

Novel and Recently Evolved MicroRNA Clusters Regulate Expansive *F-BOX* Gene Networks through Phased Small Interfering RNAs in Wild Diploid Strawberry^{1[OPEN]}

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The wild strawberry (*Fragaria vesca*) has recently emerged as an excellent model for cultivated strawberry (*Fragaria* × *ananassa*) as well as other Rosaceae fruit crops due to its short seed-to-fruit cycle, diploidy, and sequenced genome. Deep sequencing and parallel analysis of RNA ends were used to identify *F. vesca* microRNAs (miRNAs) and their target genes, respectively. Thirty-eight novel and 31 known miRNAs were identified. Many known miRNAs targeted not only conserved mRNA targets but also developed new target genes in *F. vesca*. Significantly, two new clusters of miRNAs were found to collectively target 94 *F-BOX* (*FBX*) genes. One of the miRNAs in the new cluster is 22 nucleotides and triggers phased small interfering RNA production from six *FBX* genes, which amplifies the silencing to additional *FBX* genes. Comparative genomics revealed that the main novel miRNA cluster evolved from duplications of *FBX* genes. Finally, conserved trans-acting siRNA pathways were characterized and confirmed with distinct features. Our work identified novel miRNA-*FBX* networks in *F. vesca* and shed light on the evolution of miRNAs/phased small interfering RNA networks that regulate large gene families in higher plants.

The wild strawberry (*Fragaria vesca*) is an emerging model for the commercial strawberry (*Fragaria* × *ananassa*) as well as other Rosaceae fruit crops, including apple (*Malus domestica*), pear (*Pyrus communis*), peach (*Prunus persica*), and cherry (*Prunus avium*). While many of the rosaceous fruit trees require several years of juvenile growth before flowering, *F. vesca* takes only 3.5 to 5 months from seed to flower and fruit. The small genome size (240 Mb), a sequenced genome, extensive tissue- and stage-specific transcriptomes, diploidy, and ease of transformation make *F. vesca* an ideal model system for investigations into fundamental biological questions and identifications of the regulatory pathways of

important economic traits (Shulaev et al., 2011; Darwish et al., 2013; Kang et al., 2013; Pantazis et al., 2013).

MicroRNAs (miRNAs) represent an important class of regulatory molecules found in plants and animals. Processed from fold-back hairpin precursor RNAs, miRNAs are 20 to 22 nucleotides in length and negatively regulate target genes through homology-directed cleavage or translational inhibition of mRNAs (Bartel, 2004; Mallory and Vaucheret, 2006). Certain miRNAs, in addition, can trigger the biogenesis of another type of regulatory small RNA called trans-acting small interfering RNA (tasiRNA). In *Arabidopsis thaliana*, the noncoding precursor RNA of tasiRNA is first cleaved by a trigger miRNA. One of the cleavage products is then converted into a double-stranded RNA by RNA-DEPENDENT RNA POLYMERASE6. This double-stranded RNA is subsequently cleaved, in progressive 21-nucleotide phased intervals, by DICER-LIKE4 (DCL4) into 21-nucleotide tasiRNAs (Allen et al., 2005; Yoshikawa et al., 2005). Recently, genome-wide small RNA sequencing and analyses of an ever-expanding number of plant species have revealed the production of 21-nucleotide secondary small interfering RNAs (siRNAs) from hundreds of genes through a mechanism like tasiRNAs. These siRNAs may function in trans (tasiRNAs) or in cis (cis-acting siRNAs) to cleave their target genes (Johnson et al., 2009; Zhai et al., 2011; Xia et al., 2012, 2013; Fei et al., 2013). Collectively, these tasiRNAs and cis-acting siRNAs, which are produced via progressive 21-nucleotide cleavage from the trigger miRNA

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cleavage site, are termed phased small interfering RNAs (phasiRNAs). Accordingly, phasiRNA-producing loci are termed *PHAS* loci (Zhai et al., 2011). In dicots, phasiRNAs have been found to be generated from and presumably to regulate large and conserved gene families such as those coding for the NUCLEOTIDE-BINDING AND LEUCINE-RICH REPEAT (NB-LRR) proteins, MYB transcription factors, and PENTATRICOPEPTIDE REPEAT (PPR) proteins (Zhai et al., 2011; Xia et al., 2012, 2013). The vast phasiRNA networks in higher plant genomes are just beginning to be recognized and discovered, and the functional roles of these phasiRNA networks are yet to be illuminated.

One of the most interesting and unique developmental processes in strawberry is its fruit. Unlike other conventional fruits, which usually develop from the ovary of a flower, the fleshy fruit of strawberry develops from the stem tip, the receptacle. The ovary of the strawberry flower (called the achene) instead dries up and dots the surface of the receptacle. Each achene contains a single seed and can be easily removed by manual manipulation. Nitsch (1950) showed that removing all of achenes from the receptacle prevented receptacle fruit enlargement. Furthermore, exogenous auxin application acted as a substitute for achenes and stimulated receptacle fruit growth. This experiment established auxin as an essential phytohormone that promotes strawberry receptacle enlargement and suggested achenes as the source of auxin (Nitsch, 1950). Despite these important early discoveries, not much is known about the underlying molecular mechanisms of strawberry fruit development. Recently, we comprehensively profiled mRNA levels during *F. vesca* flower and fruit development and identified stage- and tissue-specific transcripts, in particular transcripts unique to the receptacle (Kang et al., 2013; Hollender et al., 2014). Analysis of the *F. vesca* fruit transcriptome revealed the endosperm and seed coat as the primary tissues for fertilization-induced auxin and GA biosynthesis. However, it remains unknown whether small RNAs play a role in regulating flower and fruit development and if small RNAs contribute to the success of receptacle fruit formation.

Here, we collected tissues of various developmental stages from *F. vesca* and dissected the fruit into separate fine parts to profile their small RNA population by deep sequencing and computational analyses. We identified two novel miRNA clusters, one bearing eight different miRNAs in an approximately 8-kb genomic region and the other with four copies of the same miRNA. All nine miRNAs were found to target a large number of *F-BOX* (*FBX*) genes, with the majority of them targeting at a conserved region coding for the F-box domain. One of the nine miRNAs is of 22 nucleotides in length, targets at least six *FBX* genes, and initiates secondary phasiRNA production, leading to amplification of the silencing effect. Hence, combining the ability to initiate phasiRNAs with target pairing at the highly conserved F-box domain, these novel miRNAs are particularly effective in regulating large gene families. Moreover, the main miRNA cluster was shown to arise only recently from duplications of the target *FBX* genes in the *F. vesca* genome. Our work defines a new miRNA-*FBX*

regulatory network in *F. vesca* and provides novel insights into the evolution of miRNAs and their mRNA targets.

RESULTS

Identification of Novel and Known miRNA Families in *F. vesca*

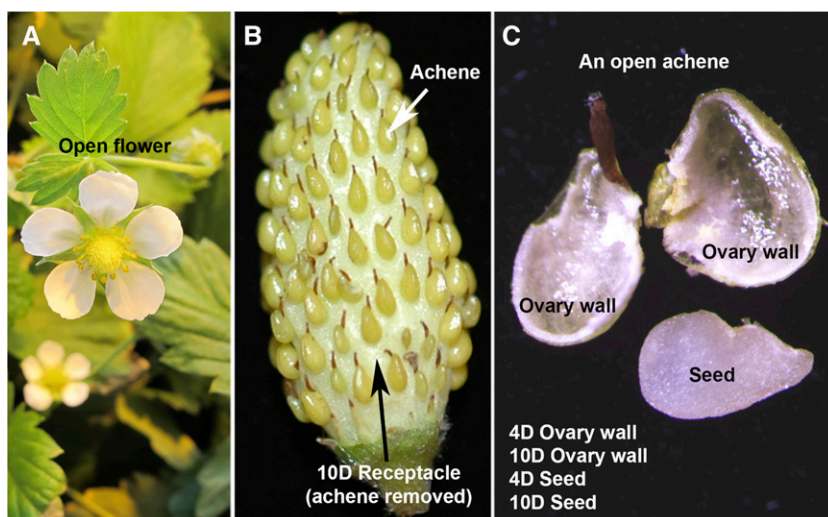
To maximize our chances of identifying *F. vesca* miRNAs, we isolated RNA from vegetative tissues (young leaves and seedlings), flowers (flower buds and newly opened flowers), and fruit tissues that separated achenes from the receptacle (Fig. 1). We further dissected each achene to harvest the ovary wall and the seed separately (Fig. 1C). Since fruit initiation occurs only after successful fertilization, the ovary wall and seed were harvested at 4 days post anthesis (DPA) as well as 10 DPA. Together, small RNAs from nine different tissues/libraries were sequenced to yield 10 to 60 million small RNA sequence reads per library (Supplemental Table S1).

miRNA identification followed an established pipeline (Supplemental Fig. S1). First, we identified 91 *MIRNA* genes, encoding 59 unique miRNA sequences and belonging to 31 known miRNA families (Supplemental Table S2). In general, known miRNA families are conserved across multiple plant species, have many family members, and are of higher abundance. For example, miR156 has eight variants (i.e. eight unique sequences) derived from 15 distinct loci in the genome. To identify novel miRNAs not previously described, all small RNAs with length of 20 to 22 nucleotides were subjected to stringent filtering criteria (Meyers et al., 2008; Supplemental Fig. S1). A total of 33 miRNA loci were identified that had greater than 75% processing accuracy as well as an miRNA star (miRNA*) present in the same library. After moving the three recently reported *F. × ananassa* miRNAs, fve-miR11283, fve-miR11284, and fve-miR11285 (Supplemental Table S2; Li et al., 2013), from our novel list, we have identified 30 novel miRNAs from *F. vesca*. They were submitted to miRBase and named fve-miR11286 to fve-miR11314 (Table I; Supplemental Table S3). Unlike the conserved miRNA families, most of the novel fve-miRNAs are coded by a single *MIRNA* locus and present at a lower abundance (Supplemental Table S3). Eleven of the 30 novel miRNAs are 22 nucleotides in length, which is a typical feature of miRNAs capable of initiating phasiRNA production (Chen et al., 2010; Cuperus et al., 2010).

Newly Evolved Target Genes of Conserved miRNAs

A high-throughput approach, called parallel analysis of RNA ends (PARE), or degradome sequencing, or genome-wide mapping of uncapped and cleaved transcripts (Addo-Quaye et al., 2008; German et al., 2008; Willmann et al., 2014) can experimentally validate target transcripts of miRNAs by capturing mRNA cleavage products. This approach has the advantage of concurrent identification of all cleavage products in the genome

Figure 1. Photographs of *F. vesca* floral and fruit tissues used as the source of small RNAs. A, An open flower. B, An image of a developing fruit, illustrating the relationship between achene, the botanical fruit, and receptacle, the accessory fruit. C, Photograph showing a dissected achene, revealing ovary wall and the white seed, normally enclosed inside the ovary wall. 4D and 10D indicate samples harvested at 4 and 10 DPA, respectively.



as well as a high detection sensitivity, compared with older techniques such as northern blotting or 5' RACE. To maximize the identification of miRNA targets, we dissected *F. vesca* tissues, mirroring the tissues used for small RNA sequencing; we then pooled RNA from these dissected tissues into three different PARE libraries: flower, fruit, and vegetative tissues. In total, approximately 86 million to approximately 108 million high-quality reads from each PARE library were mapped to the *F. vesca* transcriptome (Supplemental Table S1). Cleavage targets of 27 (of the 31) known miRNA families were found, totaling 280 target genes (Supplemental Table S4). As expected, these known miRNA families mainly targeted canonical conserved target genes, most of which encode transcription factors.

