REVEILLE Transcription Factors Contribute to the Nighttime Accumulation of Anthocyanins in 'Red Zaosu' (Pyrus Bretschneideri Rehd.) Pear Fruit Skin

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Received: 3 January 2020; Accepted: 24 February 2020; Published: 27 February 2020

Abstract: Anthocyanin biosynthesis exhibits a rhythmic oscillation pattern in some plants. To investigate the correlation between the oscillatory regulatory network and anthocyanin biosynthesis in pear, the anthocyanin accumulation and the expression patterns of anthocyanin late biosynthetic genes (ALBGs) were investigated in fruit skin of 'Red Zaosu' (Pyrus Bretschneideri Rehd.). The anthocyanin accumulated mainly during the night over three continuous days in the fruit skin, and the ALBGs’ expression patterns in 'Red Zaosu' fruit skin were oscillatory. However, the expression levels of typical anthocyanin-related transcription factors did not follow this pattern. Here, we found that the expression patterns of four PbREVEILLEs (PbRVEs), members of a class of atypical anthocyanin-regulated MYBs, were consistent with those of ALBGs in 'Red Zaosu' fruit skin over three continuous days. Additionally, transient expression assays indicated that the four PbRVEs promoted anthocyanin biosynthesis by regulating the expression of the anthocyanin biosynthetic genes encoding dihydroflavonol-4-reductase (DFR) and anthocyanidin synthase (ANS) in red pear fruit skin, which was verified using a dual-luciferase reporter assay. Moreover, a yeast one-hybrid assay indicated that PbRVE1a, 1b and 7 directly bound to PbDFR and PbANS promoters. Thus, PbRVEs promote anthocyanin accumulation at night by up-regulating the expression levels of PbDFR and PbANS in 'Red Zaosu' fruit skin.

Keywords: anthocyanin accumulation; night; red pear fruit; REVEILLEs; transient assays

1. Introduction

Anthocyanins are a group of water-soluble flavonoid metabolites that exist widely in plants [1]. Anthocyanins play various roles in plant growth and development [2–5]. In plants, the antioxidant capacities of anthocyanins rely on the extent of B-ring hydroxylation, the type and degree of acylation and glycosylation [2–4]. Anthocyanins play important roles in promoting plant reproduction by transmitting bright colors to pollinators and seed spreaders [2,5]. Additionally, anthocyanins have human health-related benefits [6–9]. Therefore, investigating anthocyanin accumulation in fruit is worthwhile.

In A. thaliana, the anthocyanin biosynthetic pathway is regulated by multiple enzymes, including early and late biosynthetic genes. The early biosynthetic genes include chalcone synthase, chalcone
isomerase, flavanone-3-hydroxylase and flavonoid 3’-hydroxylase, while the late biosynthetic genes include dihydroflavonol 4-reductase (DFR), anthocyanidin synthase (ANS) and UDP-glucoside: flavonoid glucosyltransferase (UGFT) [1,10,11]. Many transcription factors (TFs), such as MYBs, bHLHs and WD40, also participate in the anthocyanin biosynthetic pathway. R2R3–MYB proteins interact with bHLH proteins, such as TRANSPARENT TESTA8, GLABRA3 and ENHANCER OF GLABRA3, and form MYB–bHLH–WDR (MBW) complexes with the WD40 protein TRANSPARENT TESTA GLABRA1 [12–14]. MBW complexes participate in developmental regulation, physiological regulation, trichome formation, seed coat differentiation and the flavonoid biosynthetic pathway in plants [11]. This ternary, complex protein regulates anthocyanin biosynthesis in most plants [12–15]. In the dicot A. thaliana, the late biosynthetic genes are activated by the MBW complex [12–14]. In the monocot maize, the anthocyanin biosynthetic genes are activated as a unit by the MBW complex [15]. Additionally, some typical R2R3–MYB and bHLH TFs act as anthocyanin-related activators in various plants, such as A. thaliana, apple, grape and pear [1,16–22]. In A. thaliana, the MYB TFs PAP1 (AtMYB75), PAP2 (AtMYB90), AtMYB113 and AtMYB114 have been described as positive regulators in anthocyanin accumulation [1,16,17]. In apple, anthocyanin biosynthesis is promoted by MdbHLH3, MdMYB88 and MdMYB124 under low night-temperature conditions [18,19]. MYB10 and MYB10b function as activate regulators in the anthocyanin biosynthetic pathway of pear fruit [20–22].

