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The *Rosa chinensis* cv. Viridiflora Phyllody Phenotype Is Associated with Misexpression of Flower Organ Identity Genes

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Phyllody is a flower abnormality in which leaf-like structures replace flower organs in all whorls. Here, we investigated the origin and the molecular mechanism of phyllody phenotype in Rosa chinensis cv. Viridiflora, an ancient naturally occurring Chinese mutant cultivar. Reciprocal grafting experiments and microscopy analyses, demonstrated that the phyllody phenotype in Viridiflora is not associated with phytoplasmas infection. Transcriptome comparisons by the mean of RNA-Seq identified 672 up-regulated and 666 down-regulated genes in Viridiflora compared to its closely related genotype R. chinensis cv. Old Blush. A fraction of these genes are putative homologs of genes known to be involved in flower initiation and development. We show that in flower whorl 2 of Viridiflora, a down-regulation of the floral organ identity genes RcPISTILLATA (RcPI), RcAPETALA3 (RcAP3) and RcSEPALLATA3 (RcSEP3), together with an up-regulation of the putative homolog of the gene SUPPRESSOR of OVEREXPRESSION of CONSTANS1 (RcSOC1) are likely at the origin of the loss of petal identity and leaf-like structures formation. In whorl 3 of Viridiflora, ectopic expression of RCAPETALA2 (RCAP2) along with the down regulation of RCPI, RCAP3, and RCSEP3 is associated with loss of stamens identity and leaf-like structures formation. In whorl 4. the ectopic expression of RcAP2 associated with a down-regulation of RcSEP3 and of the C-class gene RcAGAMOUS correlate with loss of pistil identity. The latter also suggested the antagonist effect between the A and C class genes in the rose. Together, these data suggest that modified expression of the ABCE flower organ identity genes is associated with the phyllody phenotype in the rose Viridiflora and that these genes are important for normal flower organs development.

Keywords: rose, Viridiflora, phyllody, transcriptome analysis, ABC flower organ's identity genes, RcSOC1

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INTRODUCTION

Roses have been cultivated by humans since antiquity, as early as 3 000 B.C. in China. Today there exist more than 35 000 rose cultivars. These modern rose cultivars were established from less than 10 rose species, including Chinese species (Wylie, 1954; Liu, 1964). Chinese old roses have high ornamental, cultural, economic values, and represent an important rose germplasms resource (Zhang and Zhu, 2006; Yan et al., 2014). Among old Chinese rose species, the recurrent blooming Rosa chinensis was used in many breeding programs to select for the most advantageous traits such as recurrent blooming, scent and resistance to pathogens (Martin et al., 2001; Ku and Robertson, 2003; Bendahmane et al., 2013). R. chinensis cv. Viridiflora (hereafter Viridiflora) is a rose cultivar, commonly known as the 'green rose' (Chmelmitsky et al., 2002), in which petals, stamens and pistils are converted into leaf-like organs (Figures 1A,C). This phenomenon is also known as phyllody, a phenotype that has been described in many plant species (Meyer, 1966; Mor and Zieslin, 1992). Viridiflora is a spontaneous mutant of R. chinensis Jaquin (Krussman, 1981). This stable mutant has been maintained for over 200 years in China, Europe and America, and is widely used for ornamental horticulture.

The prevailing ABCE genetic model of floral organ identity determination, with the combinatorial activity of four classes of homeotic genes (A, B, C, and E), has been widely characterized in many flowering species. Most of the ABCE genes encode MADS-box transcription factors (Coen and Meyerowitz, 1991; Smaczniak et al., 2012). In Arabidopsis, the class A-class genes APETALA1 (AP1) and APETALA2 (AP2) specify sepal organ identity and development; A-class genes and the B-class genes APETALA3 (AP3) and PISTILLATA (PI) together determine petal organ identity and development; B-class genes together with the C-class gene AGAMOUS (AG) are required for stamen formation and the C-class gene AG is required for carpel formation (Weigel and Meyerowitz, 1994). The E-class genes SEPALLATA (SEP1, SEP2, SEP3, SEP4) interact with A, B, and C class genes and are involved in specifying floral organs in all flower whorls. The ABCE model is relatively conserved in the flowering plants with few variations (Heijmans et al., 2012; Smaczniak et al., 2012; Wellmer et al., 2014).

In *Rosa* sp., homologs of A-class (*RhAP1, RhAP2*), B-class (*RhAP3, RhP1*), C-class (*RhAG*) and E-class genes have been reported (Dubois et al., 2012; Bendahmane et al., 2013). Expression analysis and overexpression experiments of these rose MADS-encoding cDNAs in *Arabidopsis* suggested their role in flower organ identity determination in the rose (Kitahara and Matsumoto, 2000; Kitahara et al., 2004; Hibino et al., 2006; Dubois et al., 2010, 2011, 2012; Ma et al., 2015).

In *Arabidopsis*, loss of function of A, B, or C genes lead to homeotic conversion of floral organs and floral aberrations that are different from the phyllody-like phenotype (Coen and Meyerowitz, 1991). In the rose it was demonstrated that misexpression or down-regulation of the *RhAG* in whorl 3 was associated with homeotic conversion of stamens to petals and double flower formation (Dubois et al., 2010). In *Arabidopsis*, the triple *SEP1/2/3* mutant produces flowers in which all organs

develop as sepals (Pelaz et al., 2000). It was also reported in many plants that, infection by phytoplasmas was also reported to lead to phyllody (McCoy et al., 1989; Szyndel, 2003; Hogenhout et al., 2008). In *Arabidopsis*, it was recently shown that the phytoplasma-secreted protein PHYL1 is involved in the targeted protein degradation of the organ identity proteins SEP3 and AP1, which in turn lead to the transformation of flower organs into leaf-like structures (Maejima et al., 2014, 2015).