Unexpectedly, we found that many known miRNA families gained new targets, perhaps species or lineage specific (Supplemental Table S4). For example, apart from the previously known target genes encoding the growth regulation factor and the basic helix-loop-helix transcription factors (Rodriguez et al., 2010; Debernardi et al., 2012), *fve-miR396* appeared to cleave 19 new target genes (Fig. 2A; Supplemental Table S4). The PARE tags validating the miRNA cleavage sites are shown in T-plots (Fig. 2B). Interestingly, *miR396* and *miR390* acquired as targets *DCL* genes (*gene18596* and *gene15481* in *F. vesca*, respectively; Fig. 2, A and B). Given the prior instances of miRNA regulation of miRNA processing or effector machineries in higher plants (Vaucheret et al., 2004; Zhai et al., 2011), it is not surprising that *F. vesca* also involves miRNAs in potential small RNA feedback loops.

Two different 22-nucleotide miRNAs or families, *miR482/miR2118* and *miR2109*, were previously reported to target a large number of *NB-LRR* disease resistance genes in tomato (*Solanum lycopersicum*) and *Medicago truncatula* (Zhai et al., 2011; Shivaprasad et al., 2012). Most *M. truncatula* *NB-LRR* transcripts produce phasiRNAs, triggering additional layers of *NB-LRR* transcript degradation (Zhai et al., 2011). In *F. vesca*, four different variants of *fve-miR482* (*miR482a*, *miR482b*, *miR482c*, and *miR482d*) were

identified; together, they targeted 50 *F. vesca* genes, the majority of which code for *NB-LRRs* (Supplemental Table S4). However, 18 were novel targets unrelated to *NB-LRRs* (Fig. 2, A and C; Supplemental Table S4). Similarly, *miR2109* targeted 40 different *NB-LRR* genes. Our results indicate that even conserved miRNAs may rapidly acquire new family- or species-specific targets.

Two Clusters of Novel *fve-miRNAs* Targeting *FBX* Genes

Previous work hypothesized that most newly formed young miRNAs were neutrally evolving and evolutionarily transient loci. These new loci may not harbor any function but could serve as the source of novel regulatory variations, which could be captured into existing networks (Cuperus et al., 2011; Nozawa et al., 2012). Therefore, young miRNAs may not necessarily have targets or may have targets that are weakly or imprecisely cleaved. However, through analysis of the PARE data, we identified 112 target genes for 25 of the 30 novel miRNAs (Supplemental Table S5). This ratio (25 of 30) is, surprisingly, similar to that of conserved miRNAs with identifiable targets (27 of 31). In contrast, fewer target genes were identified for each of the novel miRNAs, and there was relatively low cleavage tag abundance for these targets (Supplemental Table S5). In addition, unlike conserved miRNAs that often targeted multiple members of a gene family, most novel miRNAs targeted genes from different gene families. Similar observations were made in other plant species, reflecting the properties of young miRNAs (Pantaleo et al., 2010; Xia et al., 2012; Zhu et al., 2012).

One of the most striking observations from the novel miRNA-target analyses was the unusually large number of *FBX* target genes for *miR11288a*, which targeted 11 *FBX* genes with high cleavage signals (category 0-2 in Supplemental Table S5). Further computational target prediction revealed up to 27 *FBX* genes with an *fve-miR11288a* target site (Supplemental Table S6), suggesting the existence of a previously uncharacterized yet extensive miRNA-*FBX* pathway in *F. vesca*.

Table 1. Novel miRNAs identified in *F. vesca*

Novel miRNA Name	Mature Sequence	Length	Strand	Scaffold	Position
fve-miR11286	UUGGAGAGAGAGUAGACAAUG	21	+	scf0513102	292,158
fve-miR11287	UCAGGGAUUGUUUCAUAGACC	21	-	scf0512986	31,505
fve-miR11288a(miRFBX1)	UGGGAUUUGGCGAAUUGUGGU	21	+	scf0513142	48,824
fve-miR11288c2*(miRFBX4*)	UUCGUUCGGAUUCCAAUUCAAA	22	+	scf0513142	50,334
fve-miR11289	UCGUUCGACAUCAAAGGGCACC	21	-	scf0513044	1,313,006
fve-miR11290	UGCUUCAAGUCUGGCCAAUACU	22	-	scf0513192	861,001
fve-miR11291	UUGCGGUCUUGUCUCUUCCAAU	22	-	scf0513113	286,610
fve-miR11292	UUGUAGUUCAGUCUCCGCC	21	-	scf0513157	951,427
fve-miR11293	UUCUUCUCAGGAACCUCCACC	22	+	scf0513111	73,845
fve-miR11294	UUCACCUUGACCAUAAACUGACC	22	-	scf0513110	746,561
fve-miR11295	CUCAUUCAAUUUCGGUUAUUCAG	22	-	scf0513024	360,589
fve-miR11296	UUUUUGAUGGCUUGAAUCCAGU	22	-	scf0513153	13,050
fve-miR11297	CAGACAAGAUCAUCUCGCCU	21	+	scf0513124	126,841
fve-miR11298	UUGAGGGGCUUAAACGAUUACC	21	-	scf0512969	462,110
fve-miR11299	CAAAUAGGGUUGGCUUGAUACU	21	-	scf0513061	229,454
fve-miR11300	CAACAUCACUGUUCUCUUCU	21	-	scf0513136	542,507
fve-miR11301	UCAGAGUUGUAAUUAUUAUUGAU	21	+	scf0513131	1,429,968
fve-miR11302	AGGACCGCCAUACGUUUUGG	21	+	scf0513002	843,149
fve-miR11303	UCAACAUCACUGCAGCUGUA	21	+	scf0513134	1,902,196
fve-miR11304	UAGUGUUUCAGCUUGACAAG	20	+	scf0513172	207,257
fve-miR11305	UUUUGGUCCGAAUCCGAGCUCC	22	-	scf0513080	37,252
fve-miR11306	CUACCGAAGAACUUUGCAAAAG	22	+	scf0512935	232,516
fve-miR11307	UAAGCGACGGACUCCAAUCGC	21	+	scf0513152	2,200,024
fve-miR11308	UAAGUUAGGAUUCUAGUUACC	21	-	scf0513177	3,815,717
fve-miR11309	UUUGUUUGGCAUGCAGUUGGC	21	+	scf0513081	504,742
fve-miR11310	AGGGAUCACAUCUCCAGUGCU	21	-	scf0513138	426,264
fve-miR11311	UGAGAAUGAUGUGGCAUCAGC	22	+	scf0513105	1,393,548
fve-miR11312	UGGAGCUGUUGGGGAGAGUUA	21	+	scf0513098	2,538,938
fve-miR11313	GAAAAGAAUGGACUCUCCGGGG	22	+	scf0513158	5,649,065
fve-miR11314	AGAGUUGUGGAUGCUAUGAAU	21	+	scf0513111	544,059
fve-miR11288b(miRFBX2)	UGGGAUUUGGCGAAUUUUGGU	21	+	scf0513142	49,251
fve-miR11288c1(miRFBX3)	UCCAUCGUUUCGACACGCAGG	21	+	scf0513142	50,416
fve-miR11288c2(miRFBX4)	UGAAUUGGGAUUUGGCUCAAUU	21	+	scf0513142	50,374
fve-miR11288d1(miRFBX5)	UCCAUCGUUUGGAGACACAGG	21	+	scf0513142	52,546
fve-miR11288d2(miRFBX6)	UGAAUUGGGAUUUGGCGAAUU	21	+	scf0513142	52,504
fve-miR11288e1(miRFBX7)	UGAUGGGUAGUCUGGAGAGGAU	22	+	scf0513142	55,908
fve-miR11288e2(miRFBX8)	UGAAGUGGGAUUUGGCGAAUU	21	+	scf0513142	55,825
fve-miR11315(miRFBX9)	CUAGUCAUUGGUCAUAGCAUC	21	-	scf0513106	456,307

Interestingly, manual examination of the *fve-MIR11288a* locus at Linkage Group 3 (LG3) revealed seven potential miRNAs located within 8 kb downstream of *fve-MIR11288a* (scf0513142:48000-56000 or LG3:25351470-25359470; Fig. 3A). These seven potential miRNAs reside within four good stem-loop structures and could generate miRNAs precisely (Fig. 3A). Therefore, these seven miRNAs are considered as novel miRNAs. Because of their sequence similarity to each other (Fig. 3A; Supplemental Fig. S2), miRBase named them as fve-miR11288a, fve-miR11288b, fve-miR11288c1, fve-miR11288c2, fve-miR11288d1, fve-miR11288d2, fve-miR11288e1, and fve-miR11288e2 (Fig. 3A; Table I; Supplemental Table S3). However, these eight related miRNAs are more easily remembered as miRFBX1, miRFBX2, miRFBX3, miRFBX4, miRFBX5, miRFBX6, miRFBX7, and miRFBX8 (Fig. 3A) because of their targeting at a large number of FBX genes (see below). Within this 8-kb region, the star sequence of miRFBX4 was annotated as a novel miRNA in earlier computational identification (Fig. 3A; Supplemental Fig. S3).

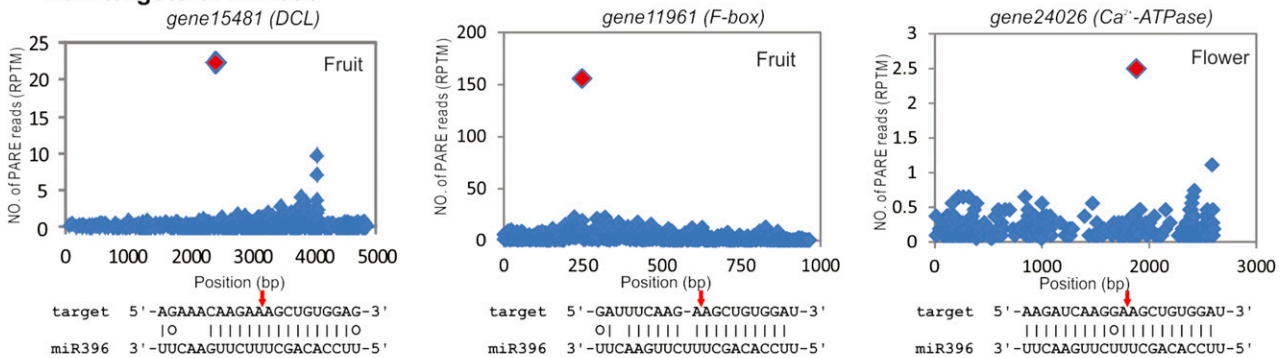
This cluster of eight miRNAs exhibited additional features. First, while fve-miRFBX1 and fve-miRFBX2 are each derived from an independent stem-loop precursor, the other three pairs, fve-miRFBX3 and fve-miRFBX4, fve-miRFBX5 and fve-miRFBX6, and fve-miRFBX7 and fve-miRFBX8, are derived from three stem-loop precursors. The derivation of multiple miRNAs from the same stem-loop precursor was observed previously in other plants (Zhang et al., 2010). Second, seven of the eight matured fve-miRNAs (except fve-miRFBX7) show sequence similarities, with a minimum identity of 15 nucleotides, to one another (Fig. 3A). Strikingly, all eight fve-miRNAs were predicted to collectively target 94 FBX genes (Supplemental Table S6). Cleavage products of 51 of the 94 FBX genes were confirmed by PARE (Fig. 3B; Supplemental Table S6). Many of these FBX genes could be cleaved by two or more miRNAs (Fig. 3B; Supplemental Table S6). For simplicity, we refer to this cluster of eight fve-miRNAs as the LG3 miRFBX cluster.

