UC Davis UC Davis Previously Published Works

Title

A pear S1-bZIP transcription factor PpbZIP44 modulates carbohydrate metabolism, amino acid, and flavonoid accumulation in fruits.

Permalink https://escholarship.org/uc/item/9bg2j65c

Journal Horticulture Research, 10(8)

ISSN 2662-6810

Authors

Wang, Hong Xu, Kexin Li, Xiaogang <u>et al.</u>

Publication Date 2023-08-01

DOI

10.1093/hr/uhad140

Peer reviewed

Article

A pear S1-bZIP transcription factor PpbZIP44 modulates carbohydrate metabolism, amino acid, and flavonoid accumulation in fruits

Hong Wang $1^{,2}$, Kexin Xu^{1,2}, Xiaogang Li², Bárbara Blanco-Ulate³, Qingsong Yang², Gaifang Yao⁴, Yiduo Wei³, Jun Wu $1^{,2}$, Baolong Sheng², Youhong Chang², Cai-Zhong Jiang $3^{,5,*}$ and Jing Lin^{1,2,*}

¹College of Horticulture, Nanjing Agricultural University, Nanjing 210014, China

²Institute of Pomology, Jiangsu Academy of Agricultural Sciences/Jiangsu Key Laboratory for Horticultural Crop Genetic Improvement, Nanjing 210014, China ³Department of Plant Sciences, University of California, Davis, Davis, CA 95616, USA

⁴School of Food and Biological Engineering, Hefei University of Technology, Hefei 230009, China

⁵Crops Pathology and Genetics Research Unit, United States Department of Agriculture, Agricultural Research Service, Davis, California, 95616, USA *Corresponding authors: Email: cjiang@ucdavis.edu, lj84390224@126.com

Abstract

Fruit quality is defined by attributes that give value to a commodity. Flavor, texture, nutrition, and shelf life are key quality traits that ensure market value and consumer acceptance. In pear fruit, soluble sugars, organic acids, amino acids, and total flavonoids contribute to flavor and overall quality. Transcription factors (TFs) regulate the accumulation of these metabolites during development or in response to the environment. Here, we report a novel TF, *PpbZIP44*, as a positive regulator of primary and secondary metabolism in pear fruit. Analysis of the transient overexpression or RNAi-transformed pear fruits and stable transgenic tomato fruits under the control of the fruit-specific E8 promoter demonstrated that *PpZIP44* substantially affected the contents of soluble sugar, organic acids, amino acids, and flavonoids. In *E8::PpbZIP44* tomato fruit, genes involved in carbohydrate metabolism, amino acid, and flavonoids biosynthesis were significantly induced. Furthermore, in *PpbZIP44* overexpression or antisense pear fruits, the expression of genes in the related pathways was significantly impacted. *PpbZIP44* directly interacted with the promoter of *PpSDH9* and *PpPrOH1* to induce their expression, thereby depleting sorbitol and proline, decreasing citrate and malate, and enhancing fructose contents. *PpbZIP44* also directly bound to the *PpADT* and *PpF3H* promoters, which led to the carbon flux toward phenylalanine metabolites and enhanced phenylalanine and flavonoid contents. These findings demonstrate that *PpbZIP44* mediates multimetabolism reprogramming by regulating the gene expression related to fruit quality compounds.

Introduction

Fruit quality influences consumer preference and market competitiveness. Therefore, improving fruit quality has been the cornerstone for breeders and scientists. Fruit quality includes taste, aroma, texture, and nutritional quality, in addition to shelf life and other visual attributes [1]. The content and species of soluble sugars and organic acids are the fundamental compounds for fruit growth, development, and ripening and determine the fruit flavor at the edible stage [2, 3]. Fatty acids, amino acids and vitamins, and secondary metabolites such as phenylpropanoids, flavonoids, and phenolics are vital factors in determining the health-promoting quality of fruits [1, 4, 5]. As with most complex agronomic traits, fruit quality is affected by primary and secondary metabolic shifts [6]. In the Rosaceae species, sorbitol and sucrose are the primary end products of photosynthesis, synthesized in leaves and transported to fruits through the phloem [7]. Then, sucrose is resynthesized via sucrose phosphate synthase (SPS) or converted to fructose and glucose by invertases (INV) or sucrose synthase (SUS) [2, 8, 9]. Sorbitol is either accumulated or converted mainly to fructose by sorbitol dehydrogenase (SDH) in fruits [7]. Unlike soluble sugar metabolites, the organic acids accumulated in the fruit mainly depend on the de novo synthesis in fruit cells [3]. Malate, an intermediate product of the tricarboxylic acid (TCA) cycle, is the most abundant organic acid determining flavor quality in pear [3]. Organic acids accumulate in young fruits to generate cellular energy for respiratory and intermediates for the biosynthesis pathway, then gradually decrease and achieve a palatable sugar: acid ratio, which is a significant indicator of fruit quality [3]. Amino acids are derived from different intermediates of glycolysis and the TCA cycle, and aromatic amino acids (AAA) are synthesized from the shikimate pathway [10]. Amino acids are essential for maintaining fruit quality and play positive roles in human health [10]. Flavonoids, as the most abundant phenolic substance in plants, regulate various plant biological processes, including fruit coloration, resistance to UV-B damage and defense against biotic and abiotic stress, and also possess a wide range of physiological features that benefit human health, including radical scavenging, anti-inflammation and anti-aging [11].

Received: 26 March 2023; Accepted: 8 July 2023; Published: 21 July 2023; Corrected and Typeset: 11 August 2023 © The Author(s) 2023. Published by Oxford University Press on behalf of Nanjing Agricultural University. This is an Open Access article distributed under the terms of the Creative Commons Attribution License (https://creativecommons.org/licenses/by/4.0/), which permits unrestricted reuse, distribution, and reproduction in any medium, provided the original work is properly cited. Elucidation of the underlying metabolic regulation responsible for changes in fruit attributes is essential for providing scientific guidance for improvement of fruit quality.

Biosynthesis and accumulation of primary and secondary metabolites are regulated by transcription factors (TFs) [12-14]. TFs can activate the collaborative expression of multiple genes, thereby effectively regulating the reprogramming of primary or secondary metabolic pathways [13]. Evidences suggest that one subgroup of the basic leucine zipper (bZIP) transcription factors (S1-bZIP group) mediates sugar-related regulatory gene expression and are required for sugar signaling and plays important roles in the regulation of fruit quality and stress response [15]. Conserved upstream open reading frames (uORFs) in the 5'untranslated regions of the S1-bZIP group genes encode a sucrose control peptide associated with sucrose-induced translation repression [16]. Genome editing FvebZIPs1.1 uORF resulted in a diverse amount increase in sugar content in strawberries [17]. Compared with the wild type (WT), overexpression of strawberry FubZIP11 in tomatoes significantly increased the total soluble solids and the contents of soluble sugars [18]. Transgenic lines with overexpressing tomato SlbZIP1 and SlbZIP2 contained 1.5-fold higher sucrose/glucose/fructose than WT control, and increased levels of several amino acids [19]. These findings suggest that the S1-bZIP TFs have a broad role in sugar accumulation and regulating carbon-nitrogen balance. However, whether or how pear S1-bZIP TFs regulate primary and secondary metabolic shifts in fruit remains unclear.

The discovery and identification of novel TFs are vital for successfully improving fruit quality, especially in woody plants that generate commercially superior fruits with better palatability, nutrient balance, and health benefits [1]. Pear is a popular fruit worldwide. For several years, there has been increased interest in the consumption of pear fruits. Hence, cultivation areas have been increasing, production has intensified, and new cultivars have been developed [20]. The great demand for pear fruit is associated with its taste, texture, and nutritional value. Enhancing fruit quality through metabolite manipulation or breeding has been an essential in increasing Asian pear quality. We have recently developed the 'Sucui 1' [Pyrus pyrifolia (Burm.f.) Nakai.,] variety by crossing 'Huasu' and 'Cuiguan'. 'Sucui 1' fruits have improved quality and extrinsic features and an extremely short growth period, and could be used as an excellent material for studying the mechanism of fruit quality or other agronomy traits in pears. In this study, we identified a critical regulator, a member of the S1-bZIP group PpbZIP44, that affected fructose accumulation, organic acids contents, amino acid, and flavonoid levels. Our finding provides new insights into understanding the regulatory mechanism for future metabolic engineering to improve fruit quality.

Results Identification of *PpbZIP44* and its expression patterns during development and ripening in pear fruit

To gain insight into the structural characteristics of the S1-bZIP TFs, we performed a phylogenic analysis using 105 pear bZIP TFs from the 'Cuiguan' pear genome database [21] and their orthologs in Arabidopsis thaliana, Solanum lycopersicum, Musa nana, Fragaria ananassa, Vitis vinifera, Oryza sativa, and Malus pumila. The phylogenetic analysis assigned 11 PpbZIPs, which were noted as red dots to S1-bZIP group (Fig. S1A and Table S2). We also constructed a comprehensive phylogenetic tree using pear

S1-bZIP proteins and S1-bZIP TFs from other species (Fig. S1B and Table S2). Pairwise protein sequence alignment found that 3 of 11 PpbZIPs, such as EVM0005411, EVM0009313, and EVM0014055, were identified as orthologs of AtbZIP11, whereas EVM0014652, EVM0000485, and EVM0042575 have the highest homology with AtbZIP44 (Fig. S1B and Table S2). In addition, EVM0041187 and EVM0041928 were identified as orthologs of AtbZIP53. To perform gene expression analysis, we collected samples from leaves, roots, and fruits, including seven developmental stages (15, 30, 45, 60, 75, 90, and 93) and one ripening stage (100 DAP) of 'Sucui 1' fruits (Fig. 1A). Our previous RNA-seq data showed that the significantly high transcription abundance of an S1bZIP gene EVM0042575 (PpbZIP44) was observed at stage 5 (S5) among the top 20 highest expressed TFs (Fig. 1B). To further examine the expression dynamics of several candidate S1-bZIP genes, we performed the quantitative real-time polymerase chain reaction (qRT-PCR) analysis. Compared with other genes, PpbZIP44 (EVM0042575) displayed much higher transcript abundances in fruits (Fig. 1C and Fig. S2). Especially, the expression level of PpbZIP44 was the highest at S5 and 3.5-fold higher than that at stage 4 (S4). A higher expression level of PpbZIP44 was also found at stage 6 (S6) and the ripening stage 8 (S8).