In plants, the circadian system coordinates physiology and metabolism with the most appropriate or favorable time of day or season [23]. Plants can save energy and resources by regulating reaction times; therefore, the circadian system is crucial to the health and survival of plants. The anthocyanin biosynthetic pattern has diurnal oscillations in some plants. In A. thaliana, the expression levels of phenylalanine ammonia lyase, chalone synthase, chalcone isomerase, and DFR are regulated by a circadian clock [24]. MYBL2 and MYBD are regulated by circadian rhythms and involved in the anthocyanin biosynthetic pathway of A. thaliana [25,26]. Single MYB TFs, named REVEILLES (LHY-CCA1-LIKE) act as important regulators of circadian clockwork [27]. Moreover, in A. thaliana, an atypical MYB (single MYB TF) named AtRVE8 regulates anthocyanin biosynthesis by binding directly to the promoters of anthocyanin structural genes, and AtRVE8 also regulates the expression levels of genes in response to diurnal oscillations [27]. However, the rhythmic regulation of anthocyanin biosynthesis has remains largely unknown in most fruit.

In this study, we isolated four RVEs of ‘Red Zaosu’ pear (Pyrus bretschneideri Rehd.) because of the linkage between their expression patterns and nighttime increases in anthocyanin biosynthesis. Moreover, we found that the expression levels of the four RVEs exhibited rhythmic oscillation patterns in ‘Red Zaosu’ fruit skin. Then, we investigated the functions of the PbRVEs in anthocyanin accumulation in pear fruit skin. This study confirmed that PbRVEs promote anthocyanin accumulation by up-regulating the expression levels of PbDFR and PbANS in pear fruit skin.

2. Results

2.1. The Anthocyanin Content Oscillated Diurnally and Mainly Increased over Night in ‘Red Zaosu’ Fruit Skin

To determine whether anthocyanin accumulates during the daytime or nighttime in red-skinned pear fruit, the anthocyanin content of ‘Red Zaosu’ fruit skin was measured during the daytime (from sunrise to sunset). Moreover, to accurately observe the anthocyanin accumulation in pear fruit skin during a short period of time, color-faded bagged fruit were used because of the low background level of anthocyanin. Color-faded fruit of ‘Red Zaosu’ were exposed to sunlight to re-accumulate anthocyanin over three continuous days. The anthocyanin content in ‘Red Zaosu’ fruit skin rhythmically increased after sunrise and then decreased from noon to sunset (Figure 1a). However, the anthocyanin content between sunrise and sunset was not significantly different during the course of a day. This phenomenon was also observed in other three red pear varieties (Supplementary Figure S1a). However, a significant increase in the anthocyanin content of ‘Red Zaosu’ fruit skin was detected from sunset to the next
sunrise over three continuous days (Figure 1a). Thus, the anthocyanin content in the skin of red pear fruit mainly accumulated during the night.

![Image of daily change patterns in anthocyanin content and expression patterns](image)

Figure 1. Daily change patterns in anthocyanin content, and the expression patterns of anthocyanin late biosynthetic genes (ALBGs) and a typical anthocyanin transport gene, PbGSTF12, in ‘Red Zaosu’ pear fruit skin. (a) The anthocyanin content in the fruit skin of ‘Red Zaosu’ over three continuous days. Images show the ‘Red Zaosu’ phenotypes at three time points over the three continuous days. (b) The expression patterns of ALBGs and PbGSTF12 in ‘Red Zaosu’ fruit skin over three continuous days. The gray boxes indicate nighttime. The NCBI accession numbers of the ALBGs and PbGSTF12 are listed in Supplementary Table S1. HAS: hours after sunrise of day 1. Error bars represent the standard errors (SEs) of the means (n = 3). Data in (a) was determined using a one-way analysis of variance (p < 0.05); the significant differences are indicated by different lowercase letters.

We further investigated the expression patterns of the anthocyanin late biosynthetic genes (ALBGs), including PbANS, PbDFR and PbUFGT, and a typical anthocyanin transporter gene, PbGSTF12, in the fruit skin of ‘Red Zaosu’ over three continuous days. The expression levels of the ALBGs in the fruit skin of ‘Red Zaosu’ rhythmically decreased from sunrise to sunset and then increased until the next sunrise over the three continuous days (Figure 1b). The significant nighttime increase in expression was observed for each of the ALBGs, but not for PbGSTF12, over the three continuous days (Figure 1b). Thus, the accumulation of anthocyanin in ‘Red Zaosu’ fruit skin occurred mainly during the night rather than during the day.
2.2. The Expression Patterns of Candidate REVEILLE (RVE) TFs Correlated with the Nighttime Increase in the Anthocyanin Level in ‘Red Zaosu’ Fruit Skin

To identify the candidate regulators of the nighttime increases in anthocyanin in ‘Red Zaosu’ pear skin, the expression levels of typical anthocyanin-related TFs, including PbMYB9, PbMYB10, PbMYB10b, PbbHLH3, PbbHLH33a and PbbHLH33b, were initially investigated in the fruit skin of ‘Red Zaosu’ over three continuous days (Figure 2). Surprisingly, none of these TFs showed an expression pattern similar to that of the ALBGs (Figure 2). Therefore, we further focused on the RVEs because of the linkage between their expression patterns and anthocyanin biosynthesis [27].

