermal cells and cuticle properties. Plant Physiol 166:1371–1386. https://doi.org/10.1104/pp.114.246405
- Feder A, Burger J, Gao S et al (2015) A kelch domain-containing F-Box coding gene negatively regulates Flavonoid accumulation in muskmelon. Plant Physiol 169:1714–1726. https://doi.org/10.1104/pp.15.01008
- Fernández-Del-Río L, Soubeyrand E, Basset GJ, Clarke CF (2020) Metabolism of the flavonol kaempferol in kidney cells liberates the B-ring to enter coenzyme Q biosynthesis. Molecules. https:// doi.org/10.3390/molecules25132955
- Fernandez-Pozo N, Menda N, Edwards JD et al (2015) The sol genomics network (SGN)–from genotype to phenotype to breeding. Nucleic Acids Res 43:D1036–D1041. https://doi.org/10.1093/nar/gku1195
- Garibay-Hernández A, Kessler N, Józefowicz AM et al (2021) Untargeted metabotyping to study phenylpropanoid diversity in crop plants. Physiol Plant 173:680–697. https://doi.org/10.1111/ppl. 13458



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- Gietz RD, Woods RA (2002) Transformation of yeast by lithium acetate/single-stranded carrier DNA/polyethylene glycol method. Meth Enzymol 350:87–96. https://doi.org/10.1016/S0076-6879(02)50957-5
- Gray WM, Estelle I (2000) Function of the ubiquitin-proteasome pathway in auxin response. Trends Biochem Sci 25:133–138. https://doi.org/10.1016/s0968-0004(00)01544-9
- Grotewold E (2006) The genetics and biochemistry of floral pigments. Annu Rev Plant Biol 57:761–780. https://doi.org/10.1146/annurev.arplant.57.032905.105248
- Gu Z, Men S, Zhu J et al (2019) Chalcone synthase is ubiquitinated and degraded via interactions with a RING-H2 protein in petals of Paeonia "He Xie." J Exp Bot 70:4749–4762. https://doi.org/10.1093/jxb/erz245
- Guo J, Wang M-H (2009) Characterization of the phenylalanine ammonia-lyase gene (SIPAL5) from tomato (Solanum lycopersicum L.). Mol Biol Rep 36:1579–1585. https://doi.org/10.1007/ s11033-008-9354-9
- Gupta S, Van Eck J (2016) Modification of plant regeneration medium decreases the time for recovery of *Solanum lycopersicum* cultivar M82 stable transgenic lines. Plant Cell Tissue Organ Cult 127:417–423. https://doi.org/10.1007/s11240-016-1063-9
- Han J, Ma K, Li H et al (2022) All-in-one: a robust fluorescent fusion protein vector toolbox for protein localization and BiFC analyses in plants. Plant Biotechnol J 20:1098–1109. https://doi.org/10.1111/pbi.13790
- Hristova V, Sun S, Zhang H, Chan DW (2020) Proteomic analysis of degradation ubiquitin signaling by ubiquitin occupancy changes responding to 26S proteasome inhibition. Clin Proteomics 17:2. https://doi.org/10.1186/s12014-020-9265-x
- Huang J, Gu M, Lai Z et al (2010) Functional analysis of the Arabidopsis PAL gene family in plant growth, development, and response to environmental stress. Plant Physiol 153:1526–1538. https://doi.org/10.1104/pp.110.157370
- Keskin O, Gursoy A, Ma B, Nussinov R (2008) Principles of proteinprotein interactions: what are the preferred ways for proteins to interact? Chem Rev 108:1225–1244. https://doi.org/10.1021/ cr040409x
- Kim HJ, Chiang Y-H, Kieber JJ, Schaller GE (2013) SCF(KMD) controls cytokinin signaling by regulating the degradation of type-B response regulators. Proc Natl Acad Sci USA 110:10028–10033. https://doi.org/10.1073/pnas.1300403110