Overexpression or silencing of PpbZIP44 modulated sugar: Acid ratio in pear fruits

We cloned the *PpbZIP44* gene from the sweet pear variety 'Sucui 1' with early maturation. The 474-bp open reading frame (ORF) of *PpbZIP44* encoded a protein of 157 amino acids with one conserved bZIP domain (Fig. 2A). To examine the biological function of *PpbZIP44* in fruit, we first carried out the transient overexpression and silencing assays in pear fruit. Fruits were collected a week before ripening and injected with the transient overexpression-, RNAi-, and empty-vector (Fig. 2B). At 7 days after injection, the expression levels of *PpbZIP44* were 59-fold higher in the OE-bZIP44 and 0.44-fold lower in the RNAi-bZIP44 fruit, when compared with that in the empty vector control (Fig. 2C).

The total soluble sugar displayed a significantly higher level in OE-PpbZIP44 and lower in RNAi-PpbZIP44 than in fruits with the empty vector (Fig. 2D). Similarly, the titratable acid content was significantly reduced in OE-PpbZIP44 but increased in RNAi-PpbZIP44 compared with the control (Fig. 2E). The sugar:acid ratio in OE-bZIP44 was 52.67, higher than 45.08 in WT and 42.43 in RNAi-bZIP44 (Fig. 2F). The fructose and sorbitol contents significantly differed between the transgenic fruits and the empty vector control. Specifically, compared with the empty vector control, the fructose contents increased significantly by 38.17% in OE-bZIP44, whereas they decreased significantly by 31.45% in RNAi-bZIP44. The sorbitol contents decreased by 26.42% in OE-bZIP44, whereas increasing significantly by 72.09% in RNAi-bZIP44. Slight changes in glucose and sucrose contents were observed on the transiently transformed fruits compared with control ones (Fig. 2G). All organic acids contents were significantly increased in RNAi-bZIP44 (Fig. 2H), indicating a role for PpbZIP44 in the TCA cycle. In agreement, the organic acids, malate, and citrate were decreased by 29.96% and 40.05%, respectively, in OE-bZIP44 compared with the empty vector control

SDH is the essential enzyme in converting sorbitol to fructose. We evaluated the PpSDH enzymatic activity in transiently transformed fruits and observed that its activity was increased approximately 1.3 times in OE-bZIP44 fruits compared with the empty vector control. In the RNAi-bZIP44, PpSDH enzyme activity was decreased by 0.6 times (Fig. 21).

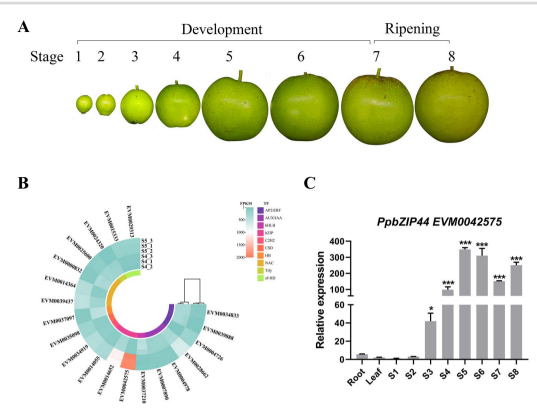


Figure 1. Fruit development stages and transcription abundance of identified genes. A. Eight fruit developmental and ripening stages were selected for analysis in pear. B. Heatmap hierarchical clustering showing top 20 TFs and FPKM value of S1-bZIP members at fruit development stages 4 and 5 based on transcriptome data in 'Sucui 1'. Colors represented different TFs types. C. Time course expression profiles of *PpbZIP44* in the root, leaf, fruit development, and ripening processes of 'Sucui 1' using qRT-PCR. Sarcocarp samples with moving peels measuring 0.4-cm thick were collected every 15 days after full bloom. Three independent biological replicates containing at least five fruits for every experiment were performed for each stage. Expression levels were normalized to the levels of pear GAPDH. The asterisks indicated values determined by the Dunnett t test to significantly differ from stage 1 (*P < 0.05, ***P < 0.001).

Effects of *PpbZIP44* on amino acid contents in transiently transformed pear fruits

We further analyzed the contents of amino acids in transiently transformed pear fruits with OE-bZIP44 and RNAi-bZIP44 constructs. The total amino acid contents of OE-bZIP44 and RNAi-bZIP44 fruits were 18.9% higher and 8.5% lower than those of fruits with the empty vector. Met and Tyr levels were approximately 2-fold higher in OE-bZIP44 than in the empty vector control. Phe, Asp, and Ala levels were 1.1- to 1.2-fold higher in OE-bZIP44 and lower in RNAi-bZIP44 compared with empty vector fruits. Val, Ser, and Pro levels were lower in OE-bZIP44 but higher in RNAi-bZIP44. The levels of Glu, Ile, and NH3 were similar in the fruits with the empty vector as that of OE-bZIP44 or RNAi-bZIP44 (Table 1).

Fruit-specific overexpression of *PpbZIP44* resulted in metabolites reprogramming in tomato fruits

To further verify the function of PpbZIP44, we attempted to overexpress the PpbZIP44 in tomatoes due to the significant challenge in pear transformation and unrealistic fruit-forming in wood plants with limited time. The tomato fruit-specific E8 promoter was used to drive PpbZIP44 expression in tomato fruit. Eight T0 lines were obtained. These lines grew and developed normally (Fig. 3A). The expression of PpbZIP44 in the transgenic lines was monitored by qRT-PCR. E8::PpbZIP44–2 (OE-2) and E8::PpbZIP44–3 (OE-3), which had higher PpbZIP44 expression in the ripening fruit compared with the WT control (Fig. 3B), were then chosen to obtain a T1 generation for further investigation.

To better characterize the effect of PpbZIP44 on the contents of compounds, we performed widely targeted metabolomics of E8::PpbZIP44–2 and WT tomato fruits using multiple reaction monitoring methods (MRM) [22]. We identified 613 metabolic compounds across all samples (Supplementary Table S3). Among these, 108 were classified as differential expression metabolites (DEMs) based on two thresholds [variable importance in projection (VIP) ≥ 1 and fold-change ≥ 2 and ≤ 0.5] when comparing the E8::PpbZIP44 lines against the WT control (Fig. 3C). Kyoto encyclopedia of genes and genomes (KEGG) enrichment analyses in the DEMs showed that the top overrepresented pathways included lipid metabolism (e.g. biosynthesis of unsaturated fatty acid, linoleic acid, and alpha-linolenic acid metabolism), biosynthesis of other secondary metabolism (e.g. isoflavonoid/flavonoid/flavone and flavonol biosynthesis), amino acid metabolism, and carbohydrate metabolism (e.g. the TCA cycle) (Fig. 3D). Most DEMs in linoleic acid biosynthesis, flavonoidlike biosynthesis, tyrosine metabolism, and alanine, aspartate, and glutamate metabolism were upregulated in the E8::PpbZIP44 tomato fruits (Fig. 3D).

The changes of the corresponding metabolites in these pathways induced by *PpbZIP44* were further analyzed. The results of widely targeted metabolomics showed that the levels of sucrose and fructose increased >1.1-fold in transgenic fruits compared with WT fruits.

PpbZIP44 significantly induced the threose level to ~2fold compared with WT fruits (Supplementary Table S4). The levels of sorbitol, glucose 6-phosphate, fructose 6-Phosphate, fructose-1,6-biphosphate, D-sedoheptuiose 7-phosphate, and

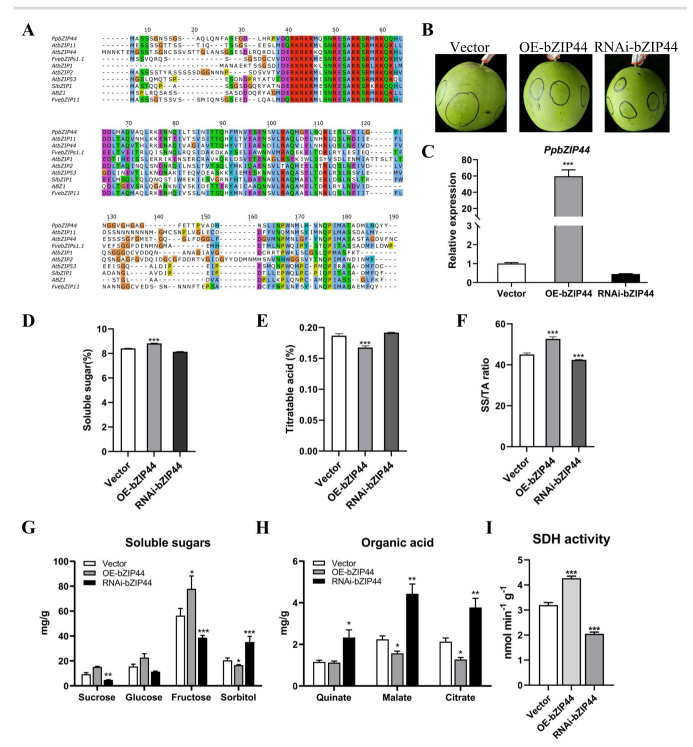


Figure 2. Transient transformation of *PpbZIP44* in pear fruits. A. Multiple protein sequence alignments of *PZIP44* and other S1-bZIP proteins in different species. B. Transient assays in 'Sucui 1' pear using overexpression (OE-) or RNAi (RNAi) or empty vector (Vector) of *PpbZIP44*. C. The expression levels of *PpbZIP44* in the fleshy tissue around the infiltration sites of transformed pear fruits using qRT-PCR. The content of total SS (D) and TA (E), SS/TA ratio (F), individual soluble sugars contents (G), individual organic acids contents (H), and SDH activity (I) in transformed pear fruits and control pear fruits. Bars represented the mean value \pm SE ($n \ge 3$). The asterisks indicated values that were determined by the Dunnett t test to significantly differ from their empty vector or WT control (*P < 0.05, **P < 0.01, ***P < 0.001).