In this study, we addressed the cause of phyllody phenotype in the rose Viridiflora. We show that phyllody phenotype in Viridiflora is not caused by Mycoplasma like organism (MLO) infection. To identify the molecular basis of the malformed flowers in Viridiflora, we used a transcriptomic approach to compare gene expression in flowers of *R. chinensis* cv. Old Blush (hereafter Old Blush) and Viridiflora. Our study identified that the green flower phenotype (phyllody) in *R. chinensis* cv. Viridiflora is associated with misexpression of the putative homologs of the flowering integrator *RcSOC1* and of ABCE flower organ identity genes *RcAP1*, *RcAP2*, *RcPI*, *RcAG*, and *RcSEP3*.

MATERIALS AND METHODS

Plant Materials and Grafting Experiment

Rosa chinensis cv. Old Blush (**Figures 1B,D**) and its variant *R. chinensis* cv. Viridiflora, in which petals, stamens, and pistils are converted to leaf-like organs (**Figures 1A,C**), were field-grown in Flower Research Institute of Yunnan, Academy of Agricultural Science. A part from the phyllody phenotype, no other phenotypic differences were observed between Old Blush and Viridiflora (**Figures 1A,C**). Flower buds at stage 8–10 mm were collected and then immediately frozen in liquid nitrogen until RNA extraction.

Grafting experiments were performed as previously described (Ohkawa, 1980). Old Blush young shoots, used as scion, were grafted on Viridiflora plants used as rootstock. As control Viridiflora young shoots, used as scion, were grafted on Old Blush plants used as rootstock (**Figures 2A,C**). Twenty shoot grafting experiments were performed for each.

Microscopy and PCR Experiment to Detect Mycoplasma

To detect mycoplasma, samples of about 2 mm² were cut from young flower buds, then crude sap extracts were prepared from these tissues in 0.01 M potassium phosphate pH 7.0 buffer and then applied to 4% Formvar-coated, carbon-stabilized copper grids (300 mesh). Grids were then stained with 2% uranyl acetate for 10 s, rinsed with 0.01 M potassium phosphate pH 7.0 buffer and water (Yin et al., 2014). Grids were dried and examined on a JEM-100CX II transmission electron microscope (JOEL, Ltd, Tokyo, Japan).

Total DNA was extracted from young flower buds using Plant DNA Isolation Reagent (TakaRa, China) and then used to PCR amplify MLO DNA using the MLO specific primers R16mF/R16mR (Lee et al., 1993; **Supplementary Table S1**). A construct harboring MLO DNA was used as positive control.

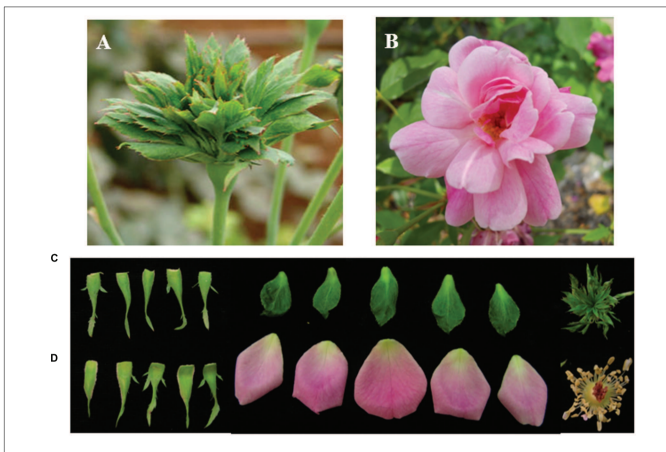


FIGURE 1 | Flower phenotype of *Rosa chinensis* cv. Viridiflora (A,C) and Old Blush (B,D). (C,D) Dissected flower organs of Viridiflora and Old Blush, respectively.

Library Construction and RNA-Sequencing

Total RNA was extracted using TRIzol reagent and contaminating DNA was then removed by treatment with RNase-free DNase. NanoDrop and Agilent Technologies 2100 Bioanalyzer were used to quantify the total RNA before RNA-Seq library construction.

Transcriptome libraries of Viridiflora and Old Blush were constructed using Illumina TruSeq RNA sample preparation Kit V2 following the manufacturer's instructions. In brief, total RNA was purified using Magnetic Oligo (dT) beads. Then, purified RNA was sheared to approximately 330 nucleotides fragments and primed for cDNA synthesis using random primers. Subsequently, the fragments were ligated to sequencing adapters. Using agarose gel electrophoresis, the suitable fragments with 400–500 nt sizes were selected as templates for PCR amplification and the final PCR products were sequenced using Illumina HiSeq 2500 system as 125 nt, paired-end Illumina reads.