- Kim JI, Dolan WL, Anderson NA, Chapple C (2015) Indole glucosinolate Biosynthesis Limits phenylpropanoid Accumulation in Arabidopsis thaliana. Plant Cell 27:1529–1546. https://doi.org/ 10.1105/tpc.15.00127
- Kim JI, Zhang X, Pascuzzi PE et al (2020) Glucosinolate and phenylpropanoid biosynthesis are linked by proteasome-dependent degradation of PAL. New Phytol 225:154–168. https://doi.org/10.1111/nph.16108
- Kong D, Li S, Smolke CD (2020) Discovery of a previously unknown biosynthetic capacity of naringenin chalcone synthase by heterologous expression of a tomato gene cluster in yeast. Sci Adv. https://doi.org/10.1126/sciadv.abd1143
- Li Y, Kim JI, Pysh L, Chapple C (2015) Four isoforms of arabidopsis 4-coumarate: CoA ligase Have overlapping yet distinct roles in phenylpropanoid metabolism. Plant Physiol 169:2409–2421. https://doi.org/10.1104/pp.15.00838
- Liang XW, Dron M, Cramer CL et al (1989) Differential regulation of phenylalanine ammonia-lyase genes during plant development and by environmental cues. J Biol Chem 264:14486–14492
- Liddington RC (2004) Structural basis of protein-protein interactions. Methods Mol Biol 261:3-14. https://doi.org/10.1385/1-59259-762-9:003

- Liu J, Osbourn A, Ma P (2015) MYB transcription factors as regulators of phenylpropanoid metabolism in plants. Mol Plant 8:689–708. https://doi.org/10.1016/j.molp.2015.03.012
- Løvdal T, Olsen KM, Slimestad R et al (2010) Synergetic effects of nitrogen depletion, temperature, and light on the content of phenolic compounds and gene expression in leaves of tomato. Phytochemistry 71:605–613. https://doi.org/10.1016/j.phytochem. 2009.12.014
- Mao W, Han Y, Chen Y et al (2022) Low temperature inhibits anthocyanin accumulation in strawberry fruit by activating FvMAPK3-induced phosphorylation of FvMYB10 and degradation of chalcone synthase 1. Plant Cell 34:1226–1249. https://doi.org/10.1093/plcell/koac006
- Micek A, Godos J, Del Rio D et al (2021) Dietary flavonoids and cardiovascular disease: a comprehensive dose-response meta-analysis. Mol Nutr Food Res 65:e2001019. https://doi.org/10.1002/mnfr.202001019
- Muhlemann JK, Younts TLB, Muday GK (2018) Flavonols control pollen tube growth and integrity by regulating ROS homeostasis during high-temperature stress. Proc Natl Acad Sci USA 115:E11188–E11197. https://doi.org/10.1073/pnas.1811492115
- Muro-Villanueva F, Mao X, Chapple C (2019) Linking phenylpropanoid metabolism, lignin deposition, and plant growth inhibition. Curr Opin Biotechnol 56:202–208. https://doi.org/10.1016/j.copbio.2018.12.008
- Nakabayashi K, Bartsch M, Ding J, Soppe WJJ (2015) Seed dormancy in arabidopsis requires self-binding ability of DOG1 protein and the presence of multiple isoforms generated by alternative splicing. PLoS Genet 11:e1005737. https://doi.org/10.1371/journal. pgen.1005737
- Novaes E, Kirst M, Chiang V et al (2010) Lignin and biomass: a negative correlation for wood formation and lignin content in trees. Plant Physiol 154:555–561. https://doi.org/10.1104/pp.110.161281
- Ohno S, Hori W, Hosokawa M et al (2018) Post-transcriptional silencing of chalcone synthase is involved in phenotypic lability in petals and leaves of bicolor dahlia (Dahlia variabilis) "Yuino." Planta 247:413–428. https://doi.org/10.1007/s00425-017-2796-3
