Graft-induced Changes in MicroRNA Expression Patterns in Citrus Leaf Petioles

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Abstract: Grafting is an important, widely used plant propagation technique but its physiological effects are as yet insufficiently understood. Recent studies indicate that movement of proteins and small RNAs through the graft union might be involved. MicroRNAs are known to play a significant role in regulation of higher plants’ developmental and metabolic traits. Extending this logic, we hypothesize that changes in activity of specific microRNAs are one of the mechanisms involved in physiological effects of grafting. The objective of the present study was to test this hypothesis. We determined the expression of a broad range of microRNAs in Citrus leaf petioles, as affected by grafting. Four stock/scion combinations (‘Merav’ mandarin and ‘Star Ruby’ grapefruit scions X ‘Troyer’ citrange and ‘Volkamer’ lemon rootstocks), rootstock auto-grafts and plants of the variety used as rootstock (= non-grafted) were examined. Grafting caused a dramatic reduction in the expression of the major microRNAs, miR156 (and miR157), which appear to be associated with reduction of juvenility in perennial woody plants. This effect was strongest in hetero-grafts but evident also in auto-grafts. Expression of miR894 also declined upon grafting. Differences in the expression of miR397 were found among grafted scion cultivars, while in non-grafted rootstocks expression of miR397 was barely detectable. Bioinformatic analysis confirmed the presence of miR397 in the citrus genome, validated its sequence and demonstrated its ability to form a stem loop. The differences in miR397 expression might be related to specific copper and other micronutrient requirements of citrus stock-scion combinations. Thus, our results support the hypothesis, indicating the involvement of specific microRNAs in engendering physiological effects of grafting in Citrus. The precise, underlying mechanism needs to be elucidated.

Keywords: Citrus, grafting effects, juvenility, rootstock, scion.

INTRODUCTION

Grafting of plants has been practiced throughout the ages [1] and is used widely even today in culture of fruit trees, ornamentals [e.g. roses] and as of recently, vegetables [2]. A major advantage of propagation by grafting is the marked shortening of the juvenile phase, which in tree seedlings may last several years or, in forest trees, up to several decades [1, 3]. The interactions between rootstocks and scions are diverse, complex and as yet not fully understood. While anatomical aspects, supply of water, mineral nutrients and transport of plant hormones have been implicated, recent work demonstrated cross-graft transfer of specific proteins [4] and RNA-induced gene silencing [5,6]. Thus, molecular mechanisms are currently assumed to be involved in physiological effects of grafting and in mediation of stock/scion interactions. MicroRNAs are currently known to be involved in regulation of a large number of developmental and metabolic plant traits [7-9]. Evidence for graft transmission of specific microRNAs has been accumulating in recent years, in various plant species [10-13].

Although grafted plants have been used extensively as experimental systems in small RNA research, a general role for microRNAs in physiological effects of grafting has not been hypothesized, to the best of our knowledge. The objective of the present study was to test this hypothesis in citrus and search for graft-related differences in expression of specific microRNAs. We used young, potted citrus trees as our experimental system, examining several grafted stock/scion combinations, rootstock auto-grafts and non-grafted controls Fig. (1). Petioles of fully expanded leaves were used as a source of RNA, the rationale being that petioles consist predominantly of vascular tissues in which transportable compounds may be abundant. Unlike studies that concentrated on a small number of presumably relevant microRNAs, we adopted a screening strategy, examining the expression of a broad spectrum of microRNAs.

Here we show that grafting induced changes in the expression pattern of several ubiquitous, physiologically significant microRNA species in citrus leaf petioles. Differences in the expression pattern of smaller, specific microRNAs among citrus’ stock/scion combinations were also demonstrated, thereby supporting our hypothesis.
MATERIAL AND METHODS

Plant Material and RNA Isolation

Grafted and non-grafted, container grown, 2 year old citrus trees were obtained from ‘Bney Dror’ Nurseries, Israel, courtesy of Mr. Shuli Rosen. Grafting was performed during spring of the second year; the graft union was smooth and healthy. Four stock/scion hetero-graft combinations (2 scions X 2 rootstocks), 2 rootstock auto-grafts and 2 non-grafted controls were tested Fig. (1). The citrus cultivars used (and their abbreviations) are listed in Table 1. Individual seedlings served as replicates.