Transcriptome Assembly and Annotation

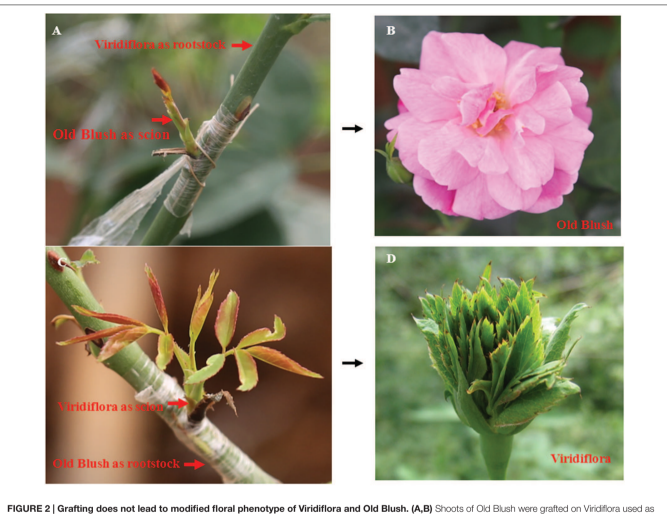
High-quality transcript sequences were obtained by *de novo* transcriptome assembly of the RNA-seq reads, using Trinity software (version: r20140413p1; Langmead and Salzberg,

2012) with default parameters. Open reading frame (ORF) identification was identified using Transdecoder (Grabherr et al., 2011) with default parameters. Homology sequences search was performed against the UniProt databases by ncbi-blast-2.2.27+ with an E-value of 10^{-6} . The GO terms were assigned to each assembled sequences based on the UniProt databases. The all information of transcriptome data is available in the database¹.

Gene Expression Analysis and DEG Identification

The read counts for each transcript were calculated after aligning the RNA-Seq reads on the assembled transcriptome using Bowtie2 2.1.0 (Langmead and Salzberg, 2012). To accurately measure gene expression level, we only retained pairs of reads having both ends matching on the same transcript. The expression level of each transcript was normalized as the fragments per kilobase of exon per million fragments mapped (FPKM), which is analogous to single-read RPKM for single reads (Mortazavi et al., 2008). The *P*-value were calculated according to the statistical R package DEGseq (Wang et al., 2010) using MA-plot-based method with random sampling model. Then, the

¹http://bioinfor.org/pub/Viridiflora



rigure 2 Gratting does not lead to modified floral phenotype of Viridifiora and Old Blush. (A,B) Shoots of Old Blush were gratted on Viridifiora used as rootstock. Mature flower of Old Blush with Viridifiora as rootstock show wild-type Old Blush flower phenotype (B); (C,D) Shoots of Viridifiora were grafted on Old Blush used as rootstock. Mature flower of Viridifiora with Old Blush as rootstock show phyllody phenotype (D).

differentially expressed genes were identified using fold-change >3 and a P<0.001 as the threshold.

Validation of Differential Expressed Genes

The sequences of the genes used for qRT-PCR validation were provided (details listed in **Supplementary Data Sheet S1**). Genespecific primers (**Supplementary Table S1**) were designed using Primer3². Total RNAs were extracted from leaves and from sepals, petals, stamens, and pistils at 8–10 mm flower buds development stage using TRIzol RNA purification kit (TaKaRa, China). One microgram of total RNA was used in reverse transcription in a total reaction volume of 20 μ L in the presence of 6-mer random primers and an oligo primer according to the protocol provided by manufacturer TaKaRa (Yan et al., 2011). The standard curve for each gene was obtained by real-time PCR with five dilutions of cDNA. The reactions were performed in 20 μ L volumes each containing 10 μ L 2× SYBR Green

²http://bioinfo.ut.ee/primer3/

Mastermix (TaKaRa), 300 nM of each primer and 2 μ L of 10-fold diluted cDNA template (Yan et al., 2014). The PCR reactions were run in a Bio-Rad Sequence Detection System. Three biological replicates were performed for each analysis. *RhGAPDH* (AB370120) was used as control. Quantification of gene relative expression in different organs was performed using the delta-delta Ct method as described by Livak and Schmittgen (2001). All data were expressed as the mean \pm standard deviation (SD) after normalization.

RESULTS

Phyllody Phenotype of *R. chinensis* cv. Viridiflora Is Not Associated with Phytoplasma

Viridiflora refers to a floral aberration in which petals, stamens, and pistils are converted to leaf-like organs (Figures 1A,C) a phenotype that resembles phytoplasmas infection-induced phyllody phenotype. Phytoplasmas are non-cultivable microorganisms transmitted by contact with infected plants. The grafting mechanical contact between the rootstock and the scion enables the spread of this disease and this approach is used generally to detect phytoplasmas (Golino et al., 1989; Aldaghi et al., 2007; Goldschmidt, 2014). To address if Viridiflora was associated with mycoplasma infection, we used grafting approach in which Viridiflora was used as rootstock and healthy Old Blush was used as scion (Figure 2A) and vice versa, used as control (Figure 2C). Twenty bud grafting experiments were performed for each. These data showed that all Old Blush buds with Viridiflora as rootstock bloomed normally and flower organs phenotype was identical to non-grafted Old Blush control plants (Figure 2B). At the same time, all Viridiflora flowers with Old Blush as rootstock exhibited phyllody phenotype similar to the none grafted Viridiflora control plants (Figure 2D). It should be noted that flowers of Old Bush used as rootstock, showed no phyllody phenotype. These data suggest that phyllody phenotype in Viridiflora is not a result of phytoplasma infection. In agreement with these data, transmission electron microscopy or PCR experiments identified no trace of phytoplasma or phytoplasma DNA respectively, in young buds of Viridiflora and Old Blush (Supplementary Figures S1 and S2). Taken together these data strongly suggest that the flower mutant phenotype of Viridiflora was not caused by phytoplasma infection.