3-phospho-D-glyceric acid were decreased; however, the levels of glucuronic-6,3-lactone, 6-phosphoglucose acid, ribulose-5phosphate, and phosphoenolpyruvate were >1.5-fold increase in the corresponding transgenic fruits (Supplementary Table S4). Overexpression of *PpbZIP44* decreased the TCA cycle intermediate contents (e.g. fumaric acid, citric acid, malic acid, citraconic acid, 3-Guanidinopropionic acid, α -Ketoglutaric acid) (Supplementary Table S4) and significantly increased the level of succinic acid and amino malonic acid (Supplementary Table S4).

Sugar and organic acids were further measured in transgenic tomato fruits. The total soluble sugar content (SS) was increased

 Table 1. Amino acid composition in PpbZIP44 transient transformed pear fruits

A	P	OF LEDAA	
Amino acids	Empty vector	OE-bZIP44	RNAi-bZIP44
Asp	244.15 ± 10.13	275.94 ± 13.08	187.62±14.53**
Thr	144.87 ± 7.13	$193.82 \pm 5.49^*$	170.18 ± 29.43
Phe	178.20 ± 2.96	$209.33 \pm 4.96^{***}$	$148.10 \pm 3.70^{***}$
Met	109.87 ± 4.76	$198.45 \pm 3.46^{***}$	$96.64 \pm 1.94^{**}$
Glu	54.71 ± 0.65	55.29 ± 3.32	$44.50 \pm 1.46^{**}$
Ala	42.17 ± 2.80	46.03 ± 1.99	38.04 ± 2.80
Cys	51.30 ± 6.72	45.99 ± 3.80	54.87 ± 1.05
Val	54.46 ± 5.44	$44.60 \pm 0.07^{*}$	55.39 ± 1.98
Ser	39.35 ± 2.11	$34.13 \pm 2.13^{*}$	43.11 ± 1.46
Tyr	27.50 ± 1.10	47.10±1.35***	$17.94 \pm 2.04^{***}$
Ile	24.10 ± 2.62	25.00 ± 1.73	23.59 ± 0.08
Pro	34.08 ± 2.60	12.70±0.36***	$39.87 \pm 1.52^*$
NH3	7.22 ± 0.51	8.79 ± 0.46	6.39 ± 1.00
Total	1011.98	1279.35	926.24

Note: Amino acid levels were given as $\mu g \cdot g^{-1}$ fresh weight of pear fruits (±SE). All measurements were performed in triplicate. Asterisks in the same row indicate significant differences relative to empty control (*P < 0.05, **P < 0.01, ***P < 0.001).

significantly, whereas the titratable acid (TA) content was decreased significantly in both two transgenic tomato lines compared with WT fruits, as shown in Fig. 4A and 4B. The sugar:

acid ratio in OE-2 and OE-3 was 13.45 and 11.57, respectively, which was much higher than that in WT (9.10) (Fig. 4C). We further examined the content of soluble sugars and organic acids using high performance liquid chromatography (HPLC). Fructose was significantly induced in *E8::PpbZIP44* fruits. However, sorbitol was significantly reduced. There were no significant changes in sucrose content in OE-3 (Fig. 4D). Malate and citrate contents were significantly reduced (Fig. 4E).

The amino acid biosynthesis and metabolism in transgenic and WT tomato fruits were significantly different. Tyramine and nicotianamine (NA) had significantly higher contents in transgenic tomato fruits. Hcy, Met, Glu, Gsh, Phe, and Val were increased ~1.45- to 1.7-fold in the E8::PpbZIP44 tomato fruits. Tyr and Asp were also slightly induced in corresponding E8::PpbZIP44 tomato fruits. On the other hand, Pro level was 2.3-fold lower in the transgenic tomato fruits (Supplementary Table S5).

The contents of most unsaturated fatty acids were increased by 1- to 3-fold in the E8:PpbZIP44 fruits, compared with WT (Table 2). For examples, punicic acid (9Z,11E,13Z-octadecatrienoic acid), 9Shydroxy-10E,12Z-octadecadienoic acid, and 9-hydroxy-13-oxo-10octadecenoic acid were significantly increased >2-fold in the E8:PpbZIP44 fruits. In addition, some other unsaturated fatty acids such as linoleoylglycerol, crepenynic acid, linolenic acid, (9Z,11E)-

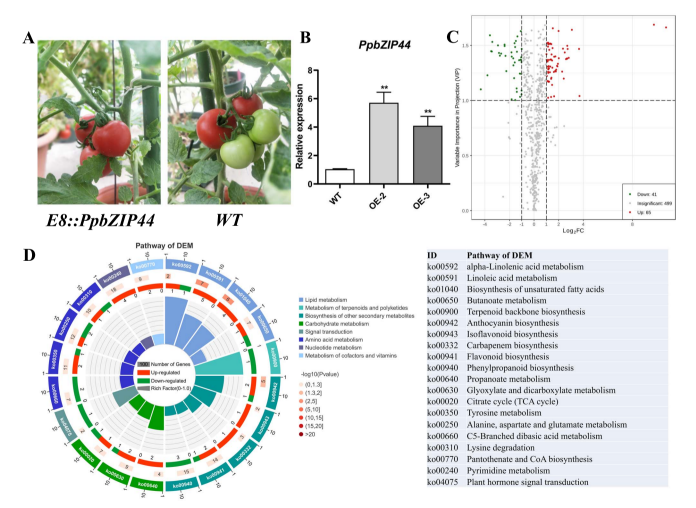


Figure 3. Fruit-specific overexpression of *PpbZIP44* in tomatoes. A. Representative phenotype of *E8::PpbZIP44* tomato fruits. B. The expression levels of *PpbZIP44* in WT and *E8::PpbZIP44* fruits. C. Volcano plot of DEMs in *E8::PpbZIP44* versus WT fruits. D. Summary of top 20 KEGG pathway, significantly overrepresented in DEMs in *E8::PpbZIP44* versus WT fruits. The results are averages \pm SE (n \geq 3) of WT and transgenic lines, each with three technical replicates. The asterisks indicated values determined by the Dunnett t test to significantly differ from the WT control (**P < 0.01).

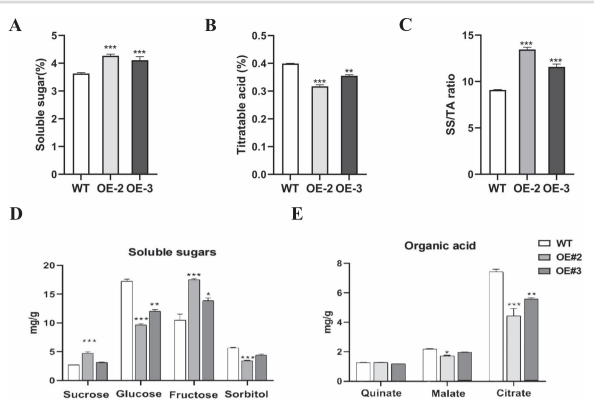


Figure 4. Overexpression of *PpbZIP44* significantly affected the contents of soluble sugars and organic acids in tomato fruits. A. SS contents. B. TA contents. C. SS/TA ratio. D. Individual soluble sugar content. E. Individual organic acid content. The results are averages \pm SE ($n \ge 3$) of WT and transgenic lines, each with three technical replicates. The asterisks indicated values that were determined by the Dunnett t test to be significantly different from the WT control (*P < 0.05, **P < 0.01, ***P < 0.001).

Table 2. Differentially expressed unsaturated fatty acids inE8::PpbZIP44 versus WT fruits

Compounds	VIP	Log2FC	Туре
1-α-Linolenoyl-glycerol	1.37	1.37	up
2-Linoleoylglycerol	1.02	1.14	up
1-Linoleoylglycerol	1.14	1.47	up
2-Dodecenedioic acid	1.44	-1.84	down
Crepenynic acid	1.51	1.16	up
γ-Linolenic acid	1.49	1.13	up
α-Linolenic acid	1.52	1.16	up
Punicic acid (9Z,11E,13Z-octadecatrienoic acid)	1.38	2.60	up
Linoleic acid	1.26	1.14	up
(9Z,11E)-Octadecadienoic acid	1.26	1.29	up
Elaidic Acid	1.28	1.17	up
11-Octadecanoic acid(Vaccenic acid)	1.31	1.32	up
9S-Hydroxy-10E,12Z-octadecadienoic acid	1.38	2.68	up
Eicosadienoic acid	1.63	1.99	up
9-Hydroxy-13-oxo-10-octadecenoic acid	1.39	2.82	up

octadecadienoic acid, and eicosadienoic acid were also increased in E8:PpbZIP44 fruits. However, 2-Dodecenedioic acid was slightly decreased in the fruits of E8:PpbZIP44 than that in the fruit of WT (Table 2).

Chalcone and flavonoid compounds were much more abundant in the fruits of E8:PpbZIP44 than WT, with increase of 1- to 12fold changes. 3,4,2',4',6'-Pentahydroxychalcone-4'-O-glucoside, isorhamnetin-3-O-(6"-acetylglucoside), homoeriodictyol, and cinnamic acid were increased >10-fold in E8::PpbZIP44 versus WT. The overexpression of PpbZIP44 also strongly increased other dihydro flavones such as hesperetin-7-O-(6"-malonyl) glucoside, poncirin, eucalyptin, naringin, neohesperidin, hesperidin, naringenin, and butin; flavonols such as kaempferol-3-O-(2"-o-acetyl) glucuronide, rutin, quercetin-3-o-robinobioside, 6-hydroxykaempferol-7-O-glucoside, brassicin, isohyperoside, and hyperin; isoflavones like genistin and calycosin-7-Oglucoside, and flavones like chrysoeriol-7-O-rutinoside. However, delphinidin-3-O-glucoside (Mirtillin) and Isorhamnetin-3,7-O-diglucoside were significantly decreased in the fruits of E8:PpbZIP44 than in the fruit of WT (Table 3).