Petioles of fully expanded leaves were used for RNA analysis. Leaf petioles were collected during mid-summer (July-August) from actively growing trees by late morning (9 – 10 am), when leaf activity was already on. The effect of wounding was also examined. The wounding treatment consisted of scratching 20 cm along stems of rootstock seedlings 48h before the sampling of leaf petioles. Fully expanded leaves were plucked using sterile gloves, the petioles removed and instantaneously frozen in liquid nitrogen. Total RNAs were extracted from petiole tissue using a ‘Norgen’ biotek corporation Plant/Fungal Total RNA Isolation Kit 50R 25800.

MicroRNA Microarray Assay

Microarray assay was performed using a service provider (LC Sciences, Houston TX, USA) according to their protocol and statistical outlay. The custom μprafe® microfluidic chip contained plant miRNAs of release version 13.0 (http://microrna.sanger.ac.uk).

Bioinformatics

Database searches were run at HarvEST (http://138.23.191.145/blast/index.html) against version 1.25 of HarvEST Citrus and the JGI Citrus genome database (citrus sinensis), using the default parameters with the exception of E-value, which was raised to 1000 due to the short input sequence length. Searches were also performed at NCBI blast (http://blast.ncbi.nlm.nih.gov/), against both NR and est_other databases, with default parameters, with exceptions of E-value of 1000, the blastn algorithm, and organism limited to citrus (taxid:2706). Alignments were performed with ClustalW 2.012 [14] and RNA secondary structure was predicted with both Mfold [15] and the Vienna package [16].

TaqMan miRNA Assay

For validation of miRNA microarray results, TaqMan miRNA assays (Applied Biosystems) were used according to the manufacturer’s protocol [17]. Levels of the citrus miR397, a perfect match to miR397a, were normalized to the miR164a control; miR164 a was used as a normalizing gene since members of this family were highly expressed, with least variation among treatments (Suppl. Table 1). All qRT-PCR reactions were performed on ABI7300 machine. Results are presented as means and standard deviation of two biological replicates.

RESULTS

The miRNAs identified through microarray in the citrus cultivars examined are listed in Suppl. Table 1. A total of 95 miRNAs belonging to 35 families were detected. The expression rates of nine ubiquitous miRNAs in petioles of the two scion cultivars, ‘Merav’ mandarin and ‘Star Ruby’ grapefruit, grafted on ‘Volkamer’ and ‘Troyer’ rootstocks and the non-grafted controls were compared in the experiment presented in Fig. (2). A significant difference between the grafted and non-grafted petioles appeared with miR156 and miR157. The expression of these miRs was much higher in the non-grafted rootstocks than in either grafted scion cultivar, in ‘Volkamer’ more than in ‘Troyer’. A similar, although less striking trend appeared also with miR894, which expressed more strongly in the non-grafted rootstocks than in the grafted scions Fig. (2).

Table 1. Citrus cultivars used in the present study.

<table>
<thead>
<tr>
<th>Horticultural use</th>
<th>Scientific name</th>
<th>Abbreviation</th>
<th>Cultivar</th>
</tr>
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<tbody>
<tr>
<td>scion</td>
<td>Citrus reticulata Blanco</td>
<td>Me</td>
<td>‘Merav’ mandarin</td>
</tr>
<tr>
<td>scion</td>
<td>Citrus paradisi Macf.</td>
<td>SR</td>
<td>‘Star Ruby’ grapefruit</td>
</tr>
<tr>
<td>rootstock</td>
<td>Citrus volkameriana</td>
<td>Vo</td>
<td>‘Volkamer’ lemon</td>
</tr>
<tr>
<td>rootstock</td>
<td>Poncirus trifoliata X Citrus sinensis</td>
<td>Tr</td>
<td>‘Troyer’ citrange</td>
</tr>
</tbody>
</table>
expression of miRs 156, 157, and 894 appeared also in a separate comparison of the non-grafted rootstocks (data not shown). With other miRs (miRs 159, 164, 166, 167, 319, 396) the expression rates of the grafted and non-grafted were similar and did not reveal striking differences. Although the expression of miR172 in our leaf petiole system was quite low, the activity was still somewhat higher in the grafted scions than in the non-grafted plants (data not shown).

