Book Chapter

Identification of Candidate Auxin Response Factors (ARFs) Involved in Pomegranate Seed Coat Development

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Abstract

Auxin response factors (ARFs) are transcription factors, regulating the auxin signaling pathways involved in plant development and related processes. In this study, we performed the genome-wide identification and characterization of ARFs in pomegranate and compared them with ARFs from three other species. Seventeen PgrARFs were identified and clustered into four groups, according to their phylogenetic relationship with the remaining 59 ARFs. A recent whole-genome duplication event in pomegranate may have contributed to the expansion and diversification of PgrARFs. Genomic truncation and variant splicing mechanisms contributed to the divergence of PgrARFs, a conclusion that was supported by different exon-intron structures of genes and incomplete conserved domains of PgrARFs in a specific phylogenetic group (group III).
Interestingly, the absence of motifs from certain \textit{PgrARF} genes corresponded to their low transcription levels, which contrasted to the highly expressed \textit{PgrARFs} with intact motifs. Specifically, \textit{PgrARF1} and \textit{PgrARF2} highly expressed in both inner and outer seed coat, and phylogenetically related to \textit{Arabidopsis} orthologs which mediates cell divisions in seed coat. We infer these two \textit{PgrARFs} might involve in seed coat development through cell divisions in response to auxin regulation. These findings provided information on the characteristics and evolutionary relationships of \textit{PgrARFs}, but also shed lights on their potential roles during seed coat development in pomegranate.

**Keywords**

Auxin Response Factor (ARF); Gene Expression; Gene Family Evolution; Phylogenetics; Seed Coat Development

**Introduction**

Pomegranate (\textit{Punica granatum} L.) (2n = 2x = 18) belongs to the family Lythraceae, and is widely cultivated in countries with Mediterranean-like climates around the world, including Tunisia, Turkey, Spain, Egypt, Morocco, the USA, China, India, Argentina, Israel, and South Africa [1,2]. The pomegranate is widely consumed in the form of fruits, juice, wines, and medicines due to its nutritional, medicinal, and ornamental values [3]. In addition, pomegranate differs from other fruit trees in terms of its unique seed structure, with a compressed inner seed coat and an expanded fleshy outer seed coat [4]. The expanded fleshy outer seed coat is the major edible part that largely determines the yield and edible rate. Thus, genetic studies of seed coat development could benefit pomegranate improvement and production.

The development of seed coat is regulated and orchestrated by several transcription factors (TFs) such as ARF, MADS-box, and WRKY [5-8]. Among those TFs, \textit{AtARF2} from auxin response factor (ARF) family was identified involved in cell divisions of seed coat, supported by a mega integument (mnt) mutant allele of \textit{AtARF2}, which induce extra cell divisions and organ growth in
seed coat [8]. Particularly, cell divisions of seed coat in early seed development stage accompanies with the proliferation of endosperm. The process further constrains the cavity of embryo development in later stages and limit the seed size and content [9]. Moreover, auxin mediates ARF expression and activates seed coat development by removing the function of a Polycomb Group (PcG) protein-encoded gene, which epigenetically blocks seed coat development [10]. Therefore, exploration of the gene regulation of ARFs and auxin signaling pathways provide valuable evidence to understand the genetic mechanisms of seed coat development.

ARFs activators and auxin/indole acetic acid (Aux/IAA) repressors are two TFs co-regulate auxin signaling pathway. ARFs target the auxin-response genes by binding to promoters of auxin response DNA elements (AuxREs), which contain the TGTCTC element, to suppress or activate the transcription level of auxin response-related genes. ARFs contain three major domains, namely a conserved N-terminal B3-type DNA-binding domain (DBD), a variable middle region (MR), which acts as an activation or suppression region for ARFs, and a C-terminal dimerization domain (CTD) for protein dimerization [11]. Meanwhile, ARFs are mediated by Aux/IAAs in an auxin concentration-dependent manner. Low auxin concentrations induce the formation of Aux/IAA protein heterodimers, which inhibit ARF activity and repress ARF transcription, whereas higher auxin concentrations derepress ARF activity through degradation of Aux/IAAs from the SCF TIR1/AFB pathway [12,13]. Understanding the regulatory mechanisms of ARFs is key to understanding the auxin signaling pathways.