To identify FBX-targeting miRNAs elsewhere in the genome, we conducted target gene prediction for all

A A selected subset of new targets for conserved miRNAs

miRNA	Target Gene	Align score	Cleavage site	Flower		Fruit		Vegetative		Annotation
				RN_reads	Category	RN_reads	Category	RN_reads	Category	
Conserved										
fve-miR159	gene27169	4	733	9.63	0	3.41	0	2.43	0	Unknown protein
fve-miR319	gene18186	4	397	0.93	2	0.55	2	1.04	2	chloroplast-targeted copper chaperone
fve-miR390	gene05023	4.5	804	29.35	0	7.74	2	25.83	0	Unknown protein
fve-miR390	gene18596	4.5	4128	5.09	2	1.10	2	6.25	2	Endoribonuclease Dicer homolog
fve-miR393	gene16871	4.5	141	1.11	2	1.10	2	0.58	2	Transcription factor bHLH78
fve-miR396	gene03763	4.5	1650	2.78	2	2.74	0	1.97	2	Agemat and bromo-adjacent homology (BAH) domain-containing protein-like protein
fve-miR396	gene11961	4.5	247	45.00	0	155.99	0	30.69	0	F-box protein FBW2
fve-miR396	gene15481	4	2417	6.30	0	22.29	0	7.64	0	Endoribonuclease Dicer homolog 3a
fve-miR396	gene23939	4	303	1.85	0	0.49	2	3.36	2	50S ribosomal protein L6, chloroplastic
fve-miR396	gene24026	3	1881	2.50	0	1.10	1	1.16	2	Probable cadmium/zinc-transporting ATPase HMA1
fve-miR396	gene25882	3	1011	1.20	0	1.89	2	0.12	4	Sulfite oxidase
Less conserved										
fve-miR2111	gene11514	2.5	218	1.67	0	2.44	0	0.12	4	Phosphate transporter
fve-miR2111	gene25889	2	197	14.54	0	28.51	0	1.85	0	F-box/kelch-repeat protein
fve-miR2111	gene30412	4.5	3400	20.83	2	21.68	2	23.16	2	C-terminal domain phosphatase-like 1
fve-miR482	gene07624	3	151	20.09	0	35.63	0	23.05	0	putative ribonuclease H protein
fve-miR482	gene23942	2	463	13.89	0	11.21	0	4.40	0	Protein TRANSPARENT TESTA 12
fve-miR482	gene31573	2	74	170.01	0	164.58	0	99.95	0	Cyclic phosphodiesterase
fve-miR482	gene31576	2	737	148.06	0	42.45	0	57.44	0	Unknown protein
fve-miR530	gene32544	2	108	3.98	0	1.34	2	3.01	2	zinc knuckle (CCHC-type) family protein

B New targets of miR396



C New targets of miR482

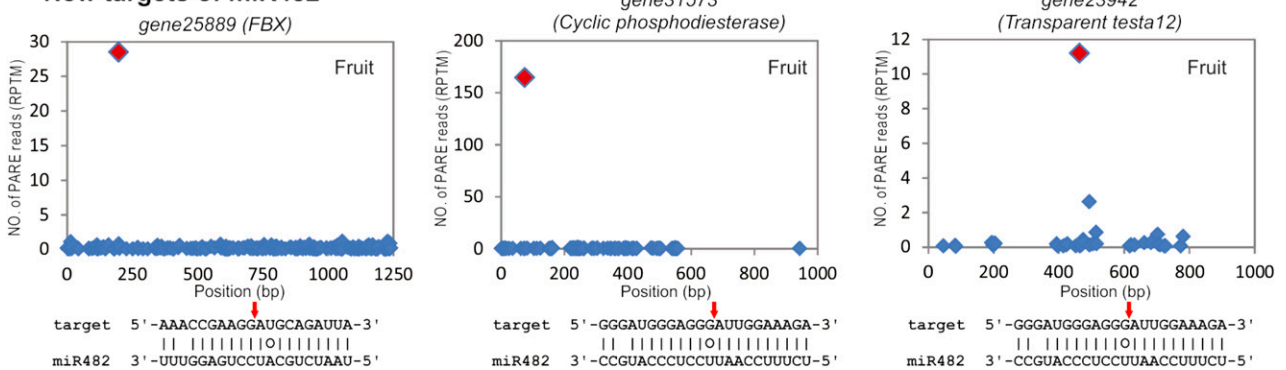


Figure 2. Novel target genes of conserved and less conserved miRNAs. A, List of selected novel target genes of conserved and less conserved miRNAs based on PARE data from flower, fruit, and vegetative tissues. Repeat normalized reads (RN_reads) are derived by dividing PARE read abundance by the number of loci in the genome to which the reads align. Alignment score (between 0 and 4.5) is calculated based on how well a miRNA pairs with its target sequence; a lower score indicates a better alignment between the miRNA and its target. The category score (between 0 and 4) considers the ratio of the PARE tag reads mapped to the

unique small RNAs with good stem-loop precursor structures (derived from earlier analyses) and found another novel miRNA, *fve-miR11315* (*miRFBX9*). This miRNA is encoded by a cluster of four stem-loop structures (Fig. 3C) located within an approximately 500-bp region (scf0513106: 456,208–456,714; or LG5: 14,533,804–14,534,310) and bears no sequence similarity to miRNAs at the LG3 *miRFBX* cluster (Fig. 3, A and C). This miRNA (*fve-miR11315* or *miRFBX9*) was predicted to target 12 *FBX* genes, and the PARE data confirmed the cleavage of seven of the 12 *FBX* genes (Supplemental Table S6). We refer to this second cluster as the LG5 *miRFBX* cluster. Collectively, these nine miRNAs (*miRFBX1*–*miRFBX9*) at the LG3 and LG5 *miRFBX* clusters are termed *fve-miRFBXs*.

***fve-miRFBX7* Cleaves *FBX* Transcripts and Initiates phasiRNA Production**

Among the nine *fve-miRFBXs*, only *fve-miRFBX7* is 22 nucleotides in length and possesses an initial uridine. In plants, 22-nucleotide uridine-initiated miRNAs are capable of triggering phasiRNAs from cleaved mRNAs of *PHAS* loci (Chen et al., 2010; Cuperus et al., 2010). We assessed whether *fve-miRFBX7* can initiate phasiRNA production. Using our *PHAS* profiling algorithm described previously (Xia et al., 2013) and the nine small RNA libraries, we identified a total of 24 *PHAS FBX* genes for which phased peaks of phasiRNAs could be detected (Supplemental Table S7). However, only six of these 24 *PHAS FBX* genes are directly cleaved by *fve-miRFBX7*, as indicated by solid alignment scores between *fve-miRFBX7* and the respective *FBX* genes (Supplemental Table S7).

In addition, *fve-miRFBX7* appears to possess common *FBX* targets with *miRFBX9*, *gene31348*, *gene31347*, *gene31349*, *gene13786*, and *gene27192* (Supplemental Table S6), indicating that *fve-miRFBX9* and *fve-miRFBX7* may participate in a two-hit cleavage to delimit the boundary of a region that gives rise to phasiRNAs. For example, *fve-miRFBX9* cleaves *FBX gene31348* at a site approximately 220 bp downstream of the *fve-miRFBX7* cleavage site (Fig. 3D). Indeed, abundant and progressively cleaved phasiRNAs were produced from the region immediately after the *fve-miRFBX7* cleavage site (Fig. 3D).

***miRFBXs* Mostly Target the Conserved F-Box Domain**

Plant *FBX* genes constitute a large family of genes (approximately 800 in *F. vesca* and approximately 690 in *Arabidopsis*) and can be classified into several distinct

subfamilies based on domain structures (Kuroda et al., 2002; Xu et al., 2009). To investigate which specific *FBX* gene subfamilies are regulated by the clustered *miRFBXs* and deduce their potential functions, we performed phylogenetic analysis using all the *FBX* genes from *Arabidopsis* (691 genes; Xu et al., 2009) and the set of 96 *F. vesca FBX* genes targeted by the clustered *miRFBXs* (Supplemental Table S6). As shown in Figure 4A, the *Arabidopsis FBX* genes are grouped into several distinct subfamilies, including FBA1, FBA3, LRR-FBD, Kelch repeats, and Domain Unknown Function295. Almost all the *F. vesca FBX* genes (except five) targeted by the *fve-miRFBXs* favorably cluster with the FBA1 and FBA3 subfamilies (Fig. 4A). The subtree for the *F. vesca FBX* genes could be further divided into three groups. Group I is targeted mainly by the LG3 *miRFBX* cluster (except *miRFBX7*). Group II is targeted by *fve-miRFBX7* and *fve-miRFBX9* and encompasses many *PHAS FBX* genes. Group III is preferentially regulated by *miRFBX7* (Fig. 4B).

Next, we examined the distribution and conservation pattern of *miRFBX* target sites within the *FBX* coding sequence. As the *fve-miRFBX*-targeted *FBX* genes are more similar to FBA1/3 subfamilies, we modeled the structure of an *FBX* gene with three typical domains, the F-box, FBA1/3, and the F-box_assoc_1 domain (Fig. 4C). The target sites of *fve-miRFBX3*, *fve-miRFBX5*, and *fve-miRFBX7* are located within the encoded F-box domain (Fig. 4C). In contrast, the target sites of *fve-miRFBX1*, *fve-miRFBX2*, *fve-miRFBX4*, *fve-miRFBX6*, and *fve-miRFBX8* are located at the C-terminal end of the F-box domain. The *fve-miRFBX9* target site is in the F-box_assoc_1 domain in some *FBX* genes (Fig. 4C). Therefore, targeting at the more conserved region of the *FBX* genes endows these *fve-miRFBXs* with the ability to regulate a large number of *FBX* family members simultaneously. However, not every *FBX* gene harbors the target sites of all members of the *miRFBXs*; therefore, each *FBX* gene may only be targeted by a subset of the *miRFBXs* as illustrated in Figure 4B.