In another experiment we compared the microRNA expression patterns in petioles of non-grafted ‘Volkamer’ and ‘Troyer’ plants with auto-grafts of the same rootstock cultivars; the effect of wounding was also tested in this experiment Fig. (3). The expression of miR156 and miR157 was considerably higher in the non-grafted plants than in the corresponding auto-grafts. Wounding of the non-grafted plants did not seem to affect the expression of miRs 156, 157. Neither auto-grafts nor wounding seemed to influence the expression of other major miRs Fig. (3), or any other miRs.

The comparison between the 4 hetero-graft, stock/scion combinations was intended as a probe for apparent rootstock effects. The microarray analysis clusters (not shown) revealed 3 miRNA families which were expressed stronger in both ‘Merav’ and ‘Star Ruby’ scions when grafted on ‘Troyer’, than when grafted on ‘Volkamer’ rootstock; miR397, miR398, and miR408. Among those, miR397 provided the most striking differences between grafts on ‘Volkamer’ and grafts on ‘Troyer’. Fig. (4) demonstrates the differences obtained in the microarray analysis with 5 members of the miR397 family; the expression on ‘Troyer’ was consistently higher than on ‘Volkamer’ rootstock. However, the difference in miR397 expression between grafts on ‘Troyer’ and ‘Volkamer’ was more prominent with ‘Star Ruby’ than with ‘Merav’ scion Fig. (4). Not of lesser significance is the finding that members of the miR397
family, which showed consistent differences between the scions grafted on ‘Volkamer’ and ‘Troyer’ rootstocks, did not show such differences between the non-grafted control plants Fig. (4). In fact, the miR397 family was either weakly expressed (ath miR397a, b) or completely undetectable (bna miR397a, bna miR397b, osa miR397a, b, ptc miR397a, c, sly miR397) in the non-grafted plants (Suppl. Table 1, and Fig. (4)). The lack of detectable miR397 expression in the petioles of non-grafted ‘Volkamer’ and ‘Troyer’ plants was reconfirmed in a separate experiment (data not shown).

Thus, miR397 appeared to be the best candidate for further validation through qRT PCR. However, as its presence in citrus was based previously on prediction from Arabidopsis [18], we performed bioinformatic analyses to confirm its presence in citrus and validate the exact sequence. Sequence based searches with the arabidopsis miR397a in citrus ESTs and mRNAs in NCBI and ESTs at HarvEST did not yield any results. However, when we searched against the JGI Citrus genome database, two perfect matches were found (BUZZ68439.b1, and BUZZ65818.g1). The two hits had a perfect overlap of 466 bp, including the miR397a sequence. The putative miR sequence was a perfect match to miR397a from arabidopsis, bdi, bna (which has an additional terminal T), osa, ptc, and vvi Fig. (5). We tested the ability of the genomic DNA in this region to form a stem-loop, required for the processing of miR. We took both 100bp up and downstream of the miR and the region equivalent to the stem-loop of arabidopsis 397a (110 bp total) and both formed stem-loops, with the miR in a stem region Fig. (6). miR397 was subsequently sequenced independently by Song et al. [9], confirming what we had found in our analyses.

The results of the qRT PCR assays confirmed the stronger activity of miR397 in both scions on ‘Troyer’ than on ‘Volkamer’ rootstock Figs. (7A and 7B). In the qRT PCR as well, the quantitative difference between the rootstocks was more prominent with ‘Star Ruby’ than with ‘Merav’ scion.

DISCUSSION

The growing interest in the role of small RNAs has resulted in a large number of publications that describe the microRNA population of various crop plants, including citrus [18-20]. Yet, studies relating to the physiological roles
Graft-induced Changes in MicroRNA Expression Patterns

The Open Plant Science Journal, 2013, Volume 7 21

of microRNAs are still scarce, and nothing of this kind has been reported so far for Citrus.

Fig. (6). Predicted folding of the citrus miR397a. The sequence equivalent to the published Arabidopsis stem-loop was taken as input to RNAfold (Gruber et al.) [16]. The minimum free energy of this conformation is -43.30 kcal/mol. The drawing is colored according to the base-pairing potential, with red meaning almost definitely in a pair. Note that the region of the stem is very significantly paired.

Grafting has been used extensively in experimental plant research, in classical studies of floral induction [21] as well as in recent investigations [22]. Grafting has played an important role in studies of gene silencing [5], mainly as a probe for the long distance movement of various kinds of small RNA [12]. Graft transmission of the tuberization inducer miR172 has been demonstrated in potato [11] and in Nicotiana benthamiana [23]. The role of graft-transmissible MicroRNA signaling has been demonstrated with P [24], S, Cu and Fe mineral starvation [10].