ARFs have been widely identified in plants along with multiple copies in each species. Twenty-three ARFs have been identified in Arabidopsis (Arabidopsis thaliana), 25 in rice (Oryza sativa), 21 in tomato (Solanum lycopersicum), and 19 in sweet orange (Citrus × sinensis) [14-16]. Nearly doubled numbers of ARFs were identified from banana (Musa acuminata), soybean (Glycine max), and rapeseed (Brassica napus), which is explained by whole-genome duplications (WGD) or polyploidizations [17-19]. Also, conserved domains exhibited
truncation or amino acid substitution in some species. For example, 14, eight, 11 and seven truncations of the DBD domain were identified from barrel medic maize, sweet orange, and tomato [20,21]. The roles of a few ARFs involved in seeds, leaves, flowers, and fruits development have been characterized by functional validation. In *Arabidopsis*, *AtARF7* and *AtARF19* control leaf expansion and lateral root growth [22,23], and *AtARF5* and *AtARF8* are critical elements related to flower formation and fruit development [24-26]. In tomato, *SlARF3* was characterized as a strong candidate for the differentiation of epidermal cells and trichomes [27], whereas *SlARF9* participated in the regulation of cell division during the process of fruit development [28]. However, the role of ARFs in seed coat development has been rarely investigated besides in *Arabidopsis* [8].

In this study, we attempted to explore potential relations between ARFs and seed coat development in pomegranate. Integrated analyses of phylogenetic classification, exon-intron structure, domain structures, of ARFs from pomegranate were conducted and compared with three other species, includes those ARFs from *Arabidopsis*, grape, and eucalyptus. We compared ARFs copy number variation to grape, a species with a recent genome triplication [29]. Also, we identified colinear *PgrARFs* in eucalyptus, a species as the same family of pomegranate, to explore potential ARF lineage specific diversification. Further, *PgrARFs* with intact structure and high expression level in seed coat were chosen and carefully studied, including their temporal expression in different growth stages, correlation between expression and seed coat content increment, and their relations to functional orthologs from *Arabidopsis*. Our study could provide fundamental information of *PgrARFs* characteristics, evolution, and structural variation, also, the candidate ARFs we chosen could provide reference to study seed coat development in pomegranate.
Materials and Methods

Identification of ARFs and Reconstruction of a Phylogenetic Tree

Protein sequences of pomegranate (*P. granatum*), *Arabidopsis* (*A. thaliana*), eucalyptus (*Eucalyptus grandis*) and grape (*Vitis vinifera*), with genome annotations, were downloaded from Phytozome (v11.1, https://genome.jgi.doe.gov) for local sequence blast. Primary genome-wide identification of ARFs from the four species was performed using the hidden Markov model (HMM) by HMMER [30] as described in protocols. Briefly, the domain profiles of B3 (PF06057) and Auxin_resp (PF02362) from the Pfam database (https://pfam.xfam.org/) were searched for, using the ‘HMMsearch’ function with a threshold E-value < 1e-05 against the local database. Furthermore, sequences identified above were confirmed by searching for their conserved domains from the Pfam database and from SMART (http://smart.embl-heidelberg.de), with sequences with incomplete annotation information or missing domains being removed manually. To construct the phylogenetic tree, full-length protein sequences of ARFs selected according to the above criteria were aligned using the multiple sequence alignment tool (MUSCLE) [31]. Conserved domains, namely the DBD, MR and CTD domains, were identified from amino acid sequences, based on alignment positions derived from previous studies [32]. Before the construction of the phylogenetic tree, an optimum amino-acid substitution selection model was selected using model-generator tools [33]. The phylogenetic tree was constructed using the PhyML tool, based on the maximum-likelihood (ML) method (starting tree: BIONJ; bootstrap:100; tree topology search: NNIs). The final tree was visualized using the interactive tree of life (iTOL) [34,35]. Based on the classification of the phylogenetic tree, variations in amino acid sequences in each ARF group were further analyzed by conducting pairwise alignments through BLASTp.
**ARF Gene Structure Analysis and Identification of Conserved Motifs of ARFs**

The gene structure of each ARF, including exon-intron distribution, was displayed, based on published gene annotation information from the four species, using the Gene Structure Display Server (GSDS) [36]. In addition, the number of genes, number of exons, average gene length and average exon length of ARFs in each group in each species were determined. The conserved motifs for each ARF protein sequence were identified by MEME (http://meme-suite.org/tools/meme), with eight as the maximum motif number for comparison. The sequence for each conserved motif identified by the MEME search was confirmed, based on the hits classification by BLAST against the conserved domain database (https://www.ncbi.nlm.nih.gov/Structure/cdd/), the location of each type of motif being represented by different symbol shapes and colors from iTOL. The final conserved motif pattern for each ARF was organized, based on groups and orders from the phylogenetic tree.