![Figure 2](image_url)

**Figure 2.** The expression patterns of typical anthocyanin-related transcription factors in ‘Red Zaosu’ pear fruit skin. The expression patterns of PbMYB9, PbMYB10, PbMYB10b, PbbHLH3, PbbHLH33a and PbbHLH33b in ‘Red Zaosu’ fruit skin over three continuous days. The gray boxes indicate nighttime. HAS: hours after sunrise of day 1. The NCBI accession numbers of the typical transcription factors are listed in Supplementary Table S1. Error bars represent SEs of the means (n = 3).

Seven selected pear RVEs were isolated from the Chinese pear genome [22] (http://peargenome.njau.edu.cn/, March 1, 2018). To analyze the relationship between PbRVE and AtRVE proteins, a phylogenetic tree was constructed using the Neighbor-Joining method (Figure 3a). These PbRVE proteins were classified into two subgroups. The PbRVE1s and PbRVE7 clustered into type I, while the PbRVE3s, PbRVE6 and PbRVE8 clustered into type II (Figure 3a).
Figure 3. Phylogenetic analysis of REVEILLEs (RVEs) and the expression patterns of PbRVEs in ‘Red Zaosu’ pear fruit skin. (a) Phylogenetic analysis of RVE proteins from pear and A. thaliana. The type I proteins are AtRVE7, AtRVE7-Like, PbRVE7, AtRVE1, AtRVE2, PbRVE1a and PbRVE1b; the type II proteins are AtRVE3, AtRVE5, PbRVE3a, PbRVE3b, AtRVE6, PbRVE6, PbRVE8, AtRVE4 and AtRVE8. (b) The expression patterns of PbRVEs in ‘Red Zaosu’ fruit skin at sunrise and sunset over three continuous days. HAS: hours after sunrise of day 1. The actual transcript abundance data of PbRVEs are listed in Supplementary Table S3. (c) The expression patterns of PbRVE1a, 1b, 7 and 8 in ‘Red Zaosu’ fruit skin over three continuous days. The gray boxes indicate nighttime. (d) Multiple sequence alignment of RVE proteins. RVE proteins were aligned using ClustalW. The DNA-binding domain is indicated by a horizontal gray bar. The phylogenetic analysis was constructed with the Neighbor-joining method (1000 replications bootstrap test and JTT model distribution) using MEGA 7.0. The protein sequences of the RVEs were obtained from the pear genome. The gene accession numbers are listed in Supplementary Table S1. Error bars represent SEs of the means (n = 3).

To identify which PbRVEs participated in anthocyanin accumulation during the night, we analyzed the expression patterns of PbRVEs in the skin of ‘Red Zaosu’ fruit at sunrise and sunset over the three
continuous days. The expression levels of PbRVE1a, 1b, 7 and 8, but not PbRVE3a, 3b and 6, significantly increased during the nighttime in skin of ‘Red Zaosu’ fruit (Figure 3b). Therefore, the expression levels of PbRVE1a, 1b, 7 and 8 in the skin of ‘Red Zaosu’ fruit were further investigated over three continuous days. PbRVE1a, 1b, 7 and 8 expression levels peaked at dawn and then decreased until sunset during each day (Figure 3c). Furthermore, this expression pattern was also found in the skins of other red pear cultivars (Supplementary Figure S1b). Moreover, at sunrise and sunset of each of three continuous days, significant correlations were observed between PbRVE and ALBG expression levels in ‘Red Zaosu’ fruit skin, while the expression levels of PbMYB9, PbMYB10, PbMYB10b and PbHLH33a were only slightly correlated with those of PbANS, PbDFR and PbUFGT (Table 1). A multiple-alignment showed that all the candidate PbRVEs and AtrVEs contained the conserved SHAQK[Y/F]F motif in the DNA-binding domains of their N-terminal regions (Figure 3d). Thus, PbRVE1a, 1b, 7 and 8 were selected as candidate genes because of the high correlations between their expression and the expression of ALBGs in pear fruit skin.