- Pawlak-Sprada S, Arasimowicz-Jelonek M, Podgórska M, Deckert J (2011) Activation of phenylpropanoid pathway in legume plants exposed to heavy metals. Part I. Effects of cadmium and lead on phenylalanine ammonia-lyase gene expression, enzyme activity and lignin content. Acta Biochim Pol 58:211–216
- Perez VC, Dai R, Bai B et al (2021) Aldoximes are precursors of auxins in arabidopsis and maize. New Phytol 231:1449–1461. https://doi.org/10.1111/nph.17447
- Potter SC, Luciani A, Eddy SR et al (2018) HMMER web server: 2018 update. Nucleic Acids Res 46:W200–W204. https://doi.org/10.1093/nar/gky448
- Prasanna P, Upadhyay A (2021) Flavonoid-based nanomedicines in alzheimer's disease therapeutics: promises made, a long way to go. ACS Pharmacol Transl Sci 4:74–95. https://doi.org/10.1021/ acsptsci.0c00224
- Rohde A, Morreel K, Ralph J et al (2004) Molecular phenotyping of the pal₁ and pal₂ mutants of *arabidopsis thaliana* reveals far-reaching consequences on phenylpropanoid, amino acid, and carbohydrate metabolism. Plant Cell 16:2749–2771. https://doi.org/10.1105/tpc. 104.023705
- Rosa-Martínez E, Bovy A, Plazas M et al (2023) Genetics and breeding of phenolic content in tomato, eggplant and pepper fruits. Front Plant Sci 14:1135237. https://doi.org/10.3389/fpls.2023.1135237
- Ruegger M, Chapple C (2001) Mutations that reduce sinapoylmalate accumulation in *arabidopsis thaliana* define loci with diverse roles in phenylpropanoid metabolism. Genetics 159:1741–1749. https://doi.org/10.1093/genetics/159.4.1741



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Ruprecht C, Proost S, Hernandez-Coronado M et al (2017) Phylogenomic analysis of gene co-expression networks reveals the evolution of functional modules. Plant J 90:447–465. https://doi.org/10.1111/tpj.13502

- Saito K, Yonekura-Sakakibara K, Nakabayashi R et al (2013) The flavonoid biosynthetic pathway in *Arabidopsis*: structural and genetic diversity. Plant Physiol Biochem 72:21–34. https://doi.org/10.1016/j.plaphy.2013.02.001
- Schijlen EGWM, de Vos CHR, Martens S et al (2007) RNA interference silencing of chalcone synthase, the first step in the flavonoid biosynthesis pathway, leads to parthenocarpic tomato fruits. Plant Physiol 144:1520–1530. https://doi.org/10.1104/pp.107.100305
- Schmelzer E, Jahnen W, Hahlbrock K (1988) In situ localization of light-induced chalcone synthase mRNA, chalcone synthase, and flavonoid end products in epidermal cells of parsley leaves. Proc Natl Acad Sci USA 85:2989–2993. https://doi.org/10.1073/pnas.85.9.2989
- Schumann N, Navarro-Quezada A, Ullrich K et al (2011) Molecular evolution and selection patterns of plant F-box proteins with C-terminal kelch repeats. Plant Physiol 155:835–850. https://doi.org/10.1104/pp.110.166579
- Shao T, Qian Q, Tang D et al (2012) A novel gene IBF1 is required for the inhibition of brown pigment deposition in rice hull furrows. Theor Appl Genet 125:381–390. https://doi.org/10.1007/s00122-012-1840-8
- Shin DH, Cho M, Choi MG et al (2015) Identification of genes that may regulate the expression of the transcription factor production of anthocyanin pigment 1 (PAP1)/MYB75 involved in Arabidopsis anthocyanin biosynthesis. Plant Cell Rep 34:805–815. https://doi.org/10.1007/s00299-015-1743-7