The LG3 Cluster of *miRFBXs* Recently Evolved in the *F. vesca* Genome from Gene Duplication

Does this regulatory network of *miRFBX-FBX* exist in other higher plants? Similar miRNA pathways have not been reported even in related species such as apple and peach (Xia et al., 2012; Zhu et al., 2012). Therefore, we searched and then identified syntenic regions of the LG3 *miRFBX* locus in many plant species (Supplemental Fig. S4). Strikingly, this region is significantly expanded in *F. vesca* (Fig. 5A) when compared with even the closely

Figure 2. (Continued.)

corresponding miRNA-mediated cleavage site to the total number of PARE tags in a given target mRNA. The lower the category score, the higher the confidence of the cleaved targets. B, T-plots for selected target genes of *miR396*. The y axis indicates the abundance of cleavage PARE reads (reads per 10 million [RPTM]). The x axis indicates the coding sequence nucleotide position (bp), with 1 marking the first nucleotide of the start codon. The red diamond denotes the cleavage site by the corresponding miRNA in the fruit or flower PARE library. The base pairing between miRNA and the target transcript is indicated beneath each plot, with the exact cleavage site indicated by a red arrow. C, T-plots for selected target genes of *miR482*.

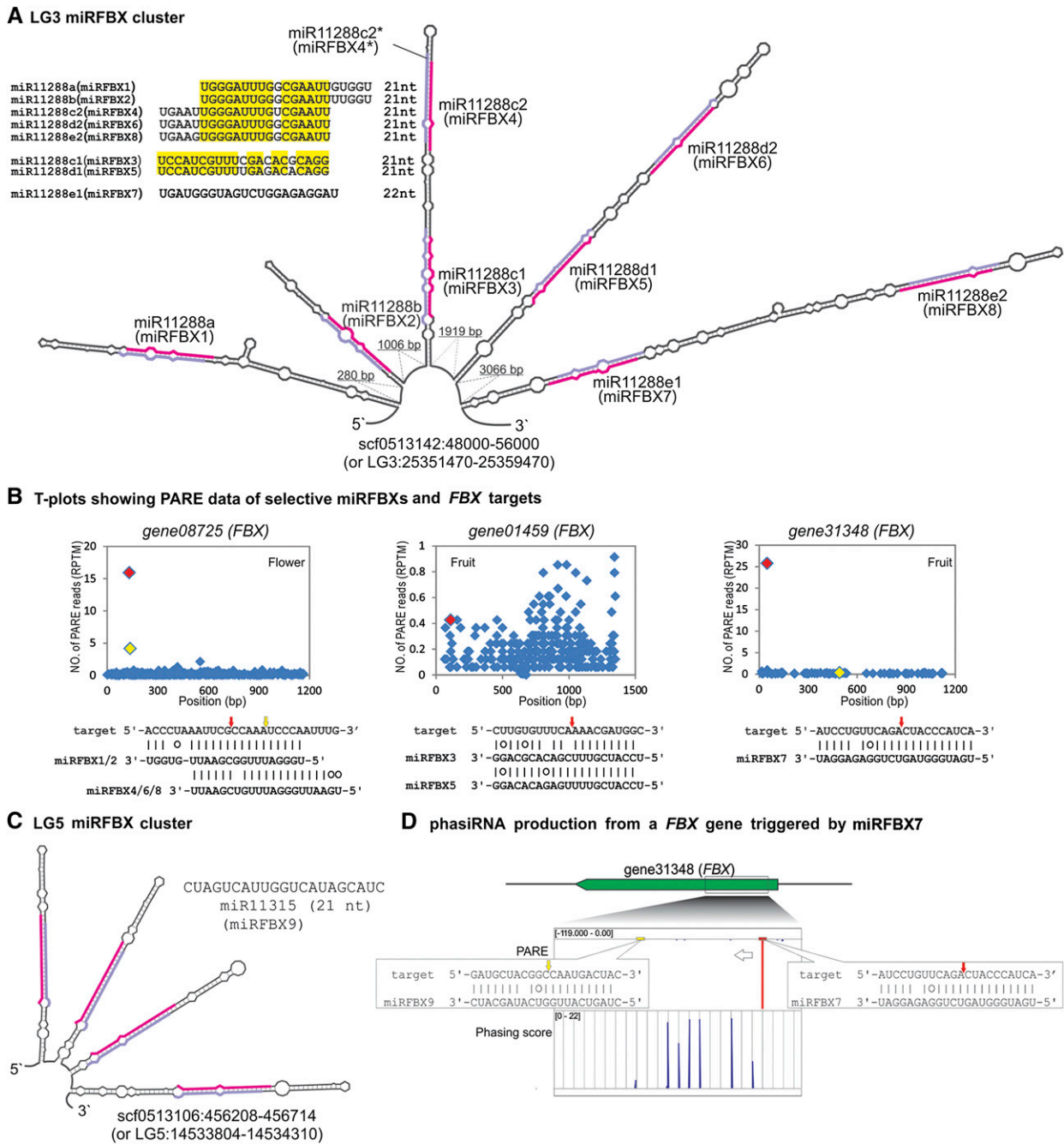


Figure 3. Two new five-miRNA clusters in the *F. vesca* genome. A, Computationally predicted multiple stem-loop structure from an 8-kb region in scf0513142 located in LG3. Eight pairs of miRNA and miRNA* are marked in pink and light purple, respectively. Their names as given by miRBase (miR11288a, miR11288b, miR11288c1, miR11288c2, miR11288d1, miR11288d2, miR11288e1, and miR11288e2) are indicated; an easier to remember name, miRFBX1 to miRFBX8, is also given for each miRNA in the cluster. Seven of these eight miRNAs are related in sequence, as shown in the top left corner; conserved nucleotides are highlighted in yellow. B, T-plots for selected five-miRFBX genes and their targeted *FBX* transcripts. The red and yellow diamonds denote the cleavage sites of miRFBXs. Yellow refers to a second cleavage event by a second miRFBX. C, Another stem-loop structure in LG5 codes for a cluster of four identical miRNAs (miR11315/miRFBX9). The miRFBX9 sequence is indicated above the structure. D, Primary and secondary cleavages of *gene31348 (FBX)* and phasiRNA production. *gene31348* shown on the top is cleaved by miRFBX7 (red arrow) and miRFBX9 (yellow arrow). In the PARE track, cleavage tag abundance is shown as a red downward-pointing peak; the negative value indicates antisense transcript (left-pointing arrow); numbers in brackets indicate the range of PARE read abundance. In the phasing score track, phasiRNA peaks are evenly distributed and upward pointing. The phasing score inside the bracket was calculated from the flower small RNA library.

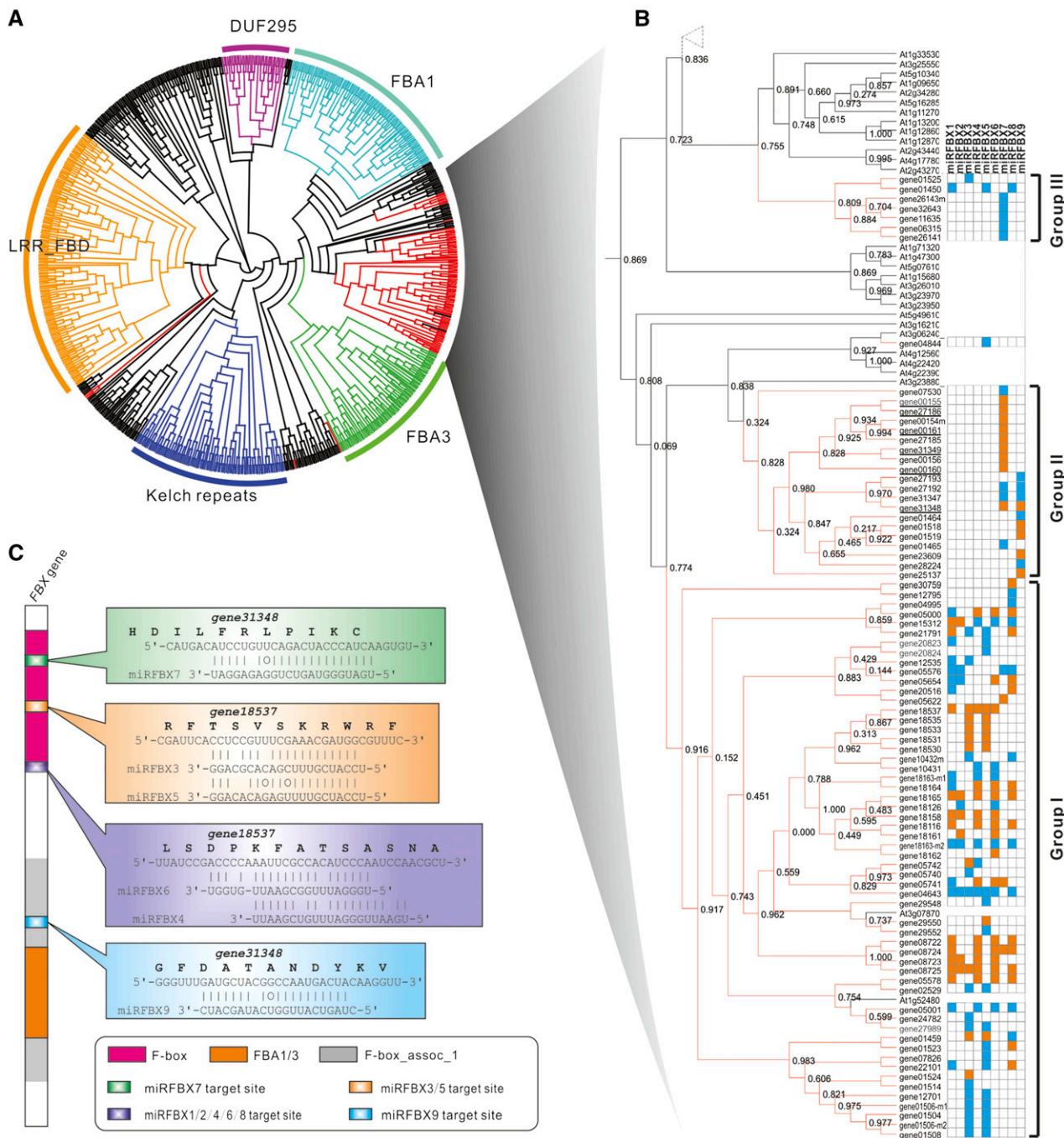


Figure 4. Analysis of *FBX* genes targeted by the nine five-miRFBXs. A, Phylogenetic tree incorporating all *Arabidopsis* *FBX* genes and miRFBX-targeted *FBX* genes in *F. vesca*. Five major classes are marked with distinct colors and indicated by group names (minor groups are in black). B, Enlarged *F. vesca* *FBX* genes targeted by the nine miRFBXs shown as orange lines. These genes are divided into three groups (I, II, and III) according to positions on the tree and the targeting miRNAs. Blue-filled grids indicate *FBX* genes bearing a predicted target site (align score ≤ 5) for a given miRNA; orange-filled grids indicate the detection of cleavage products by PARE. *PHAS* *FBX* genes directly targeted by five-miRFBX7 are underlined. C, Diagram of the *FBX* protein with typical domains characteristic of the FBA1/3 subfamilies. The positions of various miRFBX target sites are indicated, and examples of miRNA pairing to target *FBX* sequences are included on the right. Not all *FBX* genes harbor all the miRNA target sites.

related Rosaceae species apple (in the genome between MDP0000286003 and MDP000349339) and peach (between ppa026038m and ppa010681m), suggesting that

the miRFBX cluster may have recently evolved in *F. vesca* from a common ancestor of the Rosaceae. Small RNA mapping data at the LG3 miRFBX region revealed

abundant miRNAs produced from the five putative *MIRNA* precursor loci, all in the same transcriptional orientation (Fig. 5B). Between *MIRFBX5/6* and *MIRFBX7/8* is a region with repetitive sequences that gave rise to large quantities of 24-nucleotide small RNAs in both directions (Fig. 5B).