Flower Buds Transcriptome Comparison in Viridiflora and Old Blush

To investigate further the molecular basis of the phyllody phenotype, we compared the transcriptome in flowers of Viridiflora and Old Blush. Total RNA was prepared from flower buds at 8-10 mm development stage and then used to build two libraries for high-throughput sequencing. A total of 40 and 44 million reads were generated from Viridiflora and Old Blush samples, respectively. The transcriptome assembly sequences of R. chinensis cv. Viridiflora has been deposited at the Database of Transcriptome Shotgun Assembly (TSA) at DDBJ/EMBL/GenBank under the accession GETJ00000000. The transcriptome data are available in the database¹. In total, five gigabase for each library were produced. Paired-end reads were assembled into transcript sequences and yielded 68,565 unisequences with an average length of 887 bp (Figure 3A). Approximately 91% of the reads from Viridiflora and Old Blush could be successfully aligned to the assembled transcripts, used as reference sequences (Table 1). In total, over 32,224,620 and 34,200,388 reads were mapped back in pairs with the unique location for Viridiflora and Old Blush libraries, respectively (Table 1).

Pairs of reads with a unique location were then used to estimate genes expression levels. Read counts were normalized as FPKM (Mortazavi et al., 2008) and genes with a minimum expression threshold corresponding to FPKM > 3 were kept. In total, 20,017 and 20,859 genes were considered as expressed in the libraries of Viridiflora and Old Blush, respectively (**Table 1**). The average FPKMs for Viridiflora and Old Blush were 33.18 and 32.57, respectively (**Figure 3B**).

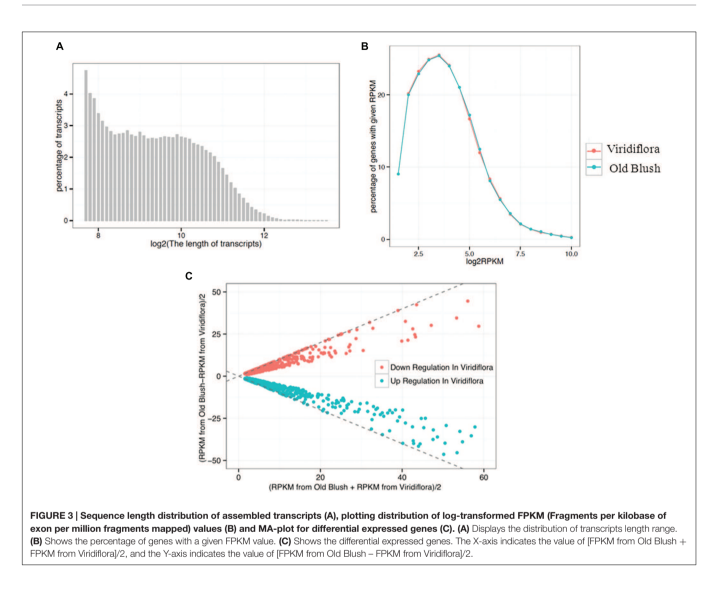
Genes differentially expressed between of Viridiflora and Old Blush were investigated using the MA-plot-based method with the random sampling model (Wang et al., 2010). 672 unisequences were up-regulated and 666 were down-regulated (Figure 3C) in Viridiflora compared to Old Blush, with a fold change of at least 3 and a P-value < 0.001 (Table 1). Flower development genes that exhibited significant expression difference between Old Blush and Viridiflora were for RTqPCR validation. Expression of 24 genes among the differentially expressed genes was analyzed using RT-qPCR (Figures 4A,B). A good correlation (Pearson correlation coefficient = 0.98) between RT-qPCR data and the RNA-Seq data was observed for the 24 analyzed genes, suggesting that our transcriptomics data are accurate in the different tissues and experimental conditions (Figure 4C). GO enrichment analysis showed that transcription factors were enriched in the up-regulated genes fraction, while GO terms associated with 'pollen wall assembly,' 'pollen exine formation,' and 'hormone catabolic process' were enriched in the down-regulated genes fraction (Figures 5A,B).

The expressions of the rose homologs of the flower development genes PI (c20259_g2), AG (c18211_g1), AGL9 (c26880_g6), AGL15 (c24715_g4; Table 2), Pollen-specific protein SF3 (Baltz et al., 1992; c15299_g3) and Gibberellin 20 oxidase 1 (c25215_g1; Jacobsen and Olszewski, 1991; Goto and Pharris, 1999) were significantly repressed in Viridiflora compared to Old Blush (Table 2). Genes expressions of putative homologs of other flower development transcription factors, such as AP1 (c16755_g1), AP2 (c27740_g2), SOC1 (c24463_g2), AGL6 (c17791_g1), AGL8 (c17660_g1; Table 2) were in contrast upregulated in Viridiflora flowers compared to Old Blush flowers. In addition, the bHLH transcription factor UNE10 (Toledo-Ortiz et al., 2003; c18395_g2) and homeotic gene BEL1 (c20518_g1) were also significantly up-regulated (Table 2). BEL1 is known to act as negative regulator of AG and has a major role in ovule patterning and in determination of integument identity via its interaction with MADS-box factors (Ray et al., 1994; Mizumoto et al., 2011; Sharma et al., 2014). Taken together these data show that the phyllody phenotype in Viridiflora is mainly associated with modified expression of flowering and flower development related genes.

Expression Profiles of MADS-Box Transcription Factor Coding Genes

To further investigate the origin of the phyllody phenotype, the expression of MADS-box flower organ identity genes *RcAP1* (c16755_g1), *RcAP2* (c27740_g2), *RcPI* (c20259_g2), *RcAP3* (RC000216; Dubois et al., 2012), *RcAG* (c18211_g1), *RcSEP1* (RC001958; Dubois et al., 2012), and *RcSEP3* (RC000799; Dubois et al., 2012) and flowering *RcSOC1* (c24463_g2) were analyzed using RT-qPCR in leaves and in sepals, petals, stamens, and pistils whorls of developing flowers of Viridiflora and Old Blush.