**Chromosomal Localization and Synteny Analysis of PgrARFs**

To investigate the gene evolution of ARFs from pomegranate and eucalyptus, another closely related species from the Lythraceae family, comprehensive gene synteny and duplication analyses were conducted between eucalyptus and pomegranate using MCScanX [37] and visualized by Circos [38]. Initially, the genomic location of 17 ARFs from eucalyptus and 17 ARFs from pomegranate were mapped to their respective chromosomal locations, based on annotation from the Phytozome database (https://phytozome.jgi.doe.gov/pz/portal.html) (v11.1). Furthermore, the protein sequences of the 34 ARFs and any ARFs within the 100-kb flanking regions were retrieved for protein sequence alignment. The pairwise alignment for each sequence was conducted by BLASTp, with an E-value > 1e-05 and an identity score > 35% as cut-offs. Synteny analysis was conducted by MCScanX (http://chibba.pgml.uga.edu/mcscan2/) with the following settings: match-score: 50; overlap-window: 5; E-value: 1e-05; max-gaps: 25. The non-synonymous mutation
rate (dN), synonymous mutation rate (dS) and the ratio of non-
synonymous to synonymous substitutions (dN/dS) were
calculated for each collinear ARF pair identified from
pomegranate. Also, duplication analysis from the two species
was conducted by classifying duplication mode using the
duplicate_gene_classifier function in MCScanX, with default
settings based on protein sequences.

**Plant Material for ARF Time-Course Gene Expression
Studies**

Pomegranate cultivar ‘Dabenzi’, a major cultivar of pomegranate
grown in Anhui Province in China, was selected to study the
relationship between seed coat development and ARF gene
expression, by sampling tissues at several time points during
fruit maturation. ‘Dabenzi’ trees were planted in an orchard in
Anhui Province in China (Huaiyuan, 32°95'N, 117°19'E), and
the flowers in full bloom were labelled and classified as 0 days
after full bloom (DAFB) in spring 2019. We sampled nine fruits
at each time point from a 30-year-old ‘Dabenzi’ tree, namely 25,
60, 90, 116 and 145 DAFB. Each fruit sample was dissected
manually and 100 seeds from each fruit were randomly selected
for weighing, with three biological replicates (with three fruits
randomly selected to represent each replicate). For gene
expression analysis, the outer seed coats from three sets of the
replication collected from 25, 66, 90, 116 and 145DAFB were
manually squeezed and frozen in liquid nitrogen, then stored at -
80 °C prior to RNA extraction. Isolation of RNA was conducted
using an OmniPlant RNA Kit (DNase I) (CwBiotech, Taizhou,
China) and cDNA synthesis was carried out by EasyScript One-
Step gDNA Removal and cDNA Synthesis SuperMix (Transgen,
Beijing, China), following the protocols described by the
manufacturers.

**Relative Expression of Candidate PgrARF Genes
During Seed Coat Development**

To identify the candidate PgrARFs potentially involved in seed
coat development, we collected the global transcriptomic data of
pomegranate from published data, which derived from several
plant tissues, namely root, flower, leaf (each organ sample collected at one stage of fruit development), peel (three stages), inner seed coat (three stages) and outer seed coats (each organ samples collected at 50, 95 and 140 days after pollination (DAP) and labelled as Stage1, Stage2 and Stage3) for screening [1]. Only those PgrARFs that exhibited high expression levels in both inner and outer seed coats were selected for further qPCR analysis. The primers used in qPCR for these selected ARFs were designed by Primer Premier 5.0 software (http://www.premierbiosoft.com), following BLAST against the reference genome of pomegranate to prevent amplification of non-specific products. To prevent false positives from qPCR, the cDNA samples from five time points, with three technical replications and three biological replications at each time point, were used, and qPCR was performed using the LightCycler 96 SYBR GREEN I Master (Roche, Indianapolis, IN, USA) in a 20-µL reaction volume, according to the manufacturer’s protocol. The relative expression levels calculated by the cycle threshold (Ct) $2^{-\Delta\Delta Ct}$ method, with a pomegranate actin gene (OWM91407) as an internal control [39]. Results from different samples were compared, using the two-tailed t-test ($\alpha = 0.05$). Further, increment of 100-seed weight and differences in relative transcriptional levels of candidate ARFs between any two representative growth stages were also compared by Pearson correlation analysis. Any PgrARF with a strong correlation may putatively involve in pomegranate seed coat development.