| Table 1. Correlation analysis between the expression of anthocyanin-related transcription factors (TFs) and the expression of ALBGs in pear fruit. |
|----------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|
| Genes           | PbMYB9 | PbMYB10 | PbMYB10b | PbHLH33a | PbRVE1a | PbRVE1b | PbRVE7 | PbRVE8 |
| PbANS           | 0.441 ** | 0.22 | −0.01 | 0.07 | 0.750 ** | 0.775 ** | 0.728 ** | 0.757 ** |
| PbDFR           | 0.416 ** | 0.17 | −0.02 | 0.08 | 0.734 ** | 0.782 ** | 0.746 ** | 0.777 ** |
| PbUFGT          | 0.355 * | 0.335 * | 0.19 | 0.06 | 0.779 ** | 0.859 ** | 0.818 ** | 0.855 ** |

The pairwise correlation coefficients between the expression levels of anthocyanin-related TFs and those of ALBGs in the fruit skin of ‘Red Zaosu’ pear over three continuous days. The NCBI accessions of the anthocyanin-related genes are listed in Supplementary Table S1. The significant correlation coefficients are indicated in bolded values. The data was analyzed using SPSS 20. *: Correlation significant at the 0.05 level (p < 0.05, two-tailed); **: Correlation significant at the 0.01 level (p < 0.01, two-tailed).

2.3. Overexpression of PbRVEs in ‘Zaosu’ Pear Fruit Promoted Anthocyanin Accumulation

To investigate the bio-functions of PbRVEs in anthocyanin regulation, these candidates were transiently overexpressed using agrobacterium-infiltration of the skin of ‘Zaosu’ fruitlets. The validity of the ‘Zaosu’ fruitlets’ infection was validated by monitoring the β-glucuronidase gene (GUS) signal (Supplementary Figure S2). The transient overexpression of PbRVE1a, 1b, 7 and 8 independently in ‘Zaosu’ fruitlet skins increased the anthocyanin accumulation (Figure 4). However, the promotive efficiency among these PbRVEs varied (Figure 4a,b). Consequently, the overexpression of PbRVE1b resulted in an intense pigmentation of the pear fruitlet skins (4.25 times darker than controls), whereas lighter pigmentation was observed when PbRVE1a, 7, and 8 were independently overexpressed in pear fruitlet skins (from 2.10 to 2.84 times darker than controls) (Figure 4c). Additionally, the virus-induced gene silencing (VIGS) of PbRVE1a, 1b, 7 and 8 independently in ‘Palacer’ (P. communis L.) fruitlet skins decreased the anthocyanin accumulation (Supplementary Figure S3). The expression levels of PbDFR and PbANS significantly increased in PbRVEs-overexpression (OE) ‘Zaosu’ fruitlet skins, especially PbRVE1b-OE ‘Zaosu’ fruitlet skins, but the expression level of PbUFGT was not affected (Figure 4c). Thus, PbRVEs promote the expression levels of PbDFR and PbANS to increase the anthocyanin accumulation in ‘Zaosu’ pear fruit.
Figure 4. Anthocyanin biosynthetic patterns in RVE-overexpression. (a) Phenotypes of ‘Zaosu’ fruitlet skins transiently overexpressing PbRVE1a, 1b, 7 and 8. (b) The overexpression levels of PbRVE1a, 1b, 7 and 8 in ‘Zaosu’ fruitlet skins. (c) The anthocyanin contents and the expression patterns of ALBGs in ‘Zaosu’ fruitlet skins which overexpressing PbRVEs. The number means the ratio of anthocyanin content in ‘Zaosu’ fruitlets skin overexpressing PbRVEs compared with the empty vector. OE: overexpression. The significant differences are based on comparisons with the empty vector. Error bars indicate the SEs of the means (n = 3). Data in (c) were analyzed using Student’s t-test: * p < 0.05, ** p < 0.01.

2.4. PbRVE1a, 1b, 7 and 8 Promoted Anthocyanin Accumulation by Activating the Promoters of ALBGs in ‘Red Zaosu’ Pear Fruit