In wild-type flowers of Old Blush, the A-class genes *RcAP1* mRNA (c16755_g1) accumulated to high levels in whorl 1 (sepals) and showed very low or no expression in whorls 2 (petals), 3 (stamens), and 4 (pistils; **Figure 6A**), thus in agreement with previously published data in *Arabidopsis* (Gustafson-Brown

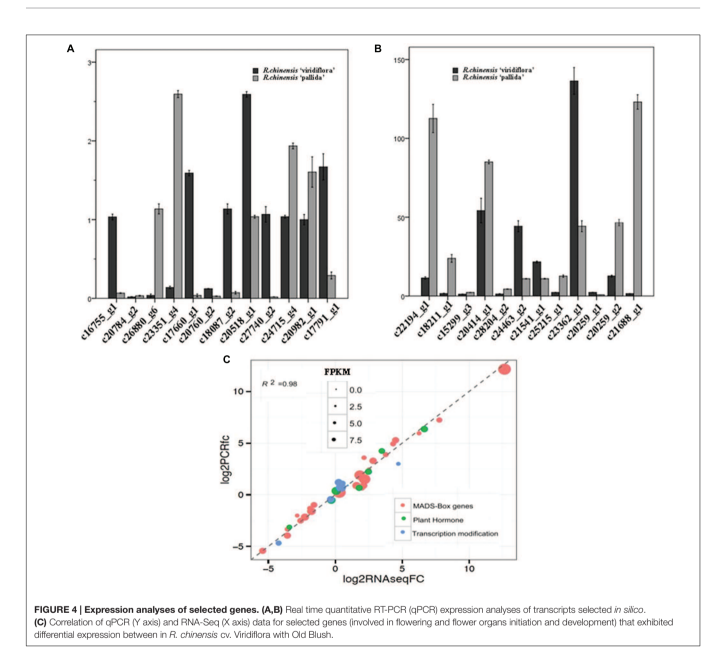


et al., 1994). mRNA of the A-class *RcAP2* was expressed in whorls 1 and 2 of Old Blush (sepal and petal, respectively; **Figure 6B**) and showed relative low expression level in the fourth whorls (pistil; **Figure 6B**), thus similar to previously reported data in *Arabidopsis* (Jofuku et al., 1994). The B-class genes *RcP1* and *RcAP3* were expressed in second and third whorls (petals and stamens respectively) of Old Blush (**Figure 6C**), thus similar pattern as their *Arabidopsis* counterparts, *PI* and *AP3* (Krizek and Meyerowitz, 1996). As expected, the C-class gene *RcAG* (c18211_g1) was expressed in stamens (whorl 3) and in pistils (whorl 4) of Old Blush. The organ identity *RcSEP1* and *RcSEP3* were expressed in all floral whorls (**Figures 6F,G**). In *Arabidopsis*, *SEP1* is expressed in all whorls while SEP3 is expressed in whorls 2, 3, and 4. These data together suggest that the ABCE genetic model of floral organ identity determination is likely conserved in the rose.

In Viridiflora flowers, the expression of RcAP1 was significantly increased in the sepal and petal whorls (**Figure 6A**). RcAP1 expression was about 1.7 times induced in whorl 1 and strikingly 20-fold induced in whorls 2 compared to that in Old Blush. The expression of RcAP2 was up-regulated in all whorls of Viridiflora flowers. Interestingly, the expression of RcAP2 was significantly up-regulated in stamens and pistils whorls of Viridiflora (**Figure 6B**) and conversely the expression

TABLE 1 Summary of Illumina sequencing and assembly for two RNA-Seq libraries.									
Sample	Reads (PE)	Length	Total bases	GC%	Mapping (ratio)	Unique (ratio)	FPKM > 3		
Viridiflora	20,640,280 × 2	125,125	5,160,070,000	46,46	37,697,457(91%)	16,112,310 × 2(78%)	20,017		
Old Blush	22,052,225 × 2	125,125	5,513,056,250	46,46	39,996,389(91%)	17,100,194 × 2(78%)	20,859		

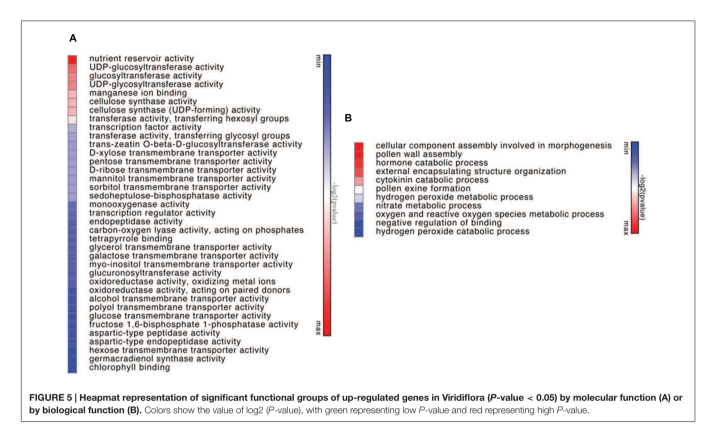
PE, paired-end reads; FPKM, fragments per kilobase of exon per million fragments mapped.



of the C-class gene RcAG was significantly down-regulated in whorl 4 compared to that in Old Blush (**Figure 6E**). It should be noted that RcAG was about eightfold over-expressed in the pistils compared to stamens (**Figure 6E**). These data are also in agreement with the antagonist effect between A and C functions, known as crucial in floral patterning (Gustafson-Brown et al., 1994). Furthermore, RcSEP3 mRNA was about sixfold under-expressed in all flower whorls of Viridiflora compared to Old Blush (**Figure 6G**). However, the expression of RcSEP1 was not strikingly different between the two roses, except for a relative low expression level in whorl 4 (pistils) of Viridiflora compared to Old Blush (**Figure 6F**). The phyllody phenotype in Viridiflora was also associated with significant down-regulation of the B class genes RcP1 and RcAP3 in the petals and stamens whorls (2 and 3, respectively) compared to Old Blush (5- and 6-fold, respectively; **Figures 6C,D**).