Transcriptome changes induced by *PpbZIP44* in transgenic tomato fruits

To elucidate how PpbZIP44 regulates the expression of downstream genes leading to the biochemical alterations described previously, we performed transcriptomic analyses using the E8::PpbZIP44-2 transgenic tomato line. The results showed that there were 1314 differential expression genes (DEGs) (false discovery rate (FDR) < 0.05 and $|Log_2FC| > 1$) when comparing the E8::PpbZIP44-2 versus WT tomato fruits (Fig. 5A). The top overrepresented KEGG pathways among the DEGs included carbohydrate metabolism, amino acid metabolism, and biosynthesis of secondary metabolism (Fig. 5B). The carbohydrate metabolism pathways included fructose and mannose metabolism, glycolysis, pentose phosphate pathway, starch and sucrose metabolism, and galactose metabolism (Fig. 5B and Supplemental table S6). Amino acids metabolism included glutathione metabolism, alanine, aspartate and glutamate metabolism, tyrosine metabolism, arginine and proline metabolism, and arginine biosynthesis (Fig. 5B and Supplemental Table S6). Most of the DEGs were upregulated in the carbohydrate and amino acid metabolism pathways (Fig. 5B).

Class	Metabolites	VIP	Log2FC	Туре
Chalcones	3,4,2′,4′,6'-Pentahydroxychalcone-4'-O-glucoside	1.62	12.88	Up
	Okanin	1.12	1.41	Up
Dihydroflavone	Homoeriodictyol	1.59	10.05	Up
	Hesperetin-7-O-(6"-malonyl)glucoside	1.54	3.35	Up
	Poncirin (Isosakuranetin-7-O-neohesperidoside)	1.23	1.93	Up
	Eucalyptin	1.49	1.86	Up
	Naringin	1.4	1.64	Up
	Neohesperidin	1.3	1.49	Up
	Hesperidin	1.03	1.39	Up
	Naringenin	1.23	1.23	Up
	Butin	1.25	1.19	Up
	6-C-Glucosyl-2-Hydroxynaringenin	1.23	1.02	Up
Flavonols	Isorhamnetin-3-O-(6''-acetylglucoside)	1.66	10.73	Up
	Kaempferol-3-O-(2"-O-acetyl)glucuronide	1.44	2.5	Up
	Rutin	1.31	1.9	Up
	Quercetin-3-O-robinobioside	1.28	1.8	Up
	6-Hydroxykaempferol-7-O-glucoside	1.27	1.72	Up
	Brassicin	1.04	1.63	Up
	Isohyperoside	1.26	1.61	Up
	Hyperin	1.36	1.47	Up
	Isorhamnetin-3,7-O-diglucoside	1.23	-3.79	Down
Isoflavones	Genistin	1.51	1.28	Up
	Calycosin-7-O-glucoside	1.46	1.24	Up
Flavones	Chrysoeriol-7-O-rutinoside	1.33	1.94	Up

Table 3. Differentially expressed flavonoid compounds in E8::PpbZIP44 versus WT

DEGs in the carbohydrate metabolism pathway were further analyzed. In starch and sucrose metabolism, the expression of an INV, two SS genes, TPS and TPP, two BGLU genes, and GLU was significantly induced by the overexpression of PpbZIP44. On the contrary, the expression of BAM, linking in starch and maltose and dextrin, was significantly downregulated by PpbZIP44 (Fig. 5C and Supplemental Table S6).

In the fructose and mannose metabolism pathway, the expression of FRK2 and MAN was significantly induced by PpbZIP44. However, MPI isomerase was significantly downregulated (Fig. 5D and Supplemental Table S6). In the glycolysis and pentose phosphate pathway (PPP) pathway, the expressions of PFP, PFK, FBA, PDC, and RBSK were significantly upregulated in the E8::PpbZIP44–2 transgenic tomato fruit, as shown in Fig. 5D. However, the expression of ENO was downregulated. In the TCA cycle, the expression of CS was repressed but the expression of ACO, ICDH, and MDH was induced in the transgenic tomato fruits (Fig. 5E).

In the amino acid biosynthesis pathway, 18 DEGs were found in E8:PpbZIP44 tomato fruits versus WT fruits. Most of those, including ALT, AST, ADT, APX, PAL, GLT1, argE, ADH2, HPD, GCCT, three GSTs, GDH, AHCY, and P4H3, were significantly upregulated (Fig. 5F and supplementary Table S6). Flavonoids biosynthesisrelated DEGs, such as C4H and F3H, were substantially upregulated by the overexpression of PpbZIP44 (Fig. 5G).

The expression changes of genes induced by *PpbZIP44* in transformed pear fruits

To further examine and verify the changes in the expression levels of DEGs identified from the transcriptome analyses in the tomato system, we measured the transcript abundances of 12 DEGs using qRT-PCR in the transient overexpression or silencing *PpbZIP44* pear fruit. The results confirmed that *PpSDH9* gene was significantly induced in OE-bZIP44 but inhibited in RNAi-bZIP44 pear fruits. The expression levels of PpSUS, PpFRK, PpACO, PpICDH, and PpMDH were significantly increased in OE-bZIP44. However, the expression level of PpCS was significantly reduced in OEbZIP44. PpProDH1 transcript accumulated approximately 13-fold in OE-bZIP44. In contrast, it reduced to 0.5-fold in RNAi-bZIP44. PpADT transcript accumulated 2-fold in OE-bZIP44, reducing to 0.45-fold in RNAi-bZIP44. The transcript abundance of PpC4H and PpF3H was strongly induced in OE-bZIP44, whereas reduced in RNAi-bZIP44 (Fig. 6).

PpbZIP44 directly binds to the promoters of PpSDH9, PpADT, PpProDH1, and PpF3H

To understand how bZIP44 regulates the expression of the downstream target genes, we performed the promoter analysis for identifying the G-boxes cis-element and the dual luciferase transient expression assays. We identified G-boxes cis-element in the 2000bp promoter of these DEGs by PpbZIP44 (Supplementary Table S7). Among the promoters of the 72 upregulated genes induced by PpbZIP44, 21 contained G-box motifs. Next, we identified typical G-boxes in the promoter of these DEGs that are involved in sorbitol and fructose metabolism, amino acids biosynthesis, and flavonoids biosynthesis, finding the promoters of PpSDH9, PpADT, PpProDH1, and PpF3H with G-boxes cis-elements (Supplementary Table S7). The regulation of PpbZIP44 on these genes was hypothesized. The promoter sequences (within 1 kb) of PpSDH9, PpADT, PpProDH1, and PpF3H from 'Sucui 1' pear fruit were cloned and analyzed using PlantCARE (Supplemental Table S8). Three to six G-box cis-acting elements (CACGTT), which were predicted to recruit S1-bZIP, were found in the promoter of these four genes. The 1.0-kb fragments were cloned into the pGreen 0800-luciferase (LUC) vector for dual luciferase transient expression assays. Compared with the control level, the coexpression of 35S::PpbZIP44 and pSDH9::LUC, pProDH1::LUC, pADT::LUC, or pF3H::LUC resulted in a 3.42-, 2.82-, 5.46-, and 7.27-fold increase in LUC activity, separately

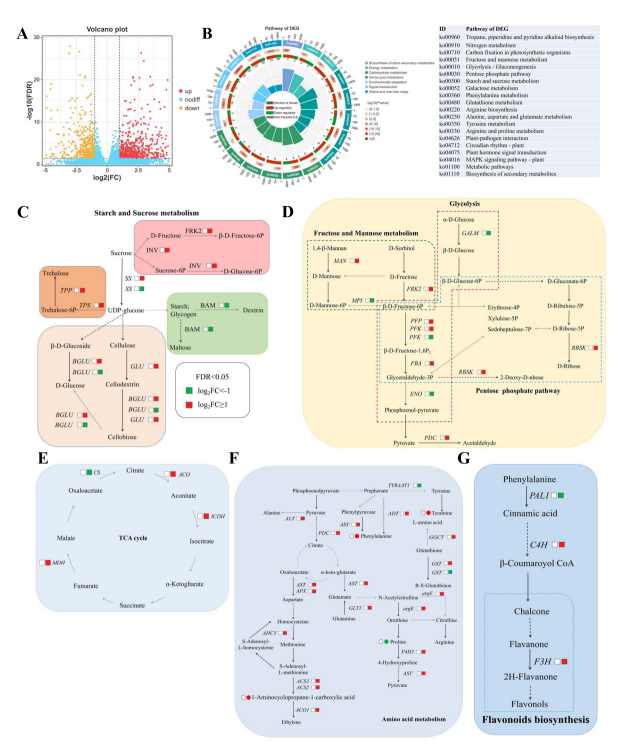


Figure 5. *PpbZIP44* significantly changed the primary and secondary metabolism pathway in tomato fruits based on transcriptome data. A. Volcano plot of DEGs in *E8::PpbZIP44* vs. WT fruits. B. Enriched top 20 KEGG pathway of DEGs in fruits of *E8:PpbZIP44* (OE-2) compared with WT. Detailed data are shown in the Support information Supplemental Table S6. C–E. Regulation mechanism analysis of *PpbZIP44* in the carbohydrate metabolism. DEGs induced by *PpbZIP44* in the starch and sucrose metabolism (C), fructose and mannose metabolism, glycolysis, and pentose phosphate pathway (D) and the TCA cycle (E). F. DEGs induced by *PpbZIP44* in the amino acid biosynthesis. G. DEGs induced by *PpbZIP44* versus WT (white squares). INV encodes *β*-fructofuranosidase; *GLU* encodes endoglucanase; *BAM* encodes *β*-amylase; *FRK2* encodes fructokinase; *MAN* encodes mannan endo-1,4-beta-mannosidase; *MPI* encodes mannose-6-phosphate isomerase; *PFP* encodes diphosphate-dependent phosphofructokinase; *PFK* encodes 6-phosphofructokinase; *APX* encodes fructose-bisphosphate aldolase; *RBSK* encodes ribokinase; *ADT* encodes acetylomithine deacetylase; *GST* encodes glutathione S-transferase; *argE* encodes acetylornithine deacetylase; *P4H3* encodes prolyl 4-hydroxylase; *AST* encodes aspartate aminotransferase.