To determine whether PbRVEs regulated PbDFR, PbANS and PbUFGT directly, yeast one-hybrid (Y1H) tests were conducted. PbRVE1a, 1b, and 7 bound directly to the PbDFR and PbANS promoters (Figure 5a). However, direct interactions between the PbRVEs and PbUFGT were not detected using the Y1H test (Figure 5a). To determine the effects of PbRVEs on PbDFR, PbANS and PbUFGT, the promoter regions of PbDFR, PbANS and PbUFGT were used in a dual-luciferase assay system in Nicotiana benthamiana leaves. Infiltration with PbRVEs activated the promoters of PbDFR, PbANS and PbUFGT, and PbRVE1b showed a strongly ability to activate the promoters of PbDFR, PbANS and PbUFGT (Figure 5b). Thus, PbRVEs appear to activate directly the promoters of PbDFR, PbANS and PbUFGT, resulting in higher anthocyanin accumulations.
Figure 5. ALBG promoters are bound and stimulated by PbRVEs. (a) Yeast one-hybrid assays of PbRVEs with the PbDFR, PbANS and PbUFGT promoters. (b) Validation of the activation effects of PbRVEs on the PbDFR, PbANS and PbUFGT promoters using a dual-luciferase assay in Nicotiana benthamiana leaves. The relative promoter activity is represented by the expression ratio of the structural luciferase (LUC) gene to the 35S Renilla (REN) gene. The ratio was further standardized based on the LUC/REN value of the empty vector. Results represent the means of five biological replicates. All significant differences are based on comparisons with the control. Error bars show the SEs of the means (n = 5).

3. Discussion

3.1. PbRVE1a, 1b, 7 and 8 Expression Levels Correlated with Anthocyanin Accumulation during the Nighttime in ‘Red Zaosu’ Pear Fruit Skin

The typical anthocyanin biosynthesis-regulating TFs, such as MYB10 and HY5, play important roles in the anthocyanin biosynthetic pathways of fruit [1,21,28]. In A. thaliana, HY5 promotes anthocyanin biosynthesis by binding directly to the promoter regions of ALBGs, such as DFR, LDOX and UF3GT [1]. MYB10 positively activates DFR in the anthocyanin biosynthesis of apple [28,29]. MYB10 and MYB10b have positive functions in regulating anthocyanin biosynthesis and accumulation in pear [20,30,31]. However, daily fluctuations in expression were not observed for the typical anthocyanin-related TFs involved in anthocyanin biosynthesis (Figure 2). Therefore, these typical anthocyanin-related TFs are not the main elements active in the anthocyanin biosynthesis pathway during the nighttime in ‘Red Zaosu’ fruit skin. Consequently, in this study, we investigated the TFs involved in the overnight accumulation of anthocyanins in ‘Red Zaosu’ fruit skin.

The single MYB-like TF, AtRVE8, has been identified as an activator in the anthocyanin biosynthetic pathway of A. thaliana [27]. Based on the phylogenetic tree and expression patterns analysis between PbRVEs and AtRVEs, PbRVE1a, 1b, 7 and 8 were selected for further investigation (Figure 3a,b). The expression patterns of PbRVE1a, 1b, 7 and 8 showed diurnal oscillations and increased during
the nighttime in ‘Red Zaosu’ fruit (Figure 3c). In other red pears, anthocyanin did not accumulate in the daytime (Supplementary Figure S1a). Moreover, the expression patterns of PbRVE1a, 1b, 7 and 8 peaked near dawn in red pear fruit skin (Figure 3c, Supplementary Figure S1b). This result was consistent with the occurrence of anthocyanin accumulation during the nighttime in ‘Red Zaosu’ pear fruit skin (Figure 1a). The data indicate that PbRVE1a, PbRVE1b, PbRVE7 and PbRVE8 are potential TFs involved in the nighttime increase in anthocyanin accumulation in ‘Red Zaosu’ pear fruit.

3.2. PbRVEs Promoted Anthocyanin Accumulation by Up-Regulating the Expression Levels of PbDFR and PbANS in Pear Fruit Skin

ALBGs (such as DFR, ANS and UFGT) are involved in the anthocyanin biosynthetic pathways of fruits [15]. The expression levels of anthocyanin-related structural genes exhibit diurnal oscillation patterns and appear to be regulated by the circadian clock in *A. thaliana* [24]. Additionally, in *A. thaliana*, the expression levels of the anthocyanin-related genes appear to change during light/dark cycles [27]. *AtRVE8* up-regulates the expression of anthocyanin biosynthetic genes in RVE8-OE *A. thaliana* plants [27]. In this study, the expression levels of *PbDFR*, *PbANS* and *PbUFGT* exhibited diurnal oscillations and increased during the night in ‘Red Zaosu’ pear fruit skin (Figure 1b). The transient over-expression assay showed that the *PbRVEs* had different abilities to up-regulate the expression levels of structural genes (Figure 4). According to the transient overexpression assay, *PbRVE1b* had a stronger ability than the other *RVEs* to increase anthocyanin biosynthesis in pear fruitlet skins (Figure 4a,b). Furthermore, transient VIGS assays indicated that anthocyanin did not accumulate when *PbRVE1b* was silenced in ‘Palacer’ pear fruitlet skin (Supplementary Figure S3).