- Shin D, Perez VC, Dickinson GK et al (2023) Altered methionine metabolism impacts phenylpropanoid production and plant development in *Arabidopsis thaliana*. BioRxiv. https://doi.org/10.1101/2023.05.29.542770
- Shirley BW, Kubasek WL, Storz G et al (1995) Analysis of Arabidopsis mutants deficient in flavonoid biosynthesis. Plant J 8:659–671. https://doi.org/10.1046/j.1365-313x.1995.08050
- Shomali A, Das S, Arif N et al (2022) Diverse physiological roles of flavonoids in plant environmental stress responses and tolerance. Plants. https://doi.org/10.3390/plants11223158
- Slika H, Mansour H, Wehbe N et al (2022) Therapeutic potential of flavonoids in cancer: ROS-mediated mechanisms. Biomed Pharmacother 146:112442. https://doi.org/10.1016/j.biopha.2021.112442
- Soubeyrand E, Johnson TS, Latimer S et al (2018) The peroxidative cleavage of kaempferol contributes to the biosynthesis of the benzenoid moiety of ubiquinone in plants. Plant Cell 30:2910–2921. https://doi.org/10.1105/tpc.18.00688
- Soubeyrand E, Latimer S, Bernert AC et al (2021) 3-O-glycosylation of kaempferol restricts the supply of the benzenoid precursor of ubiquinone (coenzyme Q) in *Arabidopsis thaliana*. Phytochemistry 186:112738. https://doi.org/10.1016/j.phytochem.2021.112738
- Steiner E, Triana MR, Kubasi S et al (2021) KISS ME DEADLY F-box proteins modulate cytokinin responses by targeting the transcription factor TCP14 for degradation. Plant Physiol 185:1495–1499. https://doi.org/10.1093/plphys/kiab033
- Struk S, Jacobs A, Sánchez Martín-Fontecha E et al (2019) Exploring the protein-protein interaction landscape in plants. Plant Cell Environ 42:387–409. https://doi.org/10.1111/pce.13433
- Sun C, Deng L, Du M et al (2020) A transcriptional network promotes anthocyanin biosynthesis in tomato flesh. Mol Plant 13:42–58. https://doi.org/10.1016/j.molp.2019.10.010
- Tan H, Man C, Xie Y et al (2019) A crucial role of GA-regulated flavonol biosynthesis in root growth of *Arabidopsis*. Mol Plant 12:521–537. https://doi.org/10.1016/j.molp.2018.12.021

- Teale WD, Pasternak T, Dal Bosco C et al (2021) Flavonol-mediated stabilization of PIN efflux complexes regulates polar auxin transport. EMBO J 40:e104416
- Tohge T, de Souza LP, Fernie AR (2017) Current understanding of the pathways of flavonoid biosynthesis in model and crop plants. J Exp Bot 68:4013–4028. https://doi.org/10.1093/jxb/erx177
- Tomato Genome Consortium (2012) The tomato genome sequence provides insights into fleshy fruit evolution. Nature 485:635–641. https://doi.org/10.1038/nature11119
- Tu S-H, Chen L-C, Ho Y-S (2017) An apple a day to prevent cancer formation: reducing cancer risk with flavonoids. J Food Drug Anal 25:119–124. https://doi.org/10.1016/j.jfda.2016.10.016
- Verweij W, Spelt CE, Bliek M et al (2016) Functionally similar WRKY proteins regulate vacuolar acidification in petunia and hair development in arabidopsis. Plant Cell 28:786–803. https://doi.org/10.1105/tpc.15.00608
- Vogt T (2010) Phenylpropanoid biosynthesis. Mol Plant 3:2–20. https://doi.org/10.1093/mp/ssp106
- Wang Y, Liu W, Wang X et al (2020) MiR156 regulates anthocyanin biosynthesis through SPL targets and other microRNAs in poplar. Hortic Res 7:118. https://doi.org/10.1038/s41438-020-00341-w