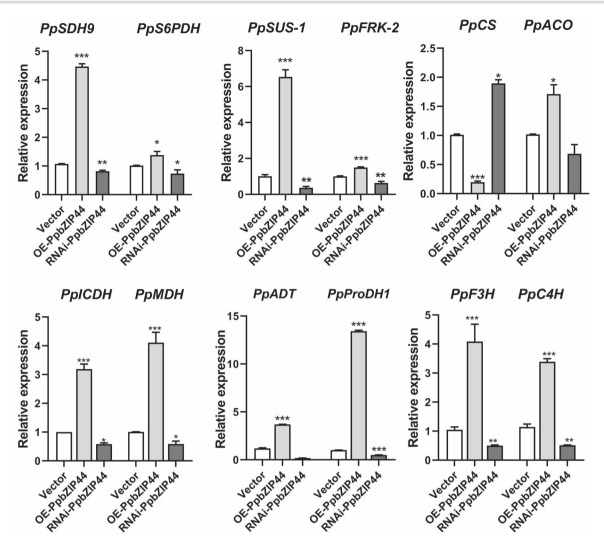


Figure 6. qRT-PCR analysis of gene expression related to sugar, amino acid, and flavonoid metabolism pathways and the TCA cycle in transiently transgenic pear fruits. Samples were collected in transformed pear fruits at 7 days after injecting overexpression (OE- *PpbZIP44*) or RNAi (RNAi-*PpbZIP44*) or empty vector (Vector) constructs. The results are average \pm SE of three biological replicates, each with three technical replicates. The asterisks indicated values that were determined by the Dunnett t test to differ from their empty vector or WT control significantly (*P < 0.05, **P < 0.01, ***P < 0.001).

(Fig. 7A). To further verify the binding of PpbZIP44 to the G-box in the promoter of these genes, we performed an electrophoretic mobility shift assay (EMSA). A 30-bp fragment (position –289 to –259 of *pSDH9*, –291 to –261 of *pADT*, –243 to –202 of *pProDH1*, and –178 to –148 of *pF3H*) was used as a probe for the EMSA (Fig. 7B). Explicit binding of the PpbZIP44 protein to the biotinlabeled target probes was detected in the polyacrylamide gel. In contrast, signals were undetectable for the protein–mutant probe complex with nucleotide changed from CACGTT to TGTACC. The signals were reduced or faded away by adding cold probes (Fig. 7B). These results suggested that *PpbZIP44* was directly bound to the cis-element of the *PpSDH9*, *PpADT*, *PpProDH1*, and *PpF3H* promoters and induced their transcript, enhancing the accumulation of fructose, phenylalanine, and flavonoids but deleting proline.

Discussion

PpbZIP44 modulates carbohydrate metabolism of fruit

Soluble sugar accumulation determines fruit sweetness and nutritional quality at harvest. Therefore, increasing sweetness

is an important breeding objective and a vital driver of consumer preference in many species [23]. Fructose is the predominant and the sweetest monosaccharide in the fruit of most pear cultivars [24, 25]. The content of fructose directly determines the quality of fresh pears. Changes in sorbitol metabolism substantially affect fructose content. In a recent study, overexpression of apple MdABI3 increased fructose content by inducing MdSDH2 expression [7]. Pear SDH genes were identified, and their expression patterns were analyzed [26]. However, it is unclear how regulating sorbitol metabolism contributes to the change in fructose content in pear fruit. Here, we found that pear PpbZIP44 was involved in regulating the expression of a sorbitol-fructose convert. During fruit development and ripening processes in pear, PpbZIP44 displayed significantly higher transcript abundance at stage 5, in which cell division and expansion occurs and the most significant transcriptional changes and fruit grow and metabolites associated with quality traits are accumulated [27]. Most DEGs involved in the primary metabolism were upregulated in this stage [27]. Our results suggest that PpbZIP44 is a positive regulator of soluble sugars accumulation in pear fruits. Transiently overexpressed and gene-silenced pear fruits

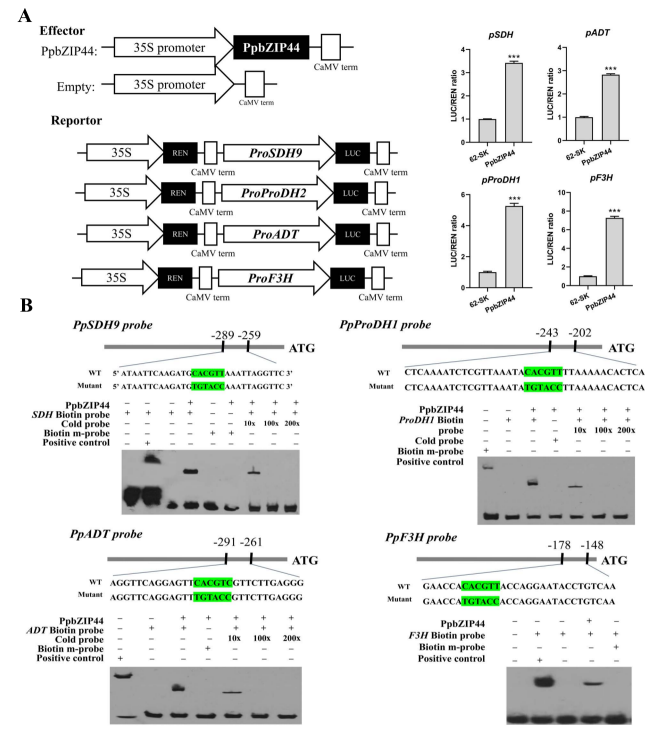


Figure 7. PpbZIP44 activated PpSDH9, PpProDH1, PpADT, and PpF3H by binding their promoters. A. Analysis of luciferase activity with pSDH9, pProDH1, pADT, and pF3H. 1.0-kb sequences upstream from ATG of these three genes were fused to the reporter gene Luciferase (LUC). LUC activities were assayed in transiently transformed tobacco leaves. Left: the vector construction for luciferase assays. Bars represent the mean value \pm SE ($n \ge 10$). The data were analyzed using the Dunnett t test (***P < 0.001). B. EMSA of PpbZIP44 binding to the pSDH9, pProDH1, pADT, and pF3H containing ACGT core site. The autoradiograph shows the DNA–protein complex of the biotin-labeled probe. Biotin probe is 30 bp of pSDH9, pProDH1, pADT, and pF3H promoter, with the CACGTT sequence; 10×, 100×, and 200× are the relative contents of the competitor probe to the detector probe. Biotin m-probe is 30 bp of pSDH9, pProDH1, pADT, and pF3H with the CACGTT mutation as TGTACC sequence.

with *PpbZIP44* ORF confirmed the role of *PpbZIP44* in promoting *PpSDH9* expression and enhancing *PpSDH* enzyme activity, fructose and sucrose accumulation, and reducing sorbitol accumulation.

SDH catalyzes the reversible sorbitol-fructose reactions and is found in core eudicot species and monocots [28]. Compatible

with the role in the initial metabolic steps of carbohydrate metabolism, *SDHs* are essential for the normal growth of plants and stress response [28–31]. Unlike pear and other Rosaceous trees, *Zea mays* does not transport sorbitol through its phloem, but instead synthesizes sorbitols in the kernel itself [32]. In maize, *Sdh1* is essential during early kernel development and

in synthesizing sorbitol from fructose [33]. Conversely, in this study, the upregulation of PpSDH9 by overexpression of PpbZIP44 demonstrated its central role in fructose accumulation from sorbitol in pear fruits, which is consistent with the role of MdSDH2 in apple fruit [7]. SDH catalyzes the conversion of fructose to sorbitol or sorbitol to fructose in a reversible manner, which might depend on species such as usual sorbitol producers or non-usual sorbitol ones [31]. In apple fruit, a large amount of sorbitol is converted into fructose and 80% of the total carbon flux goes through fructose [7], entering glycolysis and PPP. In this work, overexpression of PpbZIP44 substantially upregulated the expression of SUS, FRK2, PFP, PFK, and FBA in the glycolytic direction. G-boxes cis-elements were found in the promoter of these genes, implying the bound of PpbZIP44 to these genes (Supplementary Table S7). Accordingly, our study provides evidence supporting the notion that PpbZIP44 is involved in soluble sugars accumulation and guides carbon flux to the downstream pathway from fructose.

Interestingly, down- or up-regulation of SDH genes, as well as S6PDH and A6PR, modulates sorbitol contents, leading to stamen development, pollen tube growth, and other stress responses [34-37]. SnRK1 promoted sorbitol metabolism by activating SDH enzyme and phosphorylating PpSDH protein in peach [38]. Most recently, overexpression of MdSnRK1 increased the transcript abundance of MdSDH1 and MdA6PR through SnRK1-mediated phosphorylation of MdbZIP39 [39]. Whether bZIP39 and bZIP44 interact with each other and whether or how SnRK1 mediates this interaction require further study in the future. Increased SDH expression and SDH activities allowed a higher carbon flux through sorbitol-fructose metabolism for plant growth and development with enough sorbitol supply [39]. The various roles of PpbZIP44-mediated PpSDH9 expression on the growth and development and stress response of pear still need to be investigated.

Organic acids are significant factors determining the fruit sugar:acid ratio. Engineering organic acid metabolism substantially improved fruit traits [3]. In pear fruit, organic acids were mainly composed of quinate, citrate, and malate. Quinate can be synthesized from either 3-dehydroquinate or shikimate; these two reactions are catalyzed by QDH and QD, respectively [40]. Here, there were no significant differences in the expression level of QDH and QD and other genes in the shikimate pathway between E8::PpbZIP44 and WT. No significant differences were observed in the content of quinate either in the overexpressed PpbZIP44 pear or tomato fruits compared with WT. Citrate and malate control fruit acidity and are synthesized in the mitochondrion but accumulate in the vacuole [41, 42]. Citrate biosynthesis and conversion mediated by CS and ACO, or malate catalysis mediated by MDH, is responsible for a change in citrate or malate contents [42]. The influx and efflux and the capacity of other parts of the TCA cycle affect the import or export rate of citrate and malate [42]. In previous research, there were no apparent effects of manipulating a range of enzymes that have been proposed to control fruit citrate or malate accumulation on increasing carboxylic acids in fruits [42]. However, manipulating transcription factors such as MYB, bHLH, WRKY, and ERF family members regulated citrate or malate level [43, 44]. In Arabidopsis, induced AtbZIP11 depleted the levels of intermediates in the TCA cycle [13]. Here, fruit-specific promoter driving overexpression of PpbZIP44 led to significant enrichment of the TCA cycle in transgenic tomato fruits with decreasing content of α -Ketoglutaric acid, citrate, and malate. Transient expression of PpbZIP44 significantly decreased the level of citrate and malate in

OE transgenic pear fruits, which substantially contributed to the higher sugar:acid ratio in transgenic fruits.