In *A. thaliana*, *RVE8* directly binds and regulates the expression of anthocyanin structural gene promoters in response to the diurnal oscillation in anthocyanin accumulation [27,32]. However, the Y1H assay verified that *PbRVE1a*, *PbRVE1b* and *PbRVE7*, but not *PbRVE8*, bound directly to the promoters of *PbDFR* and *PbANS* (Figure 5a). The function of *PbRVE8* in binding to the promoters of *PbDFR* and *PbANS* in pear has been precluded by other proteins [27]. In this study, the correlation analysis indicated that the expression levels of *PbRVEs* were significantly positively correlated with the expression levels of *PbDFR* and *PbANS* (Table 1). Thus, we inferred that *PbDFR* and *PbANS* are directly downstream factors of *PbRVE1a*, *PbRVE1b* and *PbRVE7* in ‘Red Zaosu’ pear.

Using the dual-luciferase assay, we determined that the expression of *PbRVE1b* increases the activities of the *PbDFR*, *PbANS* and *PbUFGT* promoters (Figure 5b). However, *PbRVE1a* and *PbRVE1b* did not bind directly with the *PbUFGT* promoter (Figure 5a). This result was consistent with the transient overexpression of the *PbRVEs* (Figure 4c). We speculated that *PbRVE1a* and *PbRVE1b* do not directly affect the *PbUFGT* promoter in pear. In *N. benthamiana* leaves, *RVE1a* and *RVE1b* may interact with other factors to activate the *UFGT* promoter [33–35]. Thus, *PbRVE1a*, *PbRVE1b* and *PbRVE7* increase anthocyanin accumulation by directly binding and activating *PbDFR* and *PbANS* in pear fruit.

4. Materials and Methods

4.1. Plant Material and Treatments

The ‘Red Zaosu’ pear (*P. bretschneideri* Rehd.) is a bud sport of ‘Zaosu’ pear and has characteristic red fruit and leaves. The regulatory mechanism of anthocyanin biosynthesis in ‘Red Zaosu’ has been studied [20,36]. Therefore, we chose ‘Red Zaosu’ to investigate the diurnal accumulation of anthocyanin.

The fruit of ‘Red Zaosu’ was selected from a commercial orchard in Mei County, Shaanxi Province, China, in 2018. The fruit of ‘Red Zaosu’ and ‘Palacer’ (*P. communis* L.) were selected at approximately 40 d after flower blossom and bagged for 30 d until the red pigments totally faded. Then, the fruit of ‘Red Zaosu’ was exposed to daylight for three continuous days. The experiment was conducted on 12–14 June 2018. Additionally, the fruit of ‘Red Zaosu’ were harvested at 0, 3, 6, 9, 12, 24, 27, 30, 33, 36, 48, 50, 54, 57 and 60 hours after sunrise of day 1 (HAS). The faded ‘Palacer’ fruitlets were used for the
PbRVE virus-induced gene silencing (VIGS) assay. The skins of these harvested fruit were frozen in liquid nitrogen and stored at −80 °C for the subsequent measurement of the anthocyanin content and RNA extraction.

For the dual-luciferase assay infiltration, *N. benthamiana* seedlings were grown in a light incubator (16-h light/8-h dark) at 22 °C.

4.2. Anthocyanin Content Measurements

The pH differential method was used to measure the total anthocyanin contents of red skin pear fruitlets [37]. The extraction of total anthocyanins was performed using a previously reported method, with slight modifications [38,39]. Approximately 0.2 g samples of fruit skin were powderd in liquid nitrogen and mixed with PVP-K30 (Sigma, St. Louis, MI, USA), and then 1.5 mL of 1% HCL–methanol was added to the mixed sample. After centrifugation at 4 °C and 12,000× g for 5 min, 200-µL aliquots of the supernatant were transferred separately to two clear tubes for dilution. One was diluted with 400 μL 0.025 M potassium chloride buffer (pH 1.0), and the other with 400 μL 0.4 M sodium acetate buffer (pH 4.5). These solutions were placed for 15 min in the dark at room temperature before the absorbance values were measured synchronously at 520 nm and 700 nm using a Microporous plate spectrophotometer (Multiskan GO; Thermo Scientific, Waltham, MA, USA).