To investigate from where these LG3 *MIRFBX* genes could originate, BLASTX was conducted with the LG3 *MIRFBX* DNA sequence (approximately 8 kb) as a query against public protein collections. The *MIRFBX3/4* and *MIRFBX5/6* regions showed significant similarity to protein-coding genes; the reverse complement strands of *MIRFBX3/4* and *MIRFBX5/6* loci can be translated into appreciable peptides of 45 or 46 amino acids, respectively (Fig. 5C). When BLASTP was conducted against all *F. vesca* protein-coding genes, the top hits for either of these two peptides were FBX proteins (Fig. 5D), and these FBX homologs showed significantly lower *e*-values than the non-FBX hits (Student's *t* test, $2.2e-16$; Fig. 5D). Our analysis strongly suggests that *MIRFBX3/4* and *MIRFBX5/6* were derived from *FBX* genes. Considering the high level of sequence similarity among the five stem-loop sequences within the LG3 miRFBX cluster (Supplemental Fig. S2), it is likely that these five stem loops at the LG3 evolved from duplications of a common precursor (see "Discussion").

Expression of Precursor Transcripts of miRFBXs and Their Target *FBX* Genes in Various Tissues

Since our initial small RNA libraries were made for the sole purpose of discovering small RNAs in *F. vesca*, the expression of each miRNA could not be quantitatively assessed in the absence of replicates for each of the nine small RNA libraries. However, we previously generated RNA-seq data for 37 different floral, fruit, and vegetative tissues in *F. vesca* with two biological replicates for each tissue type (Kang et al., 2013; Hollender et al., 2014). The levels of miRNA precursor transcripts could be examined. Figure 6A illustrates the RNA-seq data in the LG3 *fve*-miRFBX cluster region, which corresponds to at least four different precursor transcripts (transcripts 1–4). Of particular interest to us is transcript 4, which is the precursor for both *fve*-miRFBX7 and miRFBX8. Transcript 4 (Fig. 6B, row 4) is highly abundant and specific to both the pith and the cortex of the receptacle fruit at all five fruit developmental stages (pith 1–5 and cortex 1–5). Consistent with the precursor RNA expression, the small RNA read abundance for miRFBX7 and miRFBX8 also showed highest expression in the receptacle among the nine tissues sampled (Supplemental Fig. S5). *fve*-miRFBX7/8 may thus be involved in regulating receptacle fruit development. The RNA-seq read abundance for the precursor transcript of the *MIRFBX9* cluster (transcript 5) was also examined, which showed extremely low levels in all floral and fruit tissues (Fig. 6B). This is consistent with the low expression of mature *fve*-miRFBX9 in flower and fruit tissues (Supplemental Fig. S5).

To test if there is an inverse relationship in transcript abundance between miRFBX7 and its *FBX* targets, we investigated the transcript abundance of the 24 *FBX* genes, either predicted or proven by PARE analysis to be targets of miRFBX7 (Supplemental Table S6), using the same comprehensive flower and fruit RNA-seq data sets (Kang et al., 2013; Hollender et al., 2014). Hierarchical clustering of Z-score normalized mapped reads across 37 tissues was shown for each of the 24 *FBX* genes (Fig. 6C). Interestingly, 10 *FBX* genes clustered together (at the bottom of the heat map) and showed dramatic reduction of transcript abundance in the pith and cortex tissues (Fig. 6C). This is consistent with possible degradation of these 10 *FBX* transcripts by miRFBX7, whose precursor specifically accumulates in the pith and cortex (transcript 4 in Fig. 6B) and whose mature miRNAs were most abundant in the receptacle tissue (Supplemental Fig. S5). Among these 10 *FBX* genes, five are known to produce phasiRNAs triggered by miRFBX7 (Fig. 6C; Supplemental Table S7). Hence, this analysis identified a miRFBX7-*FBX*-phasiRNA network operating in the receptacle fruit of *F. vesca*.

Conserved tasiRNA Pathways in *F. vesca*

Three major conserved tasiRNA pathways have been characterized in plants; their biogenesis is initiated by three distinct miRNAs: miR390, miR828, and the miR7122 superfamily (Fei et al., 2013; Xia et al., 2013). All three tasiRNA pathways are also found in *F. vesca* (Fig. 7). The miR390-*TAS3* pathway is one of the most well-characterized pathways in plants and was identified in mosses as well as higher plants (Axtell et al., 2007; Howell et al., 2007). In *Arabidopsis*, miR390 cleaves *TAS3* transcripts using a two-hit mechanism, generating tasiARFs that guide the degradation of transcripts encoding AUXIN RESPONSE FACTOR2 (ARF2), ARF3, and ARF4. tasiARFs regulate many important developmental processes ranging from leaf patterning and lateral root initiation to developmental timing (Adenot et al., 2006; Fahlgren et al., 2006; Garcia et al., 2006; Marin et al., 2010). *F. vesca* has two *TAS3* genes that we named *TAS3L* (L for long) and *TAS3S* (S for short; Fig. 7A). *fve*-*TAS3L* is processed similarly to *Arabidopsis* *TAS3*, resulting in two *fve*-tasiARFs. In contrast, *fve*-*TAS3S* is cleaved at both 3' and 5' sites (Fig. 7A). The phasiRNAs, including one tasiARF, are produced downstream of both cleavage sites. The demonstrated cleavage at both 3' and 5' sites of *fve*-*TAS3S* by the PARE data argues against an absolute requirement for a noncleavable 5' site in the two-hit mode of tasiARF biogenesis (Axtell et al., 2006; Allen and Howell, 2010).

miR828 is conserved in eudicots (Cuperus et al., 2011); although it was not identified in our list of known or conserved miRNAs due to its extremely low expression level, its cleavage of *fve*-*TAS4* and tasiRNA production were confirmed by the PARE data and phasing analyses, respectively (Fig. 7B). In addition to

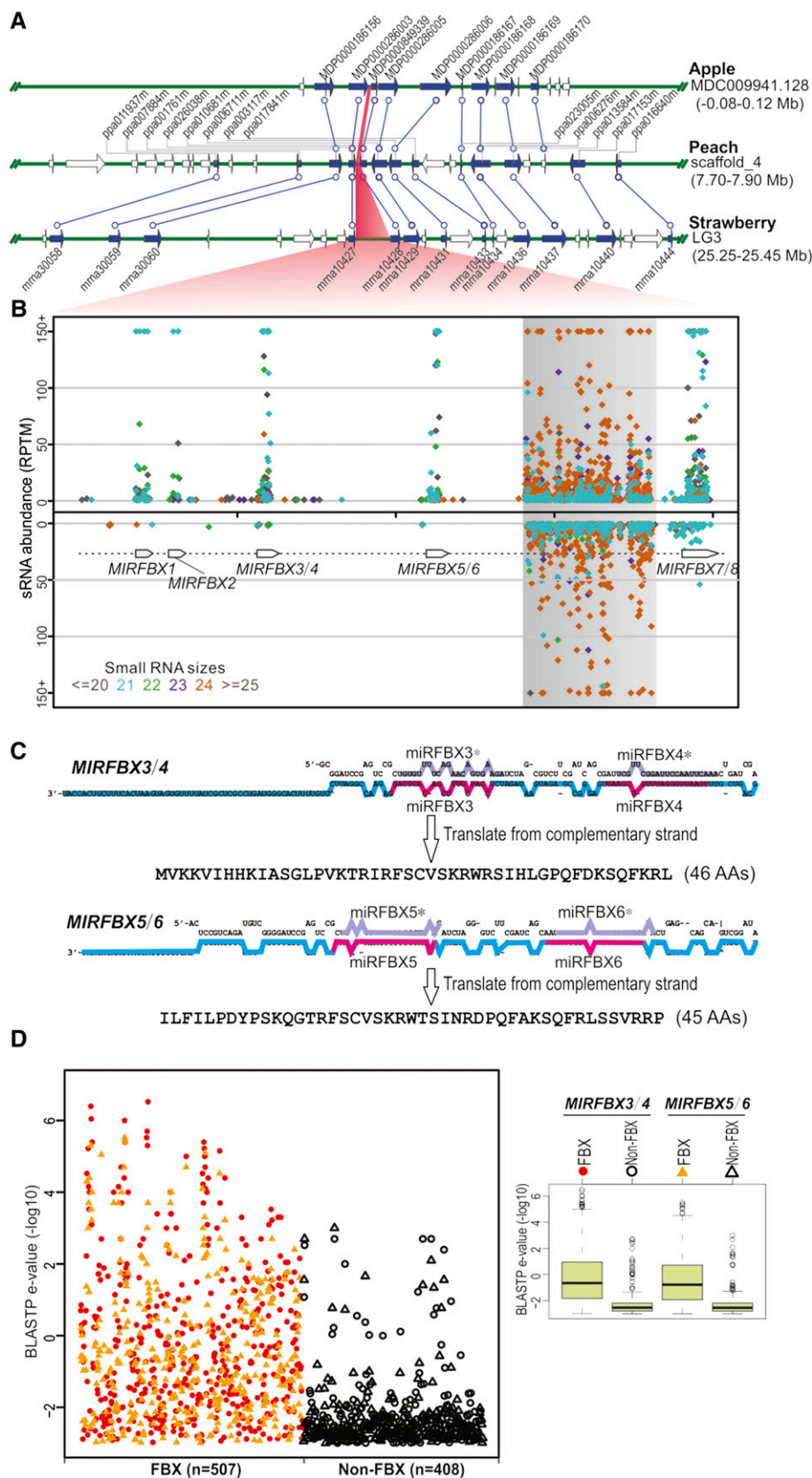


Figure 5. The LG3 miRFBX cluster likely evolved recently from *FBX* genes. **A**, Alignment of syntenic regions among apple, peach, and *F. vesca*. Genes are shown as arrows, with syntenic genes as filled blue arrows. Each syntenic gene pair is connected by a blue line. The LG3 miRFBX cluster region in strawberry and the corresponding syntenic regions in apple and peach are shaded in red. **B**, Small RNA distribution within the LG3 miRFBX region in *F. vesca*. Small RNAs of different length are denoted with different color dots. A region of repetitive DNA that yielded abundant 24-nucleotide small RNAs is shaded in gray. **C**, Stem-loop structures for *MIRFBX3/4* and *MIRFBX5/6* precursor RNAs. miRNA and miRNA* are highlighted in pink and light purple, respectively. The reverse-complementary sequence of the longer arm of the stem-loop structure (the cyan- and pink-colored region) codes for a peptide of 46 (for *MIRFBX3/4*) or 45 (for *MIRFBX5/6*) amino acids (AAs). **D**, Summary of BLASTP results for the two short peptides in **C**. BLASTP search was conducted against *F. vesca* proteins with an e-value cutoff of 1,000. All the hits (915 proteins) were classified into two groups: FBX proteins (507) and non-FBX proteins (408). Many more FBX proteins were obtained with significantly lower BLASTP e-values (show in $-\log_{10}$ scale) than the non-FBX proteins. The e-value differences between FBX and non-FBX proteins for the two peptides are shown in the box plot on the right.