**Potential Divergence of Duplicated PgrARFs**

Combining the identification of duplicated genes with transcriptomics data was used to address the problem of potential effects of gene duplication on gene function divergence. In this study, we compared the correlation of gene expression from different tissues among duplicated PgrARF gene pairs as determined by Pearson’s correlation coefficient. We proposed the use of significant correlation coefficient values to verify the degree of expression difference: $r < 0.3$ signified divergence, $0.3 < r < 0.5$ signified ongoing divergence, and $r > 0.5$ signified non-divergence, based on previous studies [40].
Results
Identification and Phylogeny of ARFs

A total of 76 ARFs were identified from the four species studied, 17 in pomegranate (*P. granatum*), 23 in *Arabidopsis* (*A. thaliana*), 17 in eucalyptus (*E. grandis*), and 19 in grape (*V. vinifera*) (Supplementary S1). Based on the sequence alignment feature, the Jones-Taylor-Thornton (JTT) amino-acid substitution model proved to be the optimum model for further phylogenetic tree reconstruction. According to the phylogenetic tree, the 76 protein sequences were clustered into four groups (Figure 1a). All ARFs were renamed, based on their potential orthologs from *Arabidopsis*, or sequentially, if no corresponding ortholog was found (Supplementary S2). The 76 ARFs were distributed unevenly among the four groups, as group I contained the smallest number of ARFs (nine), whereas group II consisted of the largest number (31). Interestingly, more than half of the ARFs from *Arabidopsis* were clustered in group II whereas the ARFs from pomegranate were distributed more evenly among the four groups, similar to the situation with grape and eucalyptus ARFs (Supplementary S3).

The pairwise alignment between the two ARF protein sequences in each pair of the 76 ARFs revealed some noteworthy features (Figure 2a). Within the same group, we found that ARFs from group IV shared a significant higher average identities (P < 2.2e-16), with those from group I, group II and group III exhibiting lower but similar average identities (P > 0.05). Further, we observed a significantly lower identities from comparison between two different groups which contains sequences from group III, including I vs III, II vs III, and III vs IV (P < 1.8e-16). We also noticed that ARFs from group II exhibited the highest degree of deviation from those from groups I, III and IV, indicating a higher level of diversity of the ARF protein sequences in group II. Additionally, we found some contrasting features of alignment coverage, compared with identities (Figure 2b). Interestingly, genes from group IV exhibited the highest identities but the lowest coverage values, which significantly differed with I vs I, II vs II, and III vs III (P < 1.4e – 15). Comparison of II vs III exhibiting the lowest identities but the
highest coverage values among all the two-group comparisons. These could be attributed to the variations in gene length and gene structure as a result of genomic deletions.

Figure 1: Phylogenetic tree and motif structure of ARFs from four plant species.

a. The phylogenetic tree was reconstructed with the maximum likelihood (ML) method in PhyML and represented in a circular fashion. The four groups are marked in four different colors, namely black, green, orange and red for groups I, II, III and IV, respectively. Branch length was marked on each branch of the tree.

b. Seven motifs corresponded to each ARF, identified as B3, ARF activation/depression or IAA/Aux motifs, are represented by different symbols as shown in the legends.
Figure 2: Pairwise alignment comparison of ARF protein sequences.

a. Comparison of pairwise sequence identity of full-length ARF proteins was conducted in ten groups, namely I vs. I, II vs. II, III vs. III, IV vs. IV, I vs. II, I vs. III, I vs. IV, II vs. III, II vs. IV, and III vs. IV. The boxplot shows the median, interquartile range, and maximum and minimum scores of each data set. Outliers are shown as black circles beyond the whiskers. The level of significance was marked in asterisk.

b. Comparison of pairwise sequence coverage of full-length ARF proteins was conducted in ten groups, namely I vs. I, II vs. II, III vs. III, IV vs. IV, I vs. II, I vs. III, I vs. IV, II vs. III, II vs. IV, and III vs. IV. The boxplot shows the median, interquartile range, and maximum and minimum scores of each data set. Outliers are shown as black circles beyond the whiskers. The level of significance was marked in asterisk.