PpbZIP44 regulated amino acid biosynthesis and mediated carbon flux toward secondary metabolism and flavonoid accumulation

Amino acids are pivotal in human nutrition as a source of nutraceutical compounds or essential dietary components [10]. The genes in the amino acid metabolism pathway are involved in plant response to abiotic stress [45]. The ProDH gene encodes a proline dehydrogenase and catalyzes the catabolism of proline [12]. In Arabidopsis, AtbZIP11 can bind to the promoters of ProDH and ASN1 to induce their expression [12]. Here, 13 more folds increase of PpProDH1 expression was induced by PpbZIP44. PpbZIP44 induced rapid proline catabolism and significant enrichment of PPP, glycolysis, and the TCA cycle. We speculated that the accelerated proline cycle was coupled to the PPP, driving sugar phosphates and glycolysis for downstream anabolic pathways. Intermediates in the glycolysis pathway and the TCA cycle provide substrates and energy for the biosynthesis of amino acids and fatty acids [10, 46]. Amino acid biosynthesis pathways were significantly enriched and a series of genes involved in the pathways were upregulated. Significantly changes in Asp, Glu, Thr, Met, Val, and Ser contents illustrated the positive role of PpbZIP44 in prompting the accumulation of these amino acids. In addition, PpbZIP44 induced significant enrichment of 'pantothenate and CoA biosynthesis', 'linoleic acid metabolism', 'alpha-Linolenic acid metabolism', and 'biosynthesis of unsaturated fatty acids' and significantly increased unsaturated fatty acids contents. All together suggested that manipulating carbohydrate metabolism and accelerating proline catalysis and the TCA cycle through PpbZIP44 are new and efficient strategies not only in sugar accumulation but also adjusting the balance between sugar, organic acids, and other primary metabolites and eventually affecting fruit quality, as well as role in balancing carbon to nitrogen metabolism in Arabidopsis [13].

The critical role of this TF on secondary metabolism was further determined by metabolism and transcriptome in transgenic tomato fruits and transient transgenic pear fruits. Phenylalanine is biosynthesized through the shikimate pathway and arogenate pathway [47]. ADT encodes arogenate dehydratase and catalyzes the final step of the arogenate pathway. Moderate expression of ADT increased flux through the phenylalanine biosynthesis [47]. Interestingly, other TFs such as PqMyb308-like can regulate the expression of genes in the shikimate pathway and promote the accumulation of shikimate, aromatic amino acids, and total lignins but reduce the contents of multiple flavonoids [48]. However, in this study, overexpression of PpbZIP44 induced the expression of ADT and the increase of phenylalanine levels in fruits rather than activating the differential expression of genes in the shikimate pathway. Phenylalanine provides the substrates for these secondary metabolites, including flavonoids. Dihydro flavone contents displayed a remarkable increase in transgenic fruits. Dual-luciferase assays and EMSA demonstrated that PpbZIP44 strongly activates PpF3H promoters and induces its expression. Notably, the content of many phenolic acids and the expression of genes in the lignin pathway, cutin, suberin and wax biosynthesis and zeatin biosynthesis were significantly downregulated, which was firstly found in the present study and need to further illustrate the molecular mechanism in the future.

Although the metabolic diversity of fruits in different species exists, e.g. soluble sugars were the major metabolites in pear fruits, whereas of considerably lower levels in tomato fruits. Large amounts of alkaloids were detected in tomato fruits but not in pear fruits. Total metabolic pathways and signatures related to quality formation, such as sugars, acids, flavonoids, phenylpropanoids, lipids, and amino acids, are highly conserved among fruits [6]. Due to the difficulty of obtaining stable transgenic pear fruits in fruit quality-related research, a stable transformation system in tomato fruits could be a supplemental function analysis of a transient transformation system. Fruit-specific promoter drivers powerful TFs, which might be involved in considerable growth and development processes, successfully avoiding the adverse effect on plant growth. RNA-Seq in *E8::PpbZIP44* tomato fruits and qRT-PCR examination in transient transformation pear fruits provide plenitude shreds of evidence for the role of *PpbZIP44* in fruit multimetabolism reprogramming and quality formation.

In summary, we generated a model to elucidate the regulation mechanism of PpbZIP44 on substantial changes in multiqualityrelated compounds. PpbZIP44 could bind to the PpSDH9 promoter to induce its expression. The enhanced activity of SDH promoted the conversion of sorbitol to fructose, resulting in a considerable accumulation of fructose. PpbZIP44 substantially induced the higher expression of SUS, FRK2, PFP, PFK, and FBA in the glycolytic direction. PpbZIP44 recruited the PpProDH1 promoter to accelerate proline catabolism, which might couple PPP to drive sugar phosphates and glycolysis for downstream anabolic pathways. Depleting the intermediate of the TCA cycle facilitated the biosynthesis of amino acids and affected the sugar:acid ratio. Moreover, PpbZIP44 bound to the PpADT promoter to induce their expression, leading to metabolic flux toward phenylalanine. PpbZIP44 bound to the PpF3H promoter and induced the increase of dihydro flavone contents. Metabolic flux toward dihydro flavone might decrease lignin formation, which is involved in cell stone formation and crucial for the perception of the quality of pear fruit. Regarding Asian pears, high sugar content, moderate acidity, crispy, few stone cells, and high nutritional value are desirable traits. Identification and functional characterization of PpbZIP44 provides a novel strategy for molecular breeding through metabolic engineering for fruit quality improvement.

Materials and methods Plant materials

Fruits at different development stages in 'Sucui 1' were collected every 15 days after full bloom (DAFB) until 90 DAFB for stage 6. Samples at 93 DAFB and 100 DAFB were collected for stages 7 and 8 from the Pear Germplasms Resources at the Institute of Pomology, Jiangsu Academy of Agricultural Sciences, Nanjing, Jiangsu, China. The trees were cultivated as follows: $3.0 \text{ m} \times 5.0 \text{ m}$ spacing, with rows oriented in the east–west direction. Trees were pruned into an arch shape. Base fertilizer was applied in the autumn and the trees were irrigated twice weekly. Alfalfa in winter and white clover in summer between rows were cut up regularly and left on the ground. The pear fruit peel (0.3 cm) was removed and sarcocarp samples were frozen in liquid nitrogen. The sarcocarps were collected with three independent biological replicates for qRT-PCR, transient transformation, or DNA extraction.

Identification of S1-bZIP proteins in the sand pear

All bZIP protein sequences of Arabidopsis were obtained from Arabidopsis Information Resource (http://arabidopsis.org/; release 10.0). Pear genome sequences were obtained from National Genomics Data Center (https://bigd.big.ac.cn/gwh.). The Hidden Markov Model profile of bZIP_1 (PF00170) and bZIP_2 (PF07716) was obtained from the Pfam database (http://pfam.sanger.ac.uk) [49] and SMART database (http://smart.embl-heidelberg.de) [50]. A local BLAST search was completed with Arabidopsis bZIP proteins as queries to remove redundancies by checking all identical sequences manually. All the confirmed pear bZIP proteins were aligned using MUSCLE (version 3.8.31) [51]. A phylogenetic tree was generated by analyzing the multiple sequence alignment with the Clustal W [51], and the alignment results were displayed using Jalview [52]. The sand pear S1-bZIP proteins were further identified through a local BLAST search of Arabidopsis, tomato, tobacco, grape, apple, strawberry, rice, cucumber, banana, and petunia protein sequences, downloaded from Solanaceae Genome Annotation Database (https://solgenomics.net) and Plant Transcription Factor Database (http://planttfdb.cbi.pku.edu.cn; release 3.0).

Transient overexpression and RNAi silencing of genes in pear fruits

The transient transformation and expression analysis were performed as Yao described [53]. Briefly, total RNA was extracted from 'Sucui 1' sarcocarps and used as the template to synthesize cDNA. The mORF sequence of PpbZIP44 (EVM0042575) was ligated to the pGreenII 62-SK or pSAK277 vectors using OE- or RNAi-primers (Table S1) under the control of the 35S promoter [53]. Agrobacterium tumefaciens-mediated transient transformation was performed in 'Sucui 1' pear fruits at 90 DAFB [53]. Fruits transformed with pGreen II 62-SK or pSAK277 empty vector were used as controls. The fruits of three biological replicates, including five fruits at least for each experiment, were collected for sugars, organic acids, amino acids, and enzyme activity measurements, and a qRT-PCR examination after 7 days.

Fruit-specific driving vector construction and tomato transformation

The CaMV 35S promoter was eliminated and E8 promoter (Gen-Bank: DQ317599) was amplified (Table S1) and linked to pBI121-GFP [18]. The fragment was then ligated into pBI121-S1-bZIP-GFP vectors. The cloned inserts, including the E8 promoter and target genes, were confirmed by sequencing. Explants were prepared and transformation was performed as described by Sagor *et al* [18]. The regenerated plantlets were then transferred to the rooting medium until the shoot grows to 2- to 4-cm long. The DNA of tomato plantlets was extracted and analyzed by RT-PCR with the bZIP44-gF2 and GFP-FCX_R. Positive transgenic plantlets were confirmed by RT-PCR and sequenced, after which they were planted in soil in a greenhouse.