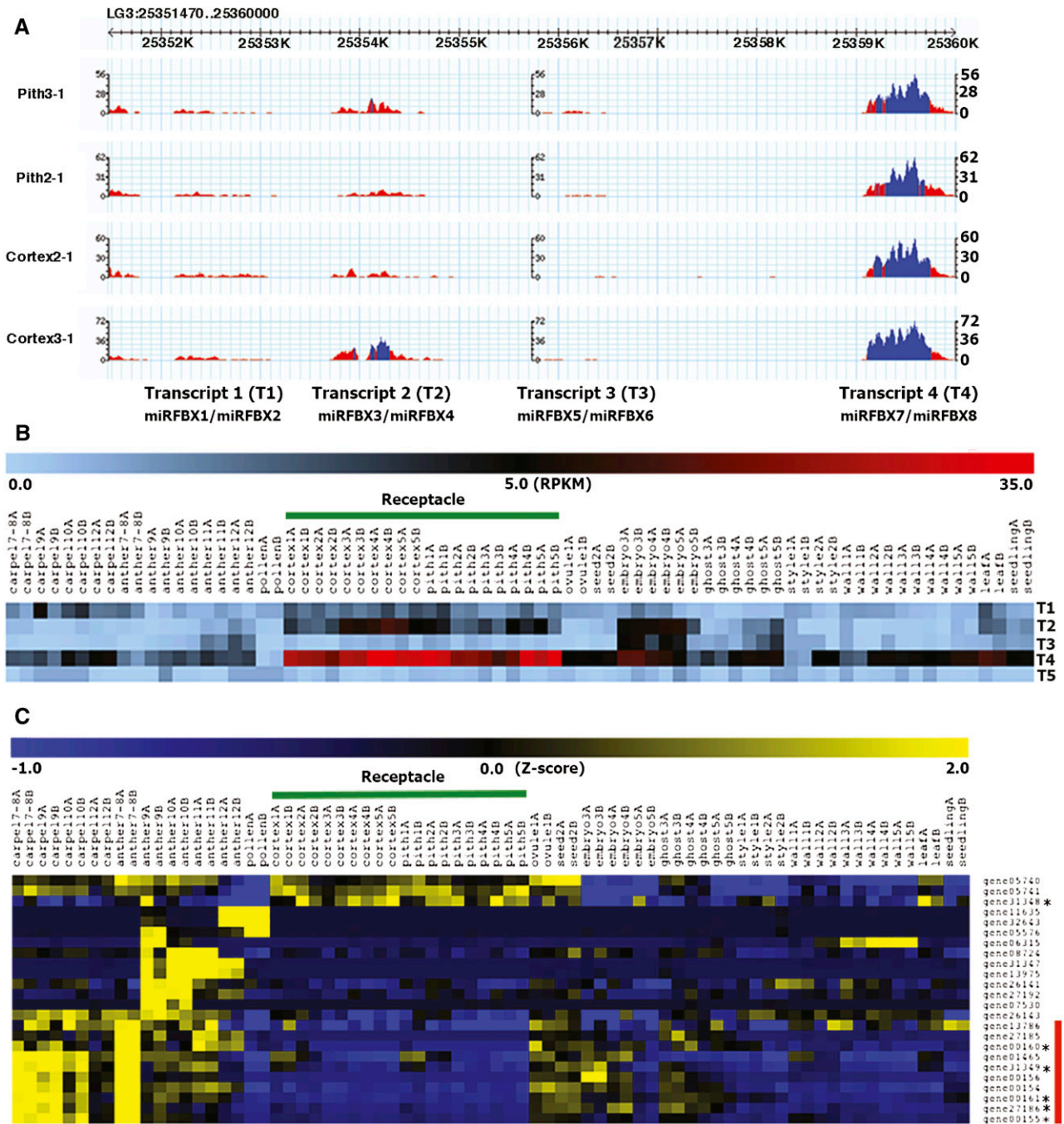


Figure 6. miRFBX7/8 precursor transcript is specifically expressed in the receptacle fruit. A, RNA-seq read abundance displayed via GBrowse at the Genome Database for Rosaceae (GDR) on the LG3 miRFBX cluster region (LG3: 25351470–25359470). There appear to be four distinct transcription units: miRFBX1/2 (T1, transcript 1), miRFBX3/4 (T2, transcript 2), miRFBX5/6 (T3, transcript 3), and miRFBX7/8 (T4, transcript 4). Blue color indicates greater than 20× read coverage, while red color is 20× or less coverage. B, Heat map showing read abundance (reads per kilobase per million [RPKM]) of the four LG3 precursor transcripts (T1, T2, T3, and T4) shown in A and the LG5 miRFBX9 precursor transcript (labeled as T5). RNA-seq data of 37 different tissues and developmental stages (labeled by number) are shown. The two biological replicates are labeled as A and B. The receptacle (highlighted by the green line) is divided into pith and cortex at fruit developmental stages 1 to 5. C, Hierarchical clustering (Pearson correlation) showing the relative expression level of 24 *FBX* genes targeted by miRFBX7. Mapped reads across the same 37 tissues/stages were normalized via the Normalize Genes/Rows function in MeV4.8. The phasiRNA-producing *FBX* genes (triggered by miRFBX7) are marked by asterisks. The red bar on the right side denotes the cluster showing reduced expression in pith and cortex of the receptacle.

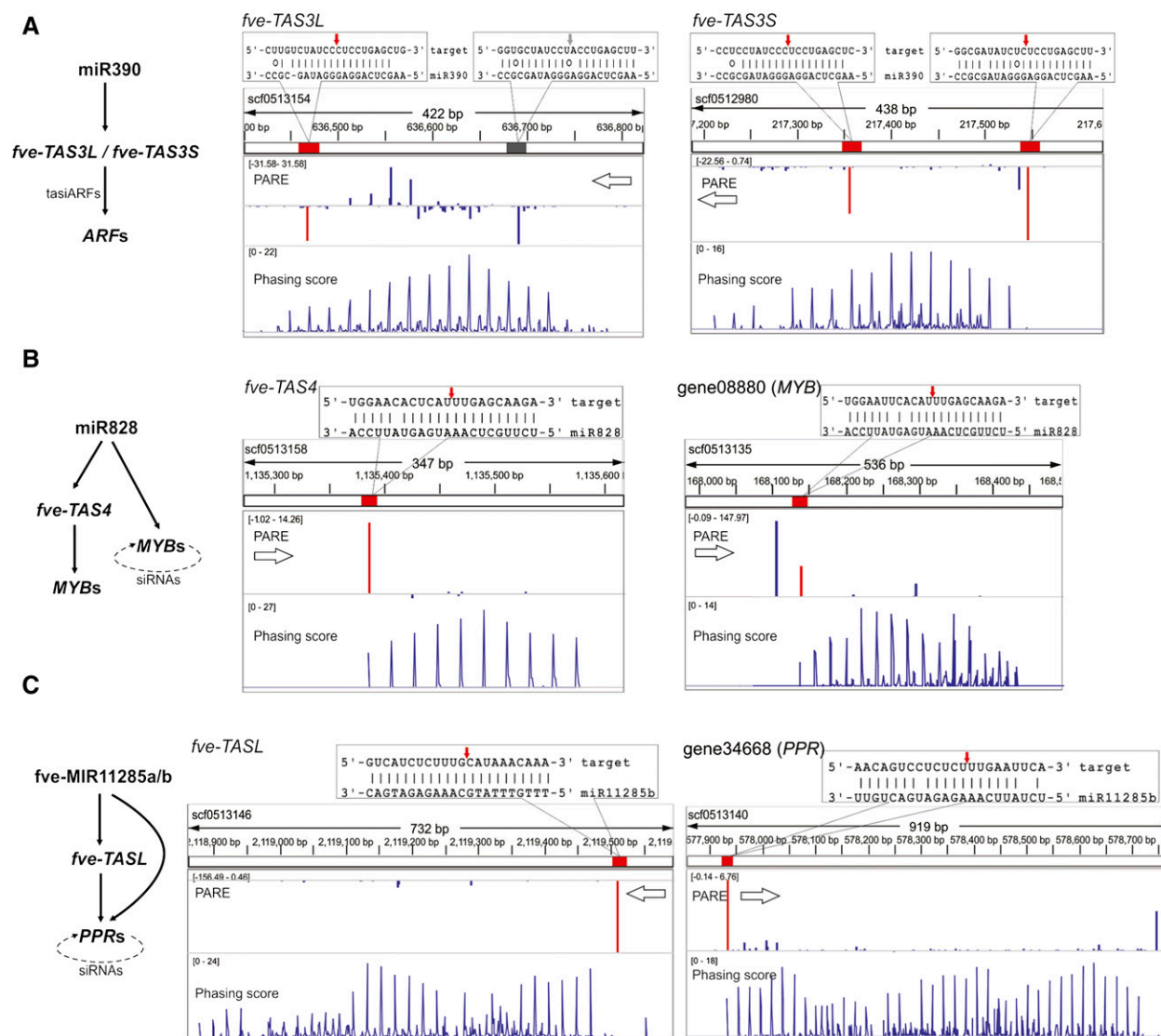


Figure 7. Conserved tasiRNA pathways in *F. vesca*. Diagrams on the left illustrate conserved trans-acting pathways in *F. vesca*. On the right are genomic configurations of *TAS* genes showing three tracks: position of the miRNA target site (red or gray rectangular boxes) within the scaffold, PARE data (middle track), and phasing score (bottom track) viewed in the Integrative Genomics Viewer. Base pairing between the miRNA and corresponding transcript is enlarged above the tracks, with the exact cleavage site marked with a red or gray arrow. The gray box indicates a miR390 target site that is not cleaved. The PARE track and the phasing score track are as described in Figure 3D. The values in brackets in the PARE track and the phasing score track were derived from the flower PARE library and the flower small RNA library, respectively. A higher phasing score indicates a higher confidence of the *PHAS* loci. **A**, *fve-TAS3L* and *fve-TAS3S* are homologous genes located on different scaffolds. Each has two miR390-binding sites, indicated by the red and gray boxes in the top track. Cleavages by miR390 are shown by downward (antisense) or upward (sense) peaks. Red peaks indicate cleavage products by the indicated miRNA. Blue peaks indicate other cleavage products. **B**, *fve-TAS4* and *MYB* genes are both cleaved by miR828 in typical one-hit mode. The dotted arrow in the diagram at left indicates self-cleavage by phasiRNAs. **C**, *fve-TASL* and *PPR* genes are each cleaved once by *fve-miR11285a/b* (previously named as *fve-PPRtri1/2*; Li et al., 2013; Xia et al., 2013). This initiates phasiRNAs revealed by the phasiRNA peaks. The dotted arrow in the diagram at left indicates self-cleavage by phasiRNAs.

targeting *fve-TAS4*, *fve-miR828* directly cleaved several *MYB* genes such as *gene08880* (Fig. 7B; Supplemental Table S8) leading to phasiRNAs, some of which target transcripts from cognate *MYB* genes. Copies of *fve-MIR11285a/b* (previously named as *fve-PPRtri1/2*)

were described to target *PPR* genes directly or via cleavage of an intergenic noncoding locus named as a *TAS-LIKE* gene (*TASL*; Xia et al., 2013). This was confirmed by our PARE data, and the ensuing phasiRNAs were detected in many small RNA libraries (Fig. 7C).