4.3. Isolation of RVE Genes and Their Phylogenetic Analysis

The sequences of selected pear RVEs were isolated from pear databases [22] (http://peargenome.njau.edu.cn/, access date 1 March 2018). The RVEs from pear and *A. thaliana* were aligned using ClustalW (MEGA 7.0, The Biodisgn Institute, Arizona State University, AZ, USA) [20]. The phylogenetic analysis was performed with the Minimum-Evolution method and the JTT model using the MEGA 7.0 program (The Biodisgn Institute, Arizona State University, AZ, USA) [20]. The GenBank accession numbers for the functionally labelled RVEs are listed in Supplementary Table S1. The complete coding DNA sequences (CDSs) of candidate RVEs were cloned using PrimeSTAR Max Premix (TaKaRa, Dalian, China) and gene-specific primers (Supplementary Table S2) from ‘Red Zaosu’ cDNA sources.

4.4. RNA Isolation and an Expression Analysis Using Quantitative Real-Time PCR (qRT-PCR)

The total RNA of skins was extracted using the RNA prep Pure Plant Kit (Tiangen, Beijing, China). The RNA concentration and quality were detected by UV spectrophotometry and a 0.8% agar gel. In total, 1 µg of total RNA was reverse-transcribed to cDNA using the PrimeScript RT reagent kit with gDNA Eraser (TaKaRa, Dalian, China). The primers used for qRT-PCR were designed with Oligo 7 software (Molecular Biology Insights, Inc., Colorado Springs, USA) and synthesized by AuGCT Biotechnology Synthesis Lab (Beijing, China). The qRT-PCR was performed on an Applied Biosystems StepOnePlus™ Real-Time PCR Systems (Applied Biosystems, Waltham, MA, USA) with TB Green Premix Ex Taq II (Tli RNaseH Plus; TaKaRa, Dalian, China) according to the manufacturer’s instructions. Data were analyzed using the 2−ΔΔCT method. All the qRT-PCR reactions were replicated three times for each biological repeat. The primers for actin, anthocyanin biosynthetic genes and candidate RVEs are listed in Supplementary Table S2.

4.5. Transient Expression Assay in Pear Fruitlet Skins

The complete CDSs of RVE TFs were cloned into the multiple cloning site (MCS) (BamHI–HindIII) of the pGreenII 0029 62-SK binary vector to form PbRVE-OE plasmids (PbRVE1a-OE, PbRVE1b-OE, PbRVE7-OE and PbRVE8-OE) [40]. The complete GUS CDS in the plBl121-GUS plasmid was cloned into the MCS of the pGreenII 0029 62-SK binary vector to form the pGreenII 0029 62-SK-GUS plasmid (described in Supplementary Figure S4a). The 400–600-bp fragments of the C-termini of RVE TFs were inserted into the MCS (BamHI--XhoI) of pTRV2 to form PbRVE VIGS vectors (PbRVE1a-TRV, PbRVE1b-TRV, PbRVE7-TRV and PbRVE8-TRV, described in Supplementary Figure S4b). The primers for amplifying the sequences are described in Supplementary Table S2. *Agrobacterium tumefaciens* strain
EHA105 independently containing the constructed plasmids was grown at 28 °C on Luria–Bertani (LB) solid medium supplemented with 50 mg/L kanamycin and 25 mg/L rifampicin. After incubating for 48 h, the A. tumefaciens was resuspended in the infiltration buffer (10 mM MgCl₂, 10 mM MES and 200 µM acetosyringone) and shaken for 3–4 h (up to a final OD600 of 0.8) at room temperature before being injected into pear fruitlet skins. The pear injection process was as described in Spolaore et al. [41] and the injection volume was as described in Zhai et al. [20]. The negative controls were infiltrated with A. tumefaciens containing pGreenII 0029 62-SK or the pTRV2 empty vector. The treated fruitlets were harvested at 5 d after injection. For GUS staining, the plant materials were stained with 5-bromo-4-chloro-3-indolyl glucuronide at 37 °C for 12 h as described previously [42].

4.6. Y1H Assay

The Y1H assays were performed following the manufacturer’s instructions for the Matchmaker Gold Yeast One-Hybrid System Kit (Clontech, Mountain View, CA, USA). We ligated independently ~800-bp fragments of the PbANS and PbDFR promoters into pAbAi to construct the pAbAi-baits. Additionally, the complete CDSs of the PbRVEs were separately inserted into the pGADT7 vector to construct the prey-AD vectors. The pAbAi-bait vectors were linearized and transformed into Y1HGold separately. The colonies were selected in the absence of uracil on selective medium-containing agar plates. The correct integration of plasmids into the genome of the Y1HGold yeast was confirmed using a colony PCR analysis (Matchmaker Insert Check PCR Mix 1; Clontech, Mountain View, CA, USA). After determining the minimal inhibitory concentration of Aureobasidin A (AbA) for the bait–reporter yeast strains, the AD-prey vectors were transformed into the bait yeast strains and selected on synthetic dextrose (SD)/−Leu/AbA plates. All the transformations and screenings were performed three times.