Widely targeted metabolomics analysis based on UPLC-MS/MS system

Compounds were extracted and examined using A UPLC (SHIMADZU Nexera X2) and Tandem mass spectrometry MS/MS (Applied Biosystems 4500 QTRAP) as previously described [22, 26]. With the methods we used, all metabolites can be measured simultaneously using widely targeted metabolomics based on MRM [22, 26]. The relative signal intensities of metabolites were normalized by dividing them according to the intensities of the internal standard (lidocaine, 0.1 mg·l⁻¹) first and after that log₂ transform them to get further normalization for normality improvement [54]. Then we calculated the area of each peak to quantify the metabolites. The PCA and orthogonal partial least squares discriminant analysis (OPLS-DA) analysis were completed to assess the differences in compounds. VIP \geq 1 and

fold-change ≥ 2 or ≤ 0.5 were used as threshold levels for selecting differentially abundant metabolites.

RNA isolation, sequencing and differential gene expression analyses

Each biological replicate consisted of at least five fruits from different trees. Total RNA from leaves and fruits was extracted from the sarcocarp or tomato flesh with Trizol (Invitrogen, USA) and the Ambion RiboPureTM Kit (Ambion, USA) [55]. Then, mRNA was prepared using Oligo (DT) system. cDNA libraries were constructed and sequenced with nova-seq6000 platform based on the manufacturer's instructions. Clean reads of the nine samples were pooled and mapped to the tomato genome (SolGenomicsNetwork, S. lycopersicum 3.0) with HISAT2 (HISAT2 (daehwankimlab.github.io)) [56]. Transcripts were assembled and quantified with StringTie [57]. Differential analysis was performed with FDR < 0.05 and |log2FC| > 1 as described [58]. Additionally, GOseq (adjusted P < 0.05) was used for the GO term enrichment analysis [59], whereas KO (KEGG Ortholog database) was used to enrich DEGs to different pathways [60].

Soluble sugar and organic acid content measurement

The content of the total SS and TA of fruits was measured according to the previous description [17]. Sucrose, glucose, fructose, sorbitol, quinate, malate, and citrate levels in pear and tomato fruits were measured using HPLC [17]. Briefly, 0.8 g of frozen pear sarcocarps were ground into powder and then resuspended with 800 μ l of extraction buffer containing 0.2% metaphosphoric acid and 80% ethanol. The mixture was treated with an ultrasonicator for 10 min, incubated overnight at 4°C, and centrifuged at 10000 rpm for 10 min at 25°C. 800 μ l of supernatant was concentrated for 3 h at 30°C, dissolved in 1.6 ml deionized water (Milli-Q water), and then filtered through 0.22- μ m Sep-Park filter paper. Organic acids and soluble sugars were analyzed with an HPLC system. We calculated the area of each peak to quantify the metabolites according to an external standard calibration method.

Amino acid content measurement

For amino acid content, pear fruit samples were directly frozen and ground in liquid nitrogen after transient transformation for 7 days. Total amino acids were extracted with a mixed 100to 200-mg sample and 750 μ l of 80% ethanol in an ultrasonic bath at 50°C for 30 min. Then we centrifuged samples, collected the supernatants and removed the solvent by freeze-drying. The extract was added with 5.7 M of hydrochloric acid, hydrolyzed overnight under a closed condition at 110°C, dissolved in water, dried again, and resuspended in 1 ml of sodium citrate buffer (pH 2.2). The composition of amino acid in all samples was tested by Hitachi amino acid automatic analyzer LA8080.

SDH enzyme activity measurement

Activities of SDH were measured using the commercial kits (catalog #:ab252902) based on the instruction. Quantifying enzyme activities were measured through a spectrophotometer with UV-VISO 2450 at 25°C (Shimadzu, Kyoto, Japan).

qRT-PCR

cDNA was synthesized using the HiScript III RT SuperMix Kit with gDNA wiper (Vazyme, China) for a qRT-PCR analysis with specific primers (Table S1). The pear GAPDH and tomato SlACTIN genes were used as internal standards for data normalization. Average

threshold cycle values were counted based on three independent biological replicates and then calculated the relative transcript level of each gene [61].

Dual luciferase assay

The dual luciferase reporter assay was performed as previously described [7]. The coding sequence of full-length PpbZIP11 (EVM0042575) was cloned into the pGreenII62-SK to produce the effector construct. A 1000-bp promoter sequence (named pSDH9, pProDH1, pADT and pF3H) of PpSDH9, PpProDH1, PpADT, and PpF3H (Supplemental Table S8) was introduced separately into the pGreenII0800-LUC vector. The transformed A. *tumefaciens*-strain GV3101 cells were used to coinoculate the young leaves of Nicotiana benthamiana tobacco plants. The LUC and renilla luciferase (REN) activities were measured with a dual luciferase assay system (Promega, USA) and an Infinite M200 (TECAN, Switzerland), recorded as a ratio of LUC to REN.

EMSA

The EMSA assay was conducted as previously described [62]. Briefly, the PpbZIP44 ORF was ligated into the pET-28a prokaryotic expression vector. The resulting plasmid was inserted into Escherichia coli Rosetta cells and cultured on LB medium with 40 mg·l⁻¹ kanamycin to overexpress the PpbZIP44 protein via the addition of isopropyl β -D-1-thiogalactopyranoside and an overnight incubation at 16°C. The recombinant protein was purified with the HisTrap HP column (GE Healthcare, Piscataway, NJ, USA). The biotin-labeled DNA fragments (pSDH9: ATAATTCAAGATGCACGTTAAATTAGGTTC; pProDH1: CTCAAAATCTCGT-. TAAATACACGTTTTTAAAAACACTCA; pADT: AGGTTCAGGAGTTCACGTCGTTCTT-. GAGGG; pF3H: GAACCA-CACGTTACCAGGAATACCTGTCAA) were synthesized, annealed, and used as WT probes, whereas the corresponding sitedirected (CACGTT) mutated DNA fragments (TGTACC) were used as mutant probes. WT probes without label served as the competitors. The interaction between PpbZIP44 and the probes and the binding signals were detected with the LightShift EMSA Optimization and Control Kit and the Chemiluminescent Nucleic Acid Detection Module Kit (Thermo Fisher, MA, USA).

Statistical analysis

Dunnett t test in the multiple comparison analysis of variance was applied to determine whether any of the differences between the mean values for treatments and genotypes were significant.

Acknowledgments

We thank Prof. Songling Bai for kindly providing the genome database of 'Cuiguan' pear. We thank Chuan-Bei Jiang (Genepioneer Biotechnologies, Nanjing 210014, China) for providing bioinformatics analysis. This work was supported by Jiangsu Agricultural Science and Technology Independent Innovation Fund [Grant No. CX(22)3010], the General Program of National Natural Science Foundation of China (Grant No. 31872078, 32272669), Jiangsu Modern Agricultural Industry Technology System (Grant No. JATS[2022]437, JATS[2022]438), Jiangsu Agricultural Key New Varieties Innovation (Grant No. PZCZ201726), Jiangsu Province Seed Industry Revitalization Unveiled Project: Breeding of New Pear Cultivars with High-quality and Extreme Early Ripening (Grant No. JBGS[2021]084). The funders had no role in study design, data collection, and analysis.

Author Contributions

H.W., L.J., and C.Z.J. designed and supervised the research. WJ., B.L.S., and Y.H.C. suggested the research. H.W. and K.X.X. wrote the manuscript. B.B.U. and C.Z.J. revised the manuscript. W.H. conducted the RNA-seq library and metabolites preparation, processed the RNA-seq data and metabolites data, gene clone, transformation, HPLC, EMSA, dual luciferase assay, and statistics. K.X.X. performed RT-qPCR and physiology experiments. H.W. and G.F.Y. performed transient transformation. Y.D.W. performed sequences analysis. Q.S.Y. and X.G.L. prepared pear fruits. All authors read and approved the final manuscript.

Data availability

All the RNA-seq data are available and have been deposited in the National Center for Biotechnology Information Sequence Reads Archive (PRJNA947778). Other relevant data are presented within the paper and its supplementary files.

Conflict of Interest

The authors declare that they have no conflict of interest.

Supplementary data

Supplementary data is available at Horticulture Research online.

References

- Adaskaveg JA, Blanco-Ulate B. Targeting ripening regulators to develop fruit with high quality and extended shelf life. *Curr Opin Biotechnol.* 2023;**79**:102872
- 2. Ruan YL. Sucrose metabolism: gateway to diverse carbon use and sugar signaling. Annu Rev Plant Biol. 2014;**65**:33–67
- 3. Batista-Silva W, Nascimento VL, Medeiros DB et al. Modifications in organic acid profiles during fruit development and ripening: correlation or causation? Front Plant Sci. 2018;**9**:1689
- Fei X, Hu H, Luo Y et al. Widely targeted metabolomic profiling combined with transcriptome analysis provides new insights into amino acid biosynthesis in green and red pepper fruits. Food Res Int. 2022;160:111718
- Sun H, Zhou X, Zhou Q et al. Disorder of membrane metabolism induced membrane instability plays important role in pericarp browning of refrigerated 'Nanguo' pears. Food Chem. 2020;320:126684
- 6. Pott DM, Osorio S, Vallarino JG. From central to specialized metabolism: an overview of some secondary compounds derived from the primary metabolism for their role in conferring nutritional and organoleptic characteristics to fruit. Front Plant Sci. 2019;**10**:835
- Wang Z, Ma B, Yang N et al. Variation in the promoter of the sorbitol dehydrogenase gene MdSDH2 affects binding of the transcription factor MdABI3 and alters fructose content in apple fruit. Plant J. 2022;109:1183–98
- Tanase K, Shiratake K, Mori H et al. Changes in the phosphorylation state of sucrose synthase during development of Japanese pear fruit. Physiol Plant. 2002;114:21–6
- Tanase K, Shiratake K, Yamaki S. The mechanisms of sucrose accumulation in Japanese pear (Pyrus pyrifolia Nakai) fruit. Acta Hortic. 2002;479-487:479–87
- Trovato M, Funck D, Forlani G et al. Editorial: amino acids in plants: regulation and functions in development and stress defense. Front Plant Sci. 2021;12:772810