Diversified ARF Protein Sequences among the Four Groups

To identify the sequence features of ARF-encoded proteins, multiple alignments of the 76 ARF protein sequences were used to identify conserved amino acid residues and the distribution of conserved domains (supplementary S4). Based on the alignments, DNA-binding domain (DBD) from the N-terminal region was identified as residues in between 190 and 310, the middle region of the ARF activation/repression domain at 370–520, and the C-terminal dimerization (CTD) domain at 1350–1450. Among the three domains of the ARF proteins, DBD was the most conserved regions, along with three conserved residues across the 76 sequences at K^{271}, G^{276}, and D^{277}, with these three residues possibly being closely associated with key functions of ARFs.
To further elucidate the variation in motif patterns among the 76 protein sequences, we performed the motif analysis by searching for conserved motif distribution in each ARF amino acid sequence. The MEME search identified eight conserved motifs which could be classified as two specific B3 motifs (namely the B3–1 and B3–2 motifs) from the DBD domain, three ARF activation/suppression-related motifs (namely the ARF-1, ARF-2, and ARF-3 motifs), and two Auxin-related motifs from CTD domains (namely the domain III and domain IV motif). Overall, motif patterns diversified and correlated with the distribution of ARFs based on the phylogenetic tree (Figure 1b). We found that all ARFs from group III lacked the ARF-1 motif and the domain IV motif. PrgARF17, EucARF17, AtARF14 and VvARF5 also lacked the domain IV motif, compared with the rest of the group III members. Interestingly, we observed a longer branch distribution for all group III ARF proteins from the phylogenetic tree, as well as ARFs bearing longer branches with the most severe motif loss. The different lengths of tree branches indicated the recent emergence of ARFs from duplications. We hypothesized that the more recent evolution of these ARF genes from group III could be associated with motif loss from the DBD domain and the CTD domain. Additionally, losses of the ARF-1, ARF-3, domain III, and domain IV motifs from PgrARF14, PgrARF15 and EucARF15 were observed, a situation which was more severe than that occurring with encoded proteins of potential orthologs from Arabidopsis and grape (AtARF3, AtARF4 and VvARF3). This might be explained by lineage-specific variation of ARF genes from pomegranate and eucalyptus, both members of the Lythraceae family. The other ARFs suffering motif loss were EucARF13, AtARF13 and AtARF13 in group II and VvARF3 and AtARF3 in group I, which were less closely related to the respective phylogenetic patterns.

Gene Truncation and Gene Structure Variations of ARFs

To explore the putative causes of protein sequence divergence, we compared the structural gene annotation from the four species. The complete gene annotation information, which
included untranslated regions (UTRs), exon sites and intron sites, presented a diversified genomic pattern. Among the 76 genomic ARF sequences, we found that the genomic sequences of ARFs from eucalyptus were longer than those from the other three species, due mostly to their longer introns (Figure 3). Most ARFs from the same group contained similar numbers of exons among the four species but different exon numbers were identified among the four phylogenetic groups. Surprisingly, significantly shorter (25–33% length) genes and substantially longer (2- to 3-times length) exons were observed in genes from group III, compared with the genes from the other three groups, with fewer exons on average per gene (Supplementary S3).

![Figure 3: Exon–intron structure of 76 ARFs full-length genomic sequences.](image)
The intron/exon structure of ARFs was visualized by gene structure display server (GSDS). Yellow rectangles represent exons and dark lines represent introns, with UTRs (untranslated regions) being marked with shaded rectangles.

**Duplication and Evolution of ARFs from Eucalyptus and Pomegranate**

Identification of collinear gene pairs and homologous genes enabled the identification of duplicated gene pairs. In the present study, pairwise comparisons between ARFs and genes from each 100-kb flanking region from the pomegranate and eucalyptus genomes revealed 32 collinear blocks from 22 genomic location
combinations (E-value: 1e-05). The 32 collinear blocks were classified between pomegranate and eucalyptus, with six blocks within the pomegranate genome, but only two blocks within the eucalyptus genome (Table 1). Collinear regions between pomegranate and eucalyptus were mainly distributed on two eucalyptus chromosomes and one pomegranate chromosome (Euc04, Euc11 and Pgr09), whereas collinear regions within the same species were distributed on Euc11 and Pgr09 (Figure 4). Analysis of duplication type revealed that whole-genome duplication (WGD) contributed to nearly all duplicated ARFs from pomegranate, similar to the finding in eucalyptus (Supplementary S5). Collinear gene-pair distribution and duplication types revealed that 12 collinear gene-pairs were categorized in group IV (Table 2). We speculated that PgrARFs from group IV played a substantial role in ARF expansion in pomegranate, caused by the recent WGD in this species.
Table 1: Summary of collinear ARFs from pomegranate.