Interestingly, in Viridiflora, the expression of the flowering time integrator MADS-box *RcSOC1* gene was induced in petals, stamens and pistils whorls, with the strongest expression in sepals and in petals whorls (**Figure 6H**). In Old Blush, the expression of *RcSOC1* was confined to leaves and although at lower levels to sepals as well, thus a similar pattern as in *Arabidopsis* (Samach et al., 2000).

To summarize, these data show that in whorl 2, the loss of petal identity is consistent with the down-regulation of the B (RcPI and RcAP3) and E-class (RcSEP3) organ identity genes. The loss of petal organ identity combined with the strong up-regulation of RcSOC1 is likely associated with the conversion of petals into leaf-like structures in whorl 2. In whorls 3 and 4, the



ectopic expression of the A-class *RcAP2* and the down-regulation of the E-class *RcSEP3* in combination with the down regulation of B-class genes *RcPI* and *RcAP3* in whorl 3 and of the C-class *RcAG* in whorl 4 is consistent with the observed loss of stamens and pistils identity, respectively, and their conversion into leaf-like structures.

DISCUSSION

Phyllody is a flower abnormality in which leaf-like structures replace flower organs. Two types of phyllody were described in the genus *Rosa*. The first type of phyllody (*R. chinensis* cv. Viridiflora) is a stable mutation known as the 'green rose' characterized by the transformation of all flower organs to leaf-like structures (Krussman, 1981). For the second type, described in *R. x hybrida* cv. Motrea, phyllody is restricted to reproductive organs (third and fourth whorls; Mor and Zieslin, 1992). It was suggested that Viridiflora phenotype might be caused by phytoplasma infection (McCoy et al., 1989; Szyndel, 2003; Zhang and Zhu, 2006). Here we used grafting, transmission electron microscopy and PCR experiments, to demonstrate that phyllody phenotype in Viridiflora is not associated with phytoplasma infection (**Figure 2**).

Gene expression analysis in flowers early during development, showed that the phyllody phenotype in Viridiflora is associated with ectopic expression of the flowering time gene *RcSOC1* and with misexpression of the ABCE flower organ identity genes leading to loss of petal, stamen and pistil organ identity

(**Figure** 7). These data indicate that in Viridiflora a shift of the expression of the A-class gene RcAP2 toward the third and fourth whorls together with significantly reduced expression of B, C, and E gene classes are associated with the observed phyllody phenotype. These data also highlights the antagonist effect of the A and C class gene.

The A-class gene AP1 is known to be involved in sepal and petal organ identity determination. In R. chinensis cv. Old Blush, RcAP1 showed high expression levels in developing sepals and low expression levels in developing petals (Figure 6A). Similar data were observed in R. hybrida (Mibus et al., 2011) and in other flowering plant species (Huijser et al., 1992; Mandel et al., 1992). Interestingly, we observed a strong down-regulation of the B-class genes RcPI and RcAP3 in petals and stamens whorls and of the E class gene RcSEP3 in all whorls. In Arabidopsis genetic and gene expression experiments showed that SEP3 is required for the up-regulation of B and C floral homeotic genes (Castillejo et al., 2005; Kaufmann et al., 2009). It is possible that the downregulation of RcAP3 and RcPI is a result of reduced expression of RcSEP3. In the flowering plants, mutation of the B-class genes leads to conversion of petals into sepals. In Viridiflora the down-regulation of RcPI and RcAP3 cannot explain the leaf-like phenotype of stamens in Viridiflora. It is likely that the up-regulation of SOC1 in the context of the B (RcPI and RcAP3) and E (RcSEP3) genes down-regulation are at least in part at the origin of the loss of petal identity and phyllody phenotype. In whorl 3, the up-regulation of RcAP2 and the down regulation of RcPI, RcAP3 and RcSEP3 are likely at the origin of loss of stamen identity and phylloid organs formation.