- Wedick NM, Pan A, Cassidy A et al (2012) Dietary flavonoid intakes and risk of type 2 diabetes in US men and women. Am J Clin Nutr 95:925–933. https://doi.org/10.3945/ajcn.111.028894
- Wen W, Alseekh S, Fernie AR (2020) Conservation and diversification of flavonoid metabolism in the plant kingdom. Curr Opin Plant Biol 55:100–108. https://doi.org/10.1016/j.pbi.2020.04.004
- Wen K, Fang X, Yang J et al (2021) Recent research on flavonoids and their biomedical applications. Curr Med Chem 28:1042–1066. https://doi.org/10.2174/0929867327666200713184138
- Xian D, Guo M, Xu J et al (2021) Current evidence to support the therapeutic potential of flavonoids in oxidative stress-related dermatoses. Redox Rep 26:134–146. https://doi.org/10.1080/13510 002.2021.1962094
- Xu W, Dubos C, Lepiniec L (2015) Transcriptional control of flavonoid biosynthesis by MYB-bHLH-WDR complexes. Trends Plant Sci 20:176–185. https://doi.org/10.1016/j.tplants.2014.12.001
- Yin R, Messner B, Faus-Kessler T et al (2012) Feedback inhibition of the general phenylpropanoid and flavonol biosynthetic pathways upon a compromised flavonol-3-O-glycosylation. J Exp Bot 63:2465–2478. https://doi.org/10.1093/jxb/err416
- Yin R, Han K, Heller W et al (2014) Kaempferol 3-O-rhamnoside-7-O-rhamnoside is an endogenous flavonol inhibitor of polar auxin transport in *Arabidopsis* shoots. New Phytol 201:466–475. https://doi.org/10.1111/nph.12558
- Zhang X, Liu C-J (2015) Multifaceted regulations of gateway enzyme phenylalanine ammonia-lyase in the biosynthesis of phenylpropanoids. Mol Plant 8:17–27. https://doi.org/10.1016/j.molp.2014. 11.001
- Zhang X, Henriques R, Lin S-S et al (2006) Agrobacterium-mediated transformation of *Arabidopsis thaliana* using the floral dip method. Nat Protoc 1:641–646. https://doi.org/10.1038/nprot. 2006.97
- Zhang X, Gou M, Liu C-J (2013) Arabidopsis kelch repeat F-box proteins regulate phenylpropanoid biosynthesis via controlling the turnover of phenylalanine ammonia-lyase. Plant Cell 25:4994–5010. https://doi.org/10.1105/tpc.113.119644
- Zhang X, Gou M, Guo C et al (2015a) Down-regulation of kelch domain-containing F-box protein in *Arabidopsis* enhances the production of (poly)phenols and tolerance to ultraviolet radiation. Plant Physiol 167:337–350. https://doi.org/10.1104/pp.114. 249136
- Zhang Y, Butelli E, Alseekh S et al (2015b) Multi-level engineering facilitates the production of phenylpropanoid compounds in tomato. Nat Commun 6:8635. https://doi.org/10.1038/ncomms9635



- Zhang X, Abrahan C, Colquhoun TA, Liu C-J (2017) A proteolytic regulator controlling chalcone synthase stability and flavonoid biosynthesis in arabidopsis. Plant Cell 29:1157–1174. https://doi.org/10.1105/tpc.16.00855
- Zhang D, Song YH, Dai R et al (2020) Aldoxime metabolism is linked to phenylpropanoid production in *Camelina sativa*. Front Plant Sci 11:17. https://doi.org/10.3389/fpls.2020.00017
- Zhao T, Huang C, Li S et al (2023) VviKFB07 F-box E3 ubiquitin ligase promotes stilbene accumulation by ubiquitinating and degrading VviCHSs protein in grape. Plant Sci 331:111687. https://doi.org/10.1016/j.plantsci.2023.111687

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