In the present study we focused on the possible involvement of microRNAs in physiological effects of grafting in young citrus trees. We had initially intended to use shoot sap, but the difficulties in obtaining substantial amounts of sap from citrus shoots [25] turned us to the use of mature, non-growing leaf petioles which consist mostly of vascular elements. Since the xylem appears to be devoid of RNAs [26] petiole extracts are presumably enriched with phloem sap RNA components. The microRNA population of phloem tissue and phloem sap has been reported for Cucumis, Lupinus [27] Brassica [26], and Malus [13].

In the study of grafting physiology it seems important to distinguish between primary, major effects, which concern any grafted plant, and secondary, minor effects which reveal differences between specific rootstock/scion combinations. Most horticultural studies have concentrated up to now on the latter, secondary effects, and paid little attention to the primary effects. In the present experimental study both types of effects became apparent.

In the discussion of our results it seems appropriate to begin with the data of the primary, major microRNAs Figs. (2 and 3). The most intriguing result is the reduced expression of miR156 and miR157 in petioles of all grafted combinations as compared with their respective non-grafted control plants. Extreme reduction was obtained in grafted scion cultivars Fig. (2) and considerable reduction appeared also in the auto-grafts of both rootstock cultivars Fig. (3). The significance of these findings becomes apparent in view of recent work on the vegetative phase change in annuals and trees [28, 29]. According to Wang et al. [29] the juvenile-to-adult transition in tree species involves a marked decline in the activity of miR156. Reduction of juvenility is one of the most significant effects of grafting in trees [1,3]. Thus, the observed reduction of miR156, and the closely related miR157 [30] activities in grafted citrus Figs. (2 and 3) fits

Fig. (7). Comparison of qRT-PCR activity of miR397 in petioles of each scion (Me, SR) on the 2 rootstocks (Vo, Tr); miR164 was used as the normalizing gene (see Methods). Averages from 2 replicates, each consisting of 3 technical replicates ± SD.
very well into the presumed role of these micro RNAs in the juvenile-to-adult transition in trees. It is conceivable that auto-grafts reveal a more moderate reduction of juvenility than hetero-grafts (that represent an interaction between two plant genomes); which explains why the reduction in the expression of miRs 156, 157 was more pronounced in hetero-grafts (Fig. 2) than in auto-grafts (Fig. 3). According to Wang et al. [29], the acquisition of the adult phase also involves a gradual increase in the expression of miR172. Slightly higher expression of miR172 in petioles of the grafted scions, as compared with non-grafted control plants, has been obtained in the present study as well (data not shown).

The search for differential expression of a specific micro RNAs between distinct rootstock/scion combinations has led to the identification of miR397, which expressed more strongly, in both scions, when grafted on ‘Troyer’ than when grafted on ‘Volkamer’ Figs. (4 and 7). The presence of miR397 in citrus has recently been demonstrated by Song et al. [19], and its targeting of IRX12 was confirmed. Expression of the miR397 family is induced by various stress treatments [31]. Members of miR397 target copper binding to proteins in Arabidopsis [32] and play a role in plant heavy metal homeostasis [33]. Growth under copper deficiency is known to induce a number of physiological responses, including the expression of miRs 397, 398, 408 [34], which were also differentially expressed in the present study. In Brassica napus copper deficiency led to a more than four-fold increase of miR397 expression in phloem sap [10]. Involvement of microRNA signaling in additional mineral nutrition elements has been demonstrated in Brassica [10, 24]. The observed differential expression of miR397 in citrus’ stock/scion combinations Figs. (4 and 7) might explain subtle differences in copper and other micronutrient requirements between citrus rootstocks and graft combinations [35-37].

All in all, the results of the present study lend support the hypothesis that microRNA expression plays a role in the mediation of both primary and secondary effects of grafting. Extension of these studies to other grafted plant systems and elucidation of the precise physiological mechanisms involved requires further research.

CONFLICT OF INTEREST

The authors confirm that this article content has no conflicts of interest.

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SUPPLEMENTARY MATERIAL

Supplementary material is available on the publishers Web site along with the published article.

REFERENCES

Graft-induced Changes in MicroRNA Expression Patterns

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