Gene ID no.	Description	FPKM fold change	P-value
Up-regulated			
c24463_g2	MADS-box protein SOC1	7.26	1.89e-11
c17716_g1	Auxin-responsive protein IAA26	10.81	9.79e-33
c18087_g2	MADS-box protein SOC1	11.71	2.97e-21
c16755_g1	Floral homeotic protein APETALA 1	11.91	7.32e-104
20554_g1	Cytochrome P450	18.81	1.76e-68
20760_g2	Ethylene-responsive transcription factor ERF105	4.81	3.06e-104
18395_g2	Transcription factor UNE10	19.88	2.17e-42
27740_g2	Floral homeotic protein APETALA2	3.43	3.67e-28
:17660_g1	Agamous-like MADS-box protein AGL8	42.98	5.07e-118
20518_g1	Homeobox protein BEL1	3.68	1.65e-12
18389_g2	Auxin-binding protein ABP19	83.77	0
17791_g1	Agamous-like MADS-box protein AGL6	6.03	4.61e-33
Down-regulated			
18211_g1	Floral homeotic protein AGAMOUS	13.63	1.20e-43
19619_g1	Transcription repressor MYB4	218.60	2.55e-49
26880_g6	Agamous-like MADS-box protein AGL9	20.10	1.59e-28
24715_g4	ABC transporter G family member 15	3.01	5.94e-130
20259_g2	Floral homeotic protein PI/GLO	3.59	0
15299_g3	Pollen-specific protein SF3	4.36	8.21e-25
21688_g1	Gibberellin 20 oxidase 1	100.50	3.24e-144
22194_g2	Floral homeotic protein AGAMOUS	7.10	4.99e-38
20259_g1	Floral homeotic protein FBP1	4.59	0
19919_g1	Transcription factor MYB44	76.32	7.49e-20
28204_g2	Agamous-like MADS-box protein AGL15	3.60	3.86e-15
20414_g1	Auxin response factor 8	4.0	4.03e-05
20982_g1	IAA-amino acid hydrolase ILR1	3.43	4.51e-48
25215_g2	Transcriptional corepressor LEUNIG	17.27	6.95e-13
18107_g1	Stamen-specific protein FIL1	6421.30	0
22194_g1	Floral homeotic protein AGAMOUS	7.11	3.86e-54

TABLE 2 | Genes known to be involved in flower initiation and development, showing either up- or down-regulated expression levels in *R. chinensis* cv. Viridiflora.

AGAMOUS (AG), the C-class floral homeotic gene is involved in the specification of stamens and pistils identity and the termination of the floral meristem in the center of the flower (Kapoor et al., 2002; ÓMaoiléidigh et al., 2013). In Viridiflora, our data show that mRNA of RcAG was down-regulated in stamens and pistils, compared to Old Blush (Figure 6D). In Arabidopsis the down-regulation of AG leads to conversion of stamens into petals (Yanofsky et al., 1990). Similarly, the silencing of pMADS3, a petunia C-class gene, resulted in homeotic conversion of stamens into petaloid structures (Kapoor et al., 2002). Interestingly, in the ranunculid Thalictrum thalictroides, down-regulation of the AG homolog ThtAG1 by the mean of virus-induced gene silencing resulted in homeotic conversion of stamens and pistils into sepaloid organs and loss of flower determinacy (Galimba et al., 2012). We observed that the putative homolog of BEL1 is up-regulated in Viridiflora flowers. In Arabidopsis BEL1 was shown to act as negative regulator of AG (Ray et al., 1994; Sharma et al., 2014). Therefore, we cannot rule out that the down-regulation of RcAG could be a consequence of increased RcBEL1 levels. In addition, RcAP2 showed ectopic expression in stamens and pistils of Viridiflora (Figure 6B). This

result was not surprising, as it is known that AP2 is a negative regulator of AG expression in the first two whorls of the flower (Drews et al., 1991). Therefore, it is likely that in Viridiflora the ectopic expression of RcAP2 in stamens and pistils whorls lead to the observed down-regulation of RcAG expression in whorl 4. It has been reported in *Arabidopsis* that AP2, besides its antagonist role with the C-class gene AG, is also involved in repressing B-class genes AP3 and PI by regulating their outer boundary expression (Krogan et al., 2012; Wuest et al., 2012). It is likely that similar role is conserved in the rose and that in Viridiflora the upregulation of RcAP2 in whorl 3 (**Figure 6B**) is responsible for the observed down-regulation of the B-class genes RcAP3 and RcPI(**Figures 6C,D**) and thus loss of petal and stamens organ identity.

We show that the phyllody phenotype in Viridiflora is associated with ectopic expression of the putative homolog of the flowering time integrator gene *SOC1*. In *Arabidopsis SOC1* plays a central role to integrate the photoperiodic, the autonomous, the vernalization, and the gibberellin pathways during flowering (Borner et al., 2000; Samach et al., 2000; Moon et al., 2003). SOC1-like genes are preferentially expressed in vegetative parts of both angiosperms and gymnosperms (Tandre et al., 1995;