- Carmona-Gutierrez D, Zimmermann A, Kainz K et al. The flavonoid 4,4'-dimethoxychalcone promotes autophagydependent longevity across species. Nat Commun. 2019;10:651
- Hanson J, Hanssen M, Wiese A et al. The sucrose regulated transcription factor bZIP11 affects amino acid metabolism by regulating the expression of ASPARAGINE SYNTHETASE1 and PROLINE DEHYDROGENASE2. Plant J. 2008;53:935–49
- Ma J, Hanssen M, Lundgren K et al. The sucrose-regulated Arabidopsis transcription factor bZIP11 reprograms metabolism and regulates trehalose metabolism. New Phytol. 2011;191:733–45
- Garg A, Kirchler T, Fillinger S et al. Targeted manipulation of bZIP53 DNA-binding properties influences Arabidopsis metabolism and growth. J Exp Bot. 2019;70:5659–71
- Wang H, Zhang Y, Norris A et al. S1-bZIP transcription factors play important roles in the regulation of fruit quality and stress response. Front Plant Sci. 2022;12:802802
- Rook F, Gerrits N, Kortstee A et al. Sucrose-specific signaling represses translation of the Arabidopsis ATB2 bZIP transcription factor gene. Plant J. 1998;15:253–63
- 17. Xing S, Chen K, Zhu H et al. Fine-tuning sugar content in strawberry. *Genome* Biol. 2020;**21**:230
- Zhang Y, Li S, Chen Y et al. Heterologous overexpression of strawberry bZIP11 induces sugar accumulation and inhibits plant growth of tomato. Sci Hortic. 2022;292:110634
- Sagor GH, Berberich T, Tanaka S et al. A novel strategy to produce sweeter tomato fruits with high sugar contents by fruitspecific expression of a single bZIP transcription factor gene. Plant Biotechnol J. 2016;14:1116–26
- Konarska A. The relationship between the morphology and structure and the quality of fruits of two pear cultivars (Pyrus communis L.) during their development and maturation. Sci World J. 2013;2013:1–13
- Gao Y, Yang Q, Yan X et al. High-quality genome assembly of 'Cuiguan' pear (Pyrus pyrifolia) as a reference genome for identifying regulatory genes and epigenetic modifications responsible for bud dormancy. Hortic Res. 2021;8:197
- Chen W, Gong L, Guo Z et al. A novel integrated method for large-scale detection, identification, and quantification of widely targeted metabolites: application in the study of rice metabolomics. Mol Plant. 2013;6:1769–80
- Jaeger SR, Andani Z, Wakeling IN et al. Consumer preferences for fresh and aged apples: a cross-cultural comparison. Food Qual Prefer. 1998;9:355–66
- Doty TE. Fructose sweetness: a new dimension. Cereal Foods World. 1976;21:62–3
- Nishio S, Hayashi T, Shirasawa K et al. Genome-wide association study of individual sugar content in fruit of Japanese pear (Pyrus spp.). BMC Plant Biol. 2021;21:378
- Dai M, Shi Z, Xu C. Genome-wide analysis of sorbitol dehydrogenase (SDH) genes and their differential expression in two sand pear (Pyrus pyrifolia) fruits. Int J Mol Sci. 2015;16: 13065–83
- 27. Xu J, Yan J, Li W et al. Integrative analyses of widely targeted metabolic profiling and transcriptome data reveals molecular insight into metabolomic variations during apple (Malus domestica) fruit development and ripening. Int J Mol Sci. 2020;21:4797
- Jia Y, Wong DC, Sweetman C et al. New insights into the evolutionary history of plant sorbitol dehydrogenase. BMC Plant Biol. 2015;15:101
- Shi XP, Ren JJ, Yu Q et al. Overexpression of SDH confers tolerance to salt and osmotic stress, but decreases ABA sensitivity in Arabidopsis. Plant Biol (Stuttg). 2018;20:327–37

- Liu X, Fan HM, Liu DH et al. Transcriptome and metabolome analyses provide insights into the watercore disorder on "Akibae" pear fruit. Int J Mol Sci. 2021;22:4911
- 31. Pleyerová I, Hamet J, Konradova H *et al*. Versatile roles of sorbitol in higher plants: luxury resource, effective defender or something else? *Planta*. 2022;**256**:13
- Zimmermann MH, Ziegler H. List of sugars and sugar alcohols in sieve-tube exudates. In: Zimmermann MH, Milburn JA (eds.), Transport in Plants I: Phloem Transport. Springer-Verlag: Berlin, 1975,158–92
- de Sousa SM, Paniago MG, Arruda P et al. Sugar levels modulate sorbitol dehydrogenase expression in maize. Plant Mol Biol. 2008;68:203–13
- Deguchi M, Bennett AB, Yamaki S et al. An engineered sorbitol cycle alters sugar composition, not growth, in transformed tobacco. Plant Cell Environ. 2006;29:1980–8
- Meng D, Li C, Park HJ et al. Sorbitol modulates resistance to Alternaria alternata by regulating the expression of an NLR resistance gene in apple. Plant Cell. 2018;30:1562–81
- 36. Meng D, He M, Bai Y et al. Decreased sorbitol synthesis leads to abnormal stamen development and reduced pollen tube growth via an MYB transcription factor, MdMYB39L, in apple (Malus domestica). New Phytol. 2018;217:641–56
- 37. He X, Meng H, Wang H et al. Quantitative proteomic sequencing of F₁ hybrid populations reveals the function of sorbitol in apple resistance to Botryosphaeria dothidea. Hortic Res. 2022;9: uhac115
- Yu W, Peng F, Wang W et al. SnRK1 phosphorylation of SDH positively regulates sorbitol metabolism and promotes sugar accumulation in peach fruit. Tree Physiol. 2021;41: 1077–86
- Meng D, Cao H, Yang Q et al. SnRK1 kinase-mediated phosphorylation of transcription factor bZIP39 regulates sorbitol metabolism in apple. Plant Physiol. 2023;192:2123–42
- Guo J, Carrington Y, Alber A et al. Molecular characterization of quinate and shikimate metabolism in *Populus trichocarpa*. J Biol Chem. 2014;289:23846–58
- Etienne A, Génard M, Lobit P et al. What controls fleshy fruit acidity? A review of malate and citrate accumulation in fruit cells. J Exp Bot. 2013;64:1451–69
- Morgan MJ, Osorio S, Gehl B et al. Metabolic engineering of tomato fruit organic acid content guided by biochemical analysis of an introgression line. Plant Physiol. 2013;161: 397–407
- Huang XY, Wang CK, Zhao YW et al. Mechanisms and regulation of organic acid accumulation in plant vacuoles. Hortic Res. 2021;8:227
- 44. Zhang L, Ma B, Wang C *et al*. MdWRKY126 modulates malate accumulation in apple fruit by regulating cytosolic malate dehydrogenase (MdMDH5). Plant Physiol. 2022;**188**:2059–72
- 45. Liu W, Wang Q, Zhang R et al. Rootstock-scion exchanging mRNAs participate in the pathways of amino acids and fatty acid metabolism in cucumber under early chilling stress. Hortic Res. 2022;9:uhac031

- Jiang L, Geng D, Zhi F et al. A genome-wide association study provides insights into fatty acid synthesis and metabolism in Malus fruits. J Exp Bot. 2022;**73**:7467–76
- Yoo H, Shrivastava S, Lynch JH et al. Overexpression of arogenate dehydratase reveals an upstream point of metabolic control in phenylalanine biosynthesis. Plant J. 2021;108:737–51
- 48. Dhakarey R, Yaritz U, Tian L et al. A Myb transcription factor, PgMyb308-like, enhances the level of shikimate, aromatic amino acids, and lignins, but represses the synthesis of flavonoids and hydrolyzable tannins, in pomegranate (Punica granatum L.). Hortic Res. 2022;9:uhac008
- Finn RD, Bateman A, Clements J et al. Pfam: the protein families database. Nucleic Acids Res. 2014;42:D222–30
- Letunic I, Doerks T, Bork P. SMART 6: recent updates and new developments. Nucleic Acids Res. 2009;37:D229–32
- Chenna R, Sugawara H, Koike T et al. Multiple sequence alignment with the Clustal series of programs. Nucleic Acids Res. 2003;31:3497–500
- 52. Waterhouse AM, Procter JB, Martin DM *et al.* Jalview version 2 a multiple sequence alignment editor and analysis workbench. *Bioinformatics.* 2009;**25**:1189–91
- 53. Yao G, Ming M, Allan AC et al. Map-based cloning of the pear gene MYB114 identifies an interaction with other transcription factors to coordinately regulate fruit anthocyanin biosynthesis. Plant J. 2017;92:437–51
- 54. Chen W, Wang W, Peng M et al. Comparative and parallel genome-wide association studies for metabolic and agronomic traits in cereals. Nat Commun. 2016;**7**:12767
- 55. Wang H, Stier G, Lin J et al. Transcriptome changes associated with delayed flower senescence on transgenic petunia by inducing expression of etr1-1, a mutant ethylene receptor. PLoS One. 2013;8:e65800
- Kim D, Langmead B, Salzberg SL. HISAT: a fast spliced aligner with low memory requirements. Nat Methods. 2015;12: 357–60
- 57. Pertea M, Pertea GM, Antonescu CM *et al*. StringTie enables improved reconstruction of a transcriptome from RNA-seq reads. Nat Biotechnol. 2015;**33**:290–5
- Robinson MD, McCarthy DJ, Smyth GK. edgeR: a bioconductor package for differential expression analysis of digital gene expression data. *Bioinformatics*. 2010;26:139–40
- Young MD, Wakefield MJ, Smyth GK et al. Gene ontology analysis for RNA-seq: accounting for selection bias. Genome Biol. 2010;11:R14
- Kanehisa M, Sato Y, Kawashima M et al. KEGG as a reference resource for gene and protein annotation. Nucleic Acids Res. 2016;44:D457–62
- Wang H, Lin J, Chang Y *et al.* Comparative transcriptomic analysis reveals that ethylene/H2O2-mediated hypersensitive response and programmed cell death determine the compatible interaction of sand pear and Alternaria alternata. *Front Plant Sci.* 2017;8:195
- 62. Rio DC. Electrophoretic mobility shift assays for RNA-protein complexes. Cold Spring Harb Protoc. 2014;**2014**:435–40