DISCUSSION

F. × ananassa is a hybrid octoploid and poses tremendous challenges in genetic and genomic studies. Past studies of cultivated strawberry, including miRNA identification (Ge et al., 2013; Li et al., 2013; Xu et al., 2013), were mostly focused on the ripening stage fruit. In this work, we focused our analysis on *F. vesca*, a diploid wild strawberry whose genome has been sequenced (Shulaev et al., 2011). Based on stringent criteria (Meyers et al., 2008), we identified 31 known and 38 novel miRNAs from *F. vesca*. Our identification and analyses of miRNAs in *F. vesca* benefited from the sequenced *F. vesca* genome (Shulaev et al., 2011), allowing accurate mapping of sequenced reads, a well-described developmental morphology of reproductive development (Hollender et al., 2012) that allowed accurate tissue and stage identification, and the availability of extensive transcriptome data sets in *F. vesca* (Kang et al., 2013; Hollender et al., 2014), which enabled us to examine the expression patterns of miRNA precursors and miRNA target genes. Importantly, the established transformation protocol and the ease of gene knockout in a diploid background will enable future functional studies.

A Majority of Conserved *fve*-miRNAs Have Evolved Novel Targets

One surprising finding is that 20 of the 31 known miRNAs exhibited not only conserved canonical gene targets but also previously unreported, presumably novel, gene targets. For example, among the 30 target genes of miR396, 11 are conserved targets (Supplemental Table S4, genes highlighted in boldface) but 19 are novel targets (Supplemental Table S4, genes not in boldface). Previous PARE analysis carried out in apple revealed that, among the 19 conserved miRNAs, two miRNAs (miR319 and miR396) were found to have new gene targets (Xia et al., 2012). By comparing our *F. vesca* list with that of apple (Xia et al., 2012), we found that the new gene targets of miR319 and miR396 in apple are distinct from those in *F. vesca*. Therefore, new miRNA-target pairs may form relatively easily and change rapidly in evolutionary time, underscoring the dynamic nature of miRNA-based regulation and its continuous evolution.

phasiRNA Production in Combination with miRNA-Target Pairing at Conserved Protein Domains Can Efficiently and Simultaneously Switch Off Members of Large Gene Families

More significantly, our work identified a novel miRNA-*FBX* network in *F. vesca*. Nine novel miRFBXs were found to cluster at two chromosomal regions: eight at LG3 within an 8-kb region and four at LG5 within a 500-bp region (Fig. 3, A and C). *MIRNA* genes are generally scattered in plant genomes. Only 17% of plant *MIRNA* genes were found to reside within clusters, and usually fewer than three *MIRNA* genes are

clustered together (Nozawa et al., 2012). This is in general much lower than the clustered *MIRNA*s in human and *Drosophila* spp. (approximately 40%; Altuvia et al., 2005; Nozawa et al., 2010). The two miRFBX clusters found in *F. vesca* demonstrated that the relatively large size of the *MIRNA* gene family could arise quickly from tandem duplications, even though this is rather rare in plants.

In legumes, the miR482/2118 superfamily regulates hundreds of *NB-LRR* genes by targeting at the conserved P-loop region and triggering profuse secondary phasiRNA production (Zhai et al., 2011; Shivaprasad et al., 2012). In apple, three conserved miRNAs, miR828, miR858, and miR159, were shown to potentially target up to 81 different *MYB* genes (Xia et al., 2012). Target sites of miR858 and miR828 are within the conserved R3 domain of *MYBs*. In this study, we discovered that nine novel miRFBXs target a total of 94 *FBX* genes and that the target sites of eight miRNAs fall either partially or fully within the region coding for the conserved F-box domain (Fig. 4C). Our work, combined with recent findings mentioned above, reveals that miRNAs targeting at conserved protein domains can be particularly powerful in coordinately regulating large gene families.

Of increasing significance are recent discoveries of 22-nucleotide miRNAs capable of initiating phasiRNAs from protein-coding *PHAS* genes in higher plants. The 22-nucleotide miRNAs were reported in legumes to trigger phasiRNA production from more than 74 loci coding for the *NB-LRR*s (Zhai et al., 2011; Arikrit et al., 2014). The resulting phasiRNAs further coregulate en masse, in trans or cis, *NB-LRR* transcripts. Similarly, 22-nucleotide miR828 reinforces its silencing of the *MYB* family of genes by instigating phasiRNAs from *MYB* transcripts (Xia et al., 2012). The superfamily of miR7122, conserved in eudicots, was also found to induce phasiRNAs from *PPR* genes in a wide variety of plants (Xia et al., 2013). In this study, we found a miRNA-phasiRNA pathway specifically targeting *FBX* genes in *F. vesca*. Although the function of these various phasiRNA networks is unknown, these prior findings together with the work reported here suggest that phasiRNA production, in combination with miRNA-target pairing at conserved protein domains, is a highly efficient and sustainable strategy evolved in higher plants to simultaneously switch off members of large gene families. With increasing analyses of plant genomes, more such miRNA-phasiRNA pathways may be discovered.

The LG3 miRFBX Cluster Recently Evolved from *FBX* Genes

To our knowledge, the miRFBX-*FBX* network has never been reported in other plant species; we are curious about how this novel miRFBX-*FBX* network evolved. Prior publications indicated that new *MIRNA*s could evolve from target genes, preexisting *MIRNA* genes, repetitive elements, or random sequences (Allen

et al., 2004; Felippes et al., 2008; Piriyaopongsa and Jordan, 2008; Xia et al., 2013). We found that the LG3 miRFBX cluster region is expanded significantly in the *F. vesca* genome when compared with its syntenic region in closely related species (Fig. 5A; Supplemental Fig. S4). Furthermore, the *MIRFBX3/4* and *MIRFBX5/6* in the LG3 cluster showed significant similarity to the *FBX* genes (Fig. 5, C and D), suggesting that the LG3 miRFBXs likely evolved from duplications of *FBX* genes. We propose a model to illustrate the evolution of miRFBXs and the miRFBX-regulated network of *FBX* genes (Fig. 8). First, inverted duplication of an *FBX* gene (or gene fragment) led to transcripts capable of forming stem-loop structures (Fig. 8A). Gradually, the stem-loop structure was adopted by the miRNA processing machinery to yield miRNA and the locus became the *MIRNA* gene (Fig. 8B). Through subsequent duplications, a single *MIRNA* became a cluster of *MIRNAs* (Fig. 8C). Over time, the clustered *MIRNAs* diverged from one another and yielded a family of miRNAs (Fig. 8D) that target different *FBX* genes (Fig. 8, E and F). The 22-nucleotide miRNAs (i.e. miRFBX7) not only cleave their target *FBX* genes (*FBX-a*) but also primed their target genes (*FBX-a*) for phasiRNA production (Fig. 8E). phasiRNAs subsequently amplify the silencing effect by targeting additional *FBX* genes in cis to regulate their cognate gene (*FBX-a*) or in trans to regulate other *FBX* genes (*FBX-b* and *FBX-n*; Fig. 8G).

The miRFBX7-*FBX*-phasiRNA Network May Function in the Receptacle of Strawberry

FBX genes encode the substrate-recognition subunits of the SCF (for S-phase kinase associated protein1-Cullin-F Box) ubiquitin ligases and constitute one of the largest gene families, with 820 members in *F. vesca* (Xu et al., 2009; Hollender et al., 2014). *FBX* proteins regulate diverse biological processes and were grouped into two general types based on whether they function in conserved processes, such as embryogenesis and circadian rhythms, or relatively specialized processes, such as pollen recognition and pathogen response (Xu et al., 2009). Xu et al. (2009) noted that those with functions in conserved processes experienced little expansion in gene numbers during the evolution from the common ancestor of eudicots and monocots, while those in specialized processes experienced frequent gene duplications, in particular tandem duplications that may be critical to the evolution of specialized functions. Indeed, most of the *FBX* genes targeted by five-miRFBXs reported here are more closely grouped with *FBA1/3*, which belongs to the second group with specialized functions. This is consistent with our finding that miRFBX-targeted *FBX* genes experienced significant gene duplications, as shown by their adjacent chromosomal locations in the *F. vesca* genome (Fig. 6C). This observation indicates that the *FBX*-targeting miRFBXs may have evolved to regulate strawberry-specific processes, which is further supported by the expression pattern of precursor RNAs

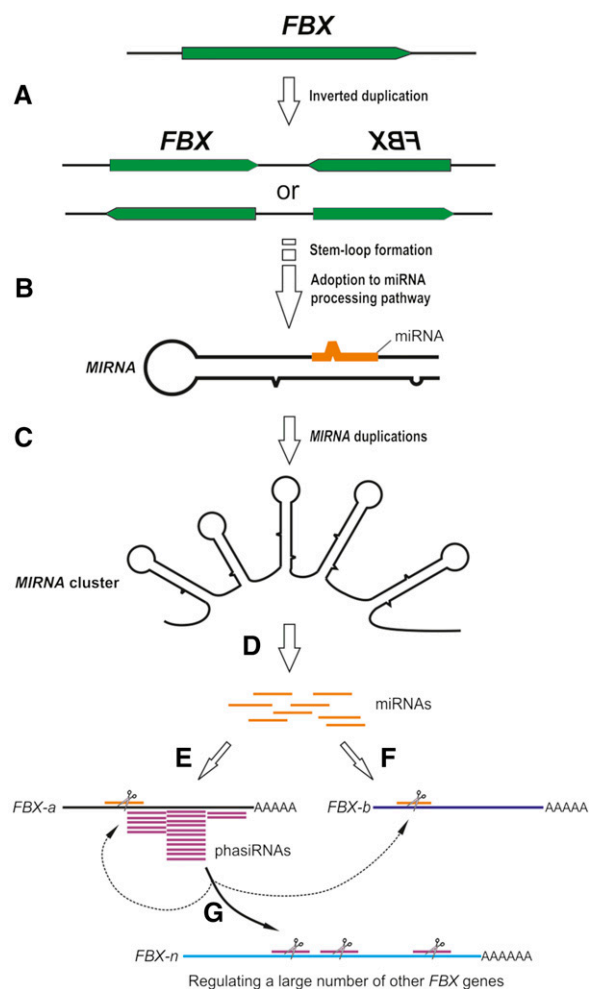


Figure 8. A model illustrating the evolution and mode of action of LG3 miRFBXs in *F. vesca*. A, The stem-loop structure of a *MIRNA* gene came from inverted duplication of an *FBX* gene (or gene fragment). B, The stem-loop structure was gradually adopted by the miRNA processing machinery to yield a miRNA, and the locus became the *MIRNA* gene. C, Subsequent local duplication of the initial *MIRNA* gene led to the *MIRNA* cluster consisting of five stem loops. D, Each *MIRNA* gene diverges over time to give rise to a family of related miRNAs (orange), which were capable of regulating a large number of *FBX* genes. E, The 22-nucleotide miRNAs (orange) produced from *MIRFBX7* guided the initial cleavage of its *FBX* target gene (*FBX-a*), which is followed by phasiRNA (purple) production. F, The 21-nucleotide miRNAs guided the cleavage of their target *FBX* genes (e.g. *FBX-b*). G, The phasiRNAs (purple) derived from E further amplified the silencing effect by guiding the degradation of additional *FBX* genes (*FBX-a*, *FBX-b*, and *FBX-n*) in cis and trans.

for miRFBX7/8. Specifically, the precursor RNA for miRFBX7/8 (transcript 4) was highly expressed in cortex and pith (Fig. 6B). The pith and cortex are the inner and outer tissues of the receptacle, which is a unique part of the strawberry flower that enlarges rapidly to form the edible fleshy fruit. Hence, the miRFBX7-*FBX*-phasiRNA network may function in the receptacle and be responsible for the unique properties of the strawberry fruit. Most of the *FBX* genes in the miRFBX7-*FBX*-phasiRNA network are homologous

to the Arabidopsis Constitutive expresser of PR1 (CPR1)/CPR30 (*At4g12560*; Supplemental Table S6). The Arabidopsis CPR1/CPR30 FBX proteins were shown to negatively regulate the accumulation of a Resistance protein, Suppressor of *npr1-1*, Constitutive1, via the SCF complex (Gou et al., 2012). By cleaving CPR1/CPR30-type *FBX* transcripts, miRFBX could act to promote Resistance protein accumulation and enhance disease resistance in the developing receptacle fruit.