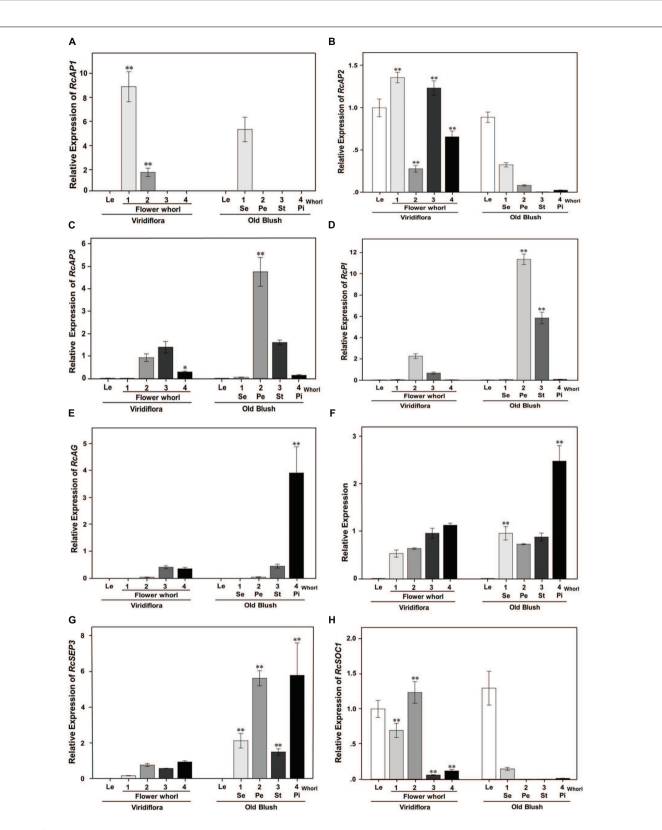
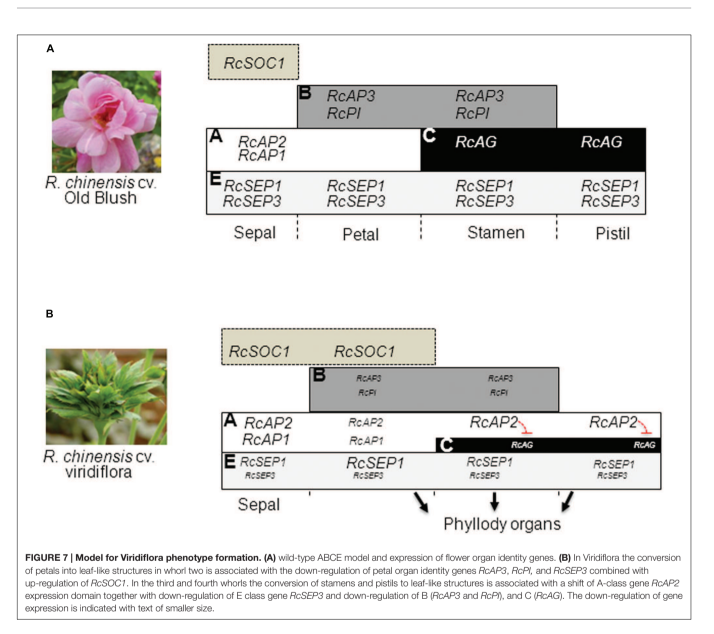


FIGURE 6 | Quantitative RT-PCR expression analyses of genes involved in flowering and in flower organ identity determination and development. Expression of RcAP1(c16755_g1; **A**), RcAP2 (c27740_g2; **B**), RcAP3 (RC000216; **C**), RcP1 (c20259_g2; **D**), RcAG (c18211_g1; **E**), RcSP1(RC001958; **F**), RcSP3 (RC000799; **G**), and RcSOC1(c24463_g2; **H**) was analyzed in developing flower buds, at stage 8–10 mm, of Viridiflora and Old Blush. Le, leaves; 1–4, Flower whorls; Se, sepals; Pe, petal; St, stamens; Pi, pistils. Values are mean \pm standard deviation (n = 3). Asterisks indicate significant differences calculated using Tukey's test (**P < 0.01; *P < 0.05).



Winter et al., 1999; Watson and Brill, 2004), but examples of SOC1-like genes expressed in reproductive organs have also been reported (Heuer et al., 2001). AtSOC1 is mainly expressed in meristems and developing leaves (Samach et al., 2000). Overexpression of AtSOC1-like genes causes early flowering in several plant species (Tadege et al., 2003; Ferrario et al., 2004; Smykal et al., 2007). Interestingly, it has also been shown that overexpression of GhSOC1 in gerbera leads to a partial loss of floral organ identity, but did not affect flowering time (Ruokolainen et al., 2010, 2011). In petunia, ectopic expression of UNSHAVEN (petunia SOC1 homolog) leads to conversion of petals to organs with leaf-like features (Ferrario et al., 2004). Here, we also show that in the rose similar conversion of petals to leaf-like structures is also associated with ectopic expression of RcSOC1. In Arabidopsis SOC1 has been shown to regulate the flower meristem identity gene LEAFY (LFY; Lee et al., 2008). LFY links floral induction and flower organs identity determination

and development via induction of the ABC genes. Whether in the rose *RcSOC1* acts in similar pathway this remains to be addressed.

CONCLUSION

Taken together, our data with those in the literature suggest that the phyllody phenotype in the rose Viridiflora is associated with an up-regulation and ectopic expression of RcSOC1 and of the A-class flower organ identity genes along with the down-regulation of the B, C, and E floral organ identity genes. Therefore, these data here represent a base for future research to deeply understand the molecular mechanisms associated with ABCE organ identity genes, and the origin of type 1 phyllody phenotype in roses.

AUTHOR CONTRIBUTIONS

HY, MB, and KT designed the entire research. HY, HZ, QW, HJ, and XQ performed experiments and helped to analyze data. LG and OR analyzed transcriptome data. SB, JJ, and JW revised the manuscript. HY and MB wrote and edited the manuscript. All authors have read and approved the final manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fpls.2016.00996

FIGURE S1 | Electron-microscopic analysis of sap extracts from young buds at stage 8–10 mm of *R. chinensis* cv. Viridiflora (A) and Old Blush used as negative control (B). The samples were negatively stained and analyzed at a magnification of x100 000. The bar represents 100 nm. No trace of mycoplasma is observed in Viridiflora or Old Blush samples.

FIGURE S2 | No trace of MLO DNA is found in Viridiflora and Old Blush: DNA was prepared from buds at stage 8–10 mm of *R. chinensis* Viridiflora and Old Blush and then used to PCR amplify MLO DNA. M, molecular marker; C, positive control using MLO plasmid; 1,Viridiflora DNA; 2,Old Blush DNA.

TABLE S1 | Primers used for real-time quantitative PCR.

DATA SHEET S1 | The sequences of genes used for qRT-PCR validation.

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

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