| Gene 1      | Gene 2      | Class   | Gene expression | Correlation | dN   | dS   | dN/dS |
|-------------|-------------|---------|-----------------|-------------|------|------|-------|
| *PgrARF11*  | *PgrARF1*   | II–II   | non–divergent   | 0.714       | 0.47 | 2.1  | 0.22  |
| *PgrARF9*   | *PgrARF1*   | II–II   | non–divergent   | 0.512       | 0.51 | 2.52 | 0.2   |
| *PgrARF17*  | *PgrARF16*  | III–II  | non–divergent   | 0.561       | 0.67 | 1.36 | 0.49  |
| *PgrARF19*  | *PgrARF6*   | IV–IV   | divergent       | 0.207       | 0.54 | 1.66 | 0.32  |
| *PgrARF19*  | *PgrARF8*   | IV–IV   | divergent       | 0.436       | 0.53 | 2.57 | 0.2   |
| *PgrARF7*   | *PgrARF5*   | IV–IV   | divergent       | 0.602       | 0.55 | 2.25 | 0.25  |
| *PgrARF7*   | *PgrARF6*   | IV–IV   | divergent       | -0.139      | 0.53 | 2.11 | 0.25  |
| *PgrARF8*   | *PgrARF12*  | IV–IV   | NA              | NA          | 0.19 | 1.03 | 0.18  |
| *PgrARF13*  | *PgrARF10*  | II–III  | divergent       | 0.332       | 0.98 | 2.7  | 0.36  |
| *PgrARF11*  | *PgrARF12*  | II–IV   | NA              | NA          | 0.68 | 1.67 | 0.41  |
| *PgrARF9*   | *PgrARF6*   | II–IV   | divergent       | 0.142       | 0.71 | 1.64 | 0.43  |
| *PgrARF17*  | *PgrARF9*   | III–II  | divergent       | 0.332       | 1.04 | 1.81 | 0.57  |
| *PgrARF5*   | *PgrARF1*   | IV–II   | divergent       | 0.351       | 0.76 | 1.48 | 0.51  |
| *PgrARF7*   | *PgrARF11*  | IV–II   | divergent       | 0.201       | 0.77 | 1.25 | 0.61  |
| *PgrARF8*   | *PgrARF1*   | IV–II   | divergent       | 0.429       | 0.75 | 1.21 | 0.62  |
| *PgrARF5*   | *PgrARF10*  | IV–III  | divergent       | 0.061       | 0.95 | 1.41 | 0.67  |
| *PgrARF7*   | *PgrARF16*  | IV–III  | divergent       | 0.063       | 1.13 | 1.1  | 1.02  |
Figure 4: Collinearity of ARFs between eucalyptus and pomegranate. Genes from 100-kb flanking genomic regions of 17 PgrARFs and 17 EucARFs were mapped on chromosomes of pomegranate and eucalyptus, based on gene annotation. The red lines connect inter-specific collinear gene pairs, whereas black lines connect collinear genes within each species.

Table 2: Summary of collinear genes between pomegranate and eucalyptus.

| Classification       | Location      | Block no. | Collinear gene pairs |
|----------------------|---------------|-----------|----------------------|
| Pomegranate–eucalyptus| Euc02-Pgr01   | 2         | 24                   |
|                      | Euc02-Pgr05   | 1         | 7                    |
|                      | Euc02-Pgr09   | 1         | 6                    |
|                      | Euc03-Pgr02   | 1         | 8                    |
|                      | Euc04-Pgr03   | 1         | 11                   |
|                      | Euc04-Pgr05   | 1         | 6                    |
|                      | Euc04-Pgr09   | 2         | 12                   |
Expression Characteristics of *PgrARF*s in different Tissues of Pomegranate

Transcriptome profiling from a variety of plant tissues may help to identify tissue-specific genes which, in turn, helps to identify candidate genes for specific biological processes. In this study, tissue-specific transcriptomic data, including root, flower, leaf, peel, and seed coat (inner and outer seed coat), revealed variations in expression patterns among the four *PgrARF* groups. We identified two highly expressed ARFs with a broad spectrum of expression during vegetative growth and reproductive developmental stages, namely *PgrARF1* and *PgrARF2* from group II (Figure 5). Expression of another three ARFs (*PgrARF5*, *PgrARF7* and *PgrARF19*) was up-regulated in most tissues except for down-regulation in the outer seed coat. However, nearly all the genes from groups I and III were transcribed at extremely low levels. The remaining genes exhibited partially tissue-specific expression. For instance, *PgrARF9*, *PgrARF11*, *PgrARF6*, and *PgrARF8* were highly expressed in root, leaf and peel. We also compared the...
transcriptome pattern of collinear genes to detect any differences in gene expression. Among 17 pairs of collinear genes from pomegranate, two pairs of collinear genes from group II and one pair from group III revealed non-divergent patterns of expression. However, expression of five gene pairs from group IV was less closely correlated, despite the similar structures shared over much of the genes. The nine gene pairs showing different expression patterns might be explained by structural variations among groups. In all, a majority of collinear PgrARFs were differentially expressed, potentially due to structural variation among different groups or some other unknown factors.