MATERIALS AND METHODS

Plant Materials

A seventh generation inbred line of *Fragaria vesca*, Yellow Wonder 5AF7 (Slovin et al., 2009), was grown in soil in growth chambers under 12 h of light as described previously (Hollender et al., 2012). Young unexpanded leaves of 6-month-old plants, unopened flowers between stages 8 and 12 (staging according to Hollender et al. [2012]), open flowers (within 1 d of opening), and receptacles (10 DPA) were harvested with hand dissection to remove the achenes from each receptacle (Fig. 1). Achenes (fertilized ovaries) were dissected open to separate the ovary wall from the seed inside. The ovary wall and seeds were harvested at 4 and 10 DPA, respectively.

For seedlings, seeds were surface sterilized, plated on Murashige and Skoog medium, stratified at 4°C for 1 week in darkness, and transferred to a growth chamber with 12 h of light. Four-week-old seedlings were collected.

RNA Extraction and Sequencing

For small RNA sequencing, total RNA was extracted using the RNeasy Plant Mini Kit (Qiagen) with certain modifications. Specifically, the supernatant, passed through a QIAshredder spin column, was precipitated in 1.5 volumes of ethanol and then centrifuged for 2 min at 12,000 rpm. The pellet was dissolved in 50 μ L of water and mixed with 500 μ L of Qiazol and then 140 μ L of chloroform. After centrifugation, the supernatant was transferred to an RNeasy mini spin column for further purification following the instructions from the RNeasy Plant Mini Kit. The quality of total RNA was checked by the Experion Automated Electrophoresis System (Bio-Rad). For library preparation following the Illumina TruSeq Small RNA Sample Preparation protocol, 10 to 20 μ g of total RNA per sample with RNA integrity number above 7.8 was sent to the Weill Cornell Medical College Genomics Resources Core Facility. The libraries were sequenced on the Illumina HiSeq 2000 platform.

For PARE analysis, total RNA from different tissues (described above) was extracted using the RNeasy Plant Mini Kit following the manufacturer's instructions. Equal amounts of RNA from different tissues were pooled to form three samples: vegetative (leaf and seedling), flower (open and unopened), and fruit (achene and receptacles). For the achene tissue, both 4- and 10-d-old achenes were scraped off the receptacle. Approximately 100 to 150 μ g of total RNA per sample was sent to the Beijing Genomics Institute for PARE library construction and sequencing on the Illumina HiSeq 2000 platform. Prior to trimming, 50-bp single-end-sequence reads were obtained (Supplemental Table S1).

miRNA Identification

miRNA identification was based on previously well-established criteria (Meyers et al., 2008; Xia et al., 2012) and is summarized in Supplemental Figure S1. The raw reads were processed by first discarding low-quality reads, then removing adaptors, and finally collapsing identical small RNA reads into one using the FASTX-Toolkit (http://hannonlab.cshl.edu/fastx_toolkit/index.html). Reads homologous to other noncoding RNAs, like tRNA and ribosomal RNAs, were removed by BLASTN alignment against Rfam 10, allowing at most two mismatches. The remaining small RNAs were subjected to miRNA identification. The prescreen for miRNAs relied on several criteria: (1) reads had to map exactly to the genome at no more than 16 loci; (2) reads had to be 20 to 22 nucleotides in length; (3) raw read counts had to be 10 or higher in at least one of the nine libraries; and (4) the novel miRNA had to derive from a well-supported stem-loop structure (Supplemental Fig. S3), defined as four or fewer mispairings and one or fewer central bulge in the region of miRNA:miRNA* pairing. Afterward, conserved miRNAs were identified using BLASTN against mature

miRNAs in miRBase (release 20; <http://www.mirbase.org/>), allowing up to two base differences. Those small RNAs, with no hit in the miRBase and combined miRNA/miRNA* reads accounting for more than 75% of reads in the precursor locus (spanning 150 bp upstream to 150 bp downstream of the small RNA), were defined as novel miRNAs. The putative precursor sequences were folded with the Vienna RNA package (Hofacker, 2003).

The total number of reads that perfectly match the *F. vesca* genome in a given library was used for the normalization of read abundance. Each miRNA was normalized to 10 million reads. *F. vesca* genome sequences (*fvesca_v1.0_scaffolds.fna.gz*) were downloaded from the GDR (<http://www.rosaceae.org/>; Shulaev et al., 2011).

miRNA-Target Analysis

Targetfinder 1.6 (Fahlgren et al., 2007) was used to predict miRNA targets, and CleaveLand version 2.0 (Addo-Quaye et al., 2009) was then applied to analyze the PARE data after adaptor trimming and genomic mapping. The *F. vesca* transcriptome (GeneMark hybrid coding sequence) was retrieved from the GDR (<http://www.rosaceae.org/>; Shulaev et al., 2011). The alignment penalty score, assigned by Targetfinder 1.6, was used in target prediction. Only matches with scores of 5 or less were retained. Two parameters, the alignment score and the category, were used to evaluate the reliability of target genes (Supplemental Tables S4 and S5). An alignment score up to 5 was used in target prediction (Li et al., 2010; Xia et al., 2012); a lower score indicates a better alignment between miRNA and its target. The category score (between 0 and 4), on the other hand, considers the ratio of the PARE tag reads mapped to the corresponding miRNA-mediated cleavage site to the total number of PARE tags in a given target mRNA (Addo-Quaye et al., 2009). The lower the category score, the higher the confidence of the cleaved targets.

RNA-seq Analysis

Seventy-four RNA-seq libraries from 37 different *F. vesca* flower, fruit, leaf, and seedling tissues (i.e. two replicates per tissue) were generated in our earlier work (Kang et al., 2013; Hollender et al., 2014). RNA-seq reads in the LG3 five-miRFBX cluster region were visualized from GBrowse hosted at the GDR (http://www.rosaceae.org/gb/gbrowse/fragaria-vesca_v1.1-1g/). A screenshot from this was incorporated into Figure 6A. Furthermore, RNA-seq mapped reads (RPKM) of the precursor transcripts at the LG3 and LG5 clusters as well as the 24 predicted *FBX* target genes of miRFBX7 (Supplemental Table S6) were extracted from the RNA-seq libraries (Kang et al., 2013; Hollender et al., 2014) and imported into MultiExperimental Viewer MeV4.8 (Saeed et al., 2003), respectively. RPKM values across the 37 tissues of the precursor transcripts at the LG3 and LG5 clusters are shown as a heat map (Fig. 6B). The Normalize Genes/Rows function in MeV4.8 was used to normalize each of the 24 *FBX* genes across all 74 libraries. Hierarchical clustering with Pearson correlation was then used to display the relative expression trend of the 24 *FBX* genes targeted by miRFBX7 (Fig. 6C).

TAS Gene and PHAS F-Box Gene Identification

Phasing analysis of secondary siRNAs was conducted as described previously (Xia et al., 2013). The same algorithms were used for *P* value and phasing score calculation. Genes with $P < 0.001$ were considered as *TAS* or *PHAS* genes. The Integrative Genomics Viewer (Thorvaldsdóttir et al., 2013) was used to view the phasing score.

F-Box Gene Classification and Phylogenetic Tree Construction

The 691 Arabidopsis (*Arabidopsis thaliana*) *FBX* gene sequences were retrieved from the data of Xu et al. (2009), and the classification of Arabidopsis *FBX* genes was based on the domain annotation in the same work. For the 94 *F. vesca* *FBX* genes targeted by miRFBXs, de novo domain structure annotation was performed using CD-search (<http://www.ncbi.nlm.nih.gov/Structure/cdd/wrpsb.cgi>). Among the 94 genes, the gene bodies of three genes (*gene26143*, *gene00154*, and *gene10432*) were corrected from the original sequences due to inaccurate annotations, and two genes (*gene18163* and *gene01506*) were separated into two additional genes. In total, 787 *FBX* genes (697 from Arabidopsis and 96 from *F. vesca*) were used for the construction of phylogenetic trees. First, their full-length amino acid sequences were aligned

using the multiple alignment tool MUSCLE (Edgar, 2004). Then, the tree was constructed by FastTree using default settings (Price et al., 2009). The resulting tree was viewed and annotated using Dendroscope 3 (Huson and Scornavacca, 2012).

Synteny Analysis of the *F. vesca* LG3 miRFBX Region

Syntenic regions among plants and their alignments were retrieved from the Plant Genome Duplication Database (<http://chibba.agtec.uga.edu/duplication/index/home>). BLASTX and BLASTP analyses were conducted locally with an e-value cutoff of 1,000, and plots of BLASTP e-value distribution were generated using the R package.

Raw and processed reads of all nine small RNA libraries and three PARE libraries are available from the Gene Expression Omnibus at the National Center for Biotechnology Information (<http://www.ncbi.nlm.nih.gov/geo>). With the exception of the open flower small RNA library (accession no. GSE44930), the eight small RNA libraries and three PARE libraries are deposited under the accession number GSE61798.

Supplemental Data

The following supplemental materials are available.

Supplemental Figure S1. Informatics pipeline for predicting miRNAs from *F. vesca*.

Supplemental Figure S2. Alignment of miRFBX precursors of the LG3 miRFBX cluster.

Supplemental Figure S3. Secondary structures of 32 novel miRNA precursors.

Supplemental Figure S4. Syntenic analysis in several plant species for the region containing the LG3 miRFBX cluster.

Supplemental Figure S5. Read abundance of mature miRFBXs.

Supplemental Table S1. Summary of sequence reads of nine small RNA libraries and three PARE libraries from *F. vesca* (in millions).

Supplemental Table S2. Known miRNAs in *F. vesca*.

Supplemental Table S3. Novel miRNAs identified in *F. vesca*.

Supplemental Table S4. Target genes of known miRNAs in *F. vesca*.

Supplemental Table S5. Target genes of novel miRNAs in *F. vesca*.

Supplemental Table S6. *FBX* target genes of miRFBX1-9.

Supplemental Table S7. A list of 24 *PHAS FBX* genes targeted by miRFBX7.

Supplemental Table S8. *PHAS MYB* genes targeted by miR828.

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