4.7. Dual-Luciferase Assay

The promoters of PbANS and PbDFR were amplified from ‘Red Zaosu’ genomic DNA using PrimeSTAR Max Premix (TaKaRa, Dalian, China) and gene-specific primers (Supplementary Table S1). These promoters were cloned into the HindIII and BamHI sites within pGreenII 0800-LUC [40]. The full-length CDS sequences of PbRVE1a, PbRVE1b, PbRVE7 and PbRVE8 were independently cloned into the MCS (BamHI-HindIII) of the pGreenII 0029 62-SK binary vector [40]. Each of these recombinant plasmids and the pSoup helper plasmid were transferred individually into A. tumefaciens strain EHA105 [40]. The A. tumefaciens cells containing PbRVE-SKs were separately mixed with proPbDFR-LUC or proPbANS-LUC at 1:1 ratio before being infiltrated into 4-week-old N. benthamiana leaves. After injection, the plants were grown for 3 d in a light incubator (16-h light/8-h dark) at 22 °C, and then, the treated leaves were collected in 1 × phosphate buffered solution for the dual-luciferase assay. The Firefly luciferase (Luc) to Renilla luciferase (Ren) activity ratios were analyzed using a Dual-Luciferase® Reporter Assay System (Promega, Madison, WI, USA) on a Tecan Infinite M200pro Full-Wavelength Multifunctional Enzyme Labelling Instrument (TECAN, Männedorf, Switzerland). Three independent experiments were carried out with at least five biological replicates per experiment.

4.8. Statistical Analysis

All the experimental data were statistically processed using GraphPad Prism 6 software (GraphPad Software Inc., San Diego, CA, USA) and are shown as means ± standard errors (SEs). Additionally, the significant differences were analyzed using a one-way analysis of variance and Student’s t-tests.

Supplementary Materials: Supplementary materials can be found at http://www.mdpi.com/1422-0067/21/5/1634/s1. Figure S1: The anthocyanin content of red pear fruit and expression patterns of candidate PbRVEs in red pear fruit; Figure S2: The GUS-stained ‘Zaosu’ fruitlets skin infiltrated by pGreen II 62-SK-GUS; Figure S3: Functional analysis of the PbRVEs using VIGS in pear fruitlets skin; Figure S4: Construction of the recombinant plasmid; Table S1: The accession numbers of the genes used in this study; Table S2: List of primers used in this study; Table S3: The actual transcript abundance data of PbRVEs.
Author Contributions: Conceptualization, X.L., R.Z. and L.X.; methodology, X.L., R.Z. and Z.W.; software, X.L., T.W., H.L., Y.W. and Q.S.; validation, X.L., R.Z., Z.W. and L.X.; data curation, X.L., T.W. and H.L.; writing-original draft, X.L.; writing-review and editing, X.L., R.Z., C.Y. and L.X.; project administration, Z.W., C.Y. and L.X.; resources, L.X. and Z.W.; supervision, R.Z., F.M. and L.X.; funding acquisition, L.X. and Z.W. All authors have read and agreed to the published version of the manuscript.

Funding: This research was funded by the National Key R&D Program of China (2019YFD1000104) and the National Natural Science Foundation of China (31572086 and 31972372).

Acknowledgments: We thank Lesley Benyon, from Liwen Bianji, Edanz Group China (www.liwenbianji.cn/ac), for editing the English text of a draft of this manuscript.

Conflicts of Interest: The authors declare no conflict of interest. The funder L.X. had the roles in conceptualization, validation, project administration, resources, supervision and the decision to publish the results. The funder Z.W. had the roles in methodology, validation, writing-review and editing, project administration and resources.

Abbreviations

ANS Anthocyanidin synthase
ALBGs Anthocyanin late biosynthetic genes
CDS Coding DNA sequence
DFR Dihydroflavonol 4-reductase
GUS the β-glucuronidase gene
HAS Hours after sunrise of day 1
LUC Firefly luciferase
MCS Multiple cloning sites
MBW MYB–bHLH–WDR
OE Overexpression
qRT-PCR Quantitative Real-Time PCR
REN Renilla luciferase
RVE REVEILLE
SD Selective synthetic dextrose medium
SEs the standard errors
TF Transcription factor
UFGT UDP-glucoside: flavonoid glucosyltransferase
VIGS Virus-induced gene silencing
Y1H Yeast one-hybrid assay

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