Figure 5: Transcriptome profiling of PgrARFs from different plant tissues. Relative expression level of 16 PgrARFs from root, flower, leaf (all sampled at one stage), peel (three stages) inner seed coat (three stages) and outer seed coat.
(three stages) of pomegranate are presented as Reads Per Kilobase of transcript per Million mapped reads (FPKM) values, based on phylogenetic grouping. Three stages for peel, inner seed coat and outer seed coat were 50, 95 and 140 days after pollution (DAP).

**Identification of ARF Candidates Involved in Seed Coat Development**

On the basis of transcriptome profiling of *PgrARFs*, we selected ARF candidates that were potentially involved in seed coat development, and performed qPCR to confirm the results, then carried out statistical analysis of the relationship between 100-seed weight and mRNA abundance level. Transcriptome data revealed that *PgrARF1* and *PgrARF2* were highly expressed in both the inner and outer seed coats, suggesting that these patterns might be related to their involvement in seed coat development. *PgrARF5*, *PgrARF7* and *PgrARF19*, which were highly expressed in the inner seed coat, were expressed at a lower rate in the outer seed coat during fruit development. It is suggested that these three genes might participate in the early stages of seed coat development while being less involved in the later stages.

Similar expression patterns of *PgrARF1* and *PgrARF2* were detected by transcriptomics and by qPCR. The expression of these two candidate *PgrARFs* peaked at 25 DAFB then fell to their lowest level at 55 DAFB, followed by a slight increase or decrease at the third time point (Figure 6a). Regression analysis revealed a significant linear relationship between 100-seed weight increment and mRNA transcription level differences in each two stage of each candidate gene, as identified by significant positive correlations for both *PgrARF1* (*r* = 0.969, *P* < 0.05) and *PgrARF2* (*r* = 0.967, *P* < 0.05) (Figure 6b and 6c). Hence, we propose that *PgrARF1* and *PgrARF2* might be involved in the outer seed coat development of pomegranate.
Figure 6: The relationship between PgrARF gene expression level and pomegranate seed weight.

a. Relative gene expression level of PgrARF1 and PgrARF2 genes for outer seed coats at 25, 55, 90, 116, 145 days after full bloom (DAFB) with three technical replicates and three biological replicates.

b. 100-seed pomegranate seed weight (g) from three biological replicates across five stages of fruit development (days after full bloom, DAFB).

c. Linear regression analysis of differences in 100-seed weight from each of two stages with respect to change in rate of gene expression from four sampling time-points.

Discussion

Gene Duplication as Trigger of PgrARF Expansion

Gene duplication is one of the major driving forces for plant genome evolution, and also impacts the expansion of and functional variation within gene families. Global identification of ARFs from multiple flowering plant lineages has identified the co-occurrence of ARF expansion and whole-genome duplication (WGD) [17,19,41,42]. It is plausible to propose gene duplication
as a trigger of PgrARF expansion. In the present study, we identified that WGD contributed to ARF gene duplication in pomegranate, which mostly occurred in groups II, III and IV, matching the distribution pattern from previous studies [43]. Interestingly, duplicated (collinear) ARFs from group IV were functionally divergent, whereas three pairs of collinear genes within groups II and III exhibited correlated expression patterns (Table 2). Similar examples of functional redundancy of arf6/arf8 mutants from Arabidopsis and duplicated AtARF3 and AtARF4 with functional divergence had been identified in previous studies [43,44]. In all, we proposed that gene duplication is closely related to gene redundancy or functional divergence during the evolution of the ARF gene family in pomegranate.

Genomic Truncation and Splicing Variation Contributed to Diversified PgrARFs

Modification at the post-transcriptional level is another major force potentially contributing to gene diversification. Alternative splicing of the ARF gene family has been identified from numerous land plants [21,43]. For instance, different functional roles were identified from two isoforms of Arabidopsis ARF4 (ARF4 and ΔARF4) during carpel development [43]. In another species in the Lythraceae family, alternative transcripts have been identified from 10 out of 17 ARFs from eucalyptus [45]. On the other hand, no alternative transcripts were found among the 17 PgrARFs, based on an exhaustive search of putative transcripts from annotation, although eucalyptus shared quite a few collinear ARFs with pomegranate, and highly similar ARF protein sequences were identified from inter-specific pairwise alignment (Table 1). Consequently, the origins of alternative splicing of ARFs might be lineage specific, and hence, less relevant to the evolution of species. In addition, reduced exon numbers, increased gene lengths and truncated genomic lengths were identified from group III (Supplementary S3). It is plausible to reason that, besides the missing residues or motifs from ARF protein sequences, some functional divergence might have occurred due to these structural gene variations. However, this hypothesis, regarding the structural variants which, in
pomegranate, appeared exclusively in group III, indicates that the variation in structure was not confined to pomegranate but was also found in the other three species in our study. We hypothesized that this specific phenomenon during the subfamily evolution might be related to the evolution of the splicing process. It would be tempting in future studies to explore the potential mechanisms involved in achieving increased exon length.

**Conserved Domains as Evidence of Intact PgrARF Function**

Conserved amino acid residues or motifs play substantial roles in maintaining intact domain functions, which is closely related to gene expression and to gene regulation. The role of each domain from the ARF protein was characterized in numbers of earlier studies [12,46,47]. The DBD achieves binding to the DNA target site in an auxin-independent manner. On the other hand, the MR in the ARF domain either activates (the Q-rich ARF domain) or represses (the S-rich ARF domain) transcription level, whereas the CTD regulates the auxin response pathway by interaction with Aux/IAAs. Numerous truncated proteins caused by motif losses were identified from several of the species under investigation, whereas such variations in domains showed a close relationship between gene expression and sequence conservation. For example, significantly reduced transcription levels in root, leaf, shoot, cotyledon and flower were exhibited by nine MtARF genes from *Medicago*, all of which exhibited the missing CTD or the partially truncated ARF domain [20]. A similar expression pattern in citrus revealed a lower relative mRNA abundance from *CiARF3* and *CiARF17*, which could be related to the missing CTD domain [14]. In our structural analysis and expression profiling (Figure 1b and Figure 5), we found that ARFs with incomplete ARF and CTD domains, lacking ARF-1, domain III and domain IV motifs from certain PgrARFs from both group I and group III, were associated with low transcription rates from a number of plant tissues.

In addition, several residues have been reported to play a substantial role in ARF transcription [48]. In *Arabidopsis*,
variations in the H170 residue reduced the binding of AtARF5 to the corresponding AuxREs, as did mutations identified from the P218, R215, T227, and S230 codons. Interrupted dimerization was identified as results of G279, A282 and A287 substitution identified in the ARF domain. In our alignments of the 76 sequences, residues among those sites were carefully scanned and we found strong associations between conserved codon patterns and transcription expression patterns (Supplementary Figure 1), with substitution from H to G at the H170 residue position or substitution of T to A at the 202 residue position resulting in reduced expression of ARFs from group III. Since motifs and some amino acid residues from the ARF (MR) domain play a substantial role in binding target DNA, it might be plausible to postulate that down-regulation of expression of ARFs was associated with truncated conserved domains or even substitution of an amino acid residue in such domains.

**Diversified PgrARFs and Potential Candidate Genes for involvement in Seed Coat Development**

ARFs regulate numerous auxin-related processes at different plant developmental stages, as evident from gene expression patterns identified from previous studies [16,17,21]. In the current study, two pairs of highly expressed ARFs (PgrARF1 and PgrARF2, and PgrARF7 and PgrARF19) exhibited broad-spectrum expression in several different plant tissues, and shared a similar expression pattern to those of EucARF1 and EucARF2, and EucARF17 and EucARF9 from eucalyptus [45]. On the other hand, tissue-specific expression patterns were identified from the four corresponding ARFs in other species, including Arabidopsis, tomato and citrus [14,21,49]. In situations where functional analysis of ARFs has been studied, functional mutant analysis in Arabidopsis provided invaluable resources for exploring ARF gene functions in other species. For orthologs from Arabidopsis, the loss-of-function double mutant revealed overlapping functions of AtARF9 and AtARF17, which participate in the key step of lateral root formation and root development [49]. ARF1 and ARF2 regulate leaf senescence and floral organ abscission, while sharing partial functional redundancy [50].
Interestingly, we found that the expression patterns of *PgrARF1* and *PgrARF2* were also very similar, as were those of *PgrARF7* and *PgrARF19*. The similarity of expression pattern might be related to functional redundancy as occurred in their respective orthologs from *Arabidopsis*. Combined with the roles of *ARF1* and *ARF2* in cell division and cellulose synthesis, specifically the role of mediating cell division in seed coat development of *AtARF2* \[8,51]\], and the similar expression patterns from a number of tissues between two genes, we proposed that *PgrARF1* and *PgrARF2* are two structurally intact candidates that participate in cell division of seed coat during seed coat development.

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**Supplementary Material**

Supplementary material can be accessed online at: https://videleaf.com/wp-content/uploads/2020/12/PAPLS2ED-20-05_Supplementary-Material.zip