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**Permalink** https://escholarship.org/uc/item/73n5t15x

**Journal** Journal of Experimental Botany, 65(12)

## ISSN

0022-0957

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## **Publication Date**

2014-07-01

## DOI

10.1093/jxb/eru148

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**RESEARCH PAPER** 

Journal of Experimental Botany www.jxb.oxfordjournals.org

# Fruit load induces changes in global gene expression and in abscisic acid (ABA) and indole acetic acid (IAA) homeostasis in citrus buds

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Received 6 November 2013; Revised 10 March 2014; Accepted 11 March 2014

#### Abstract

Many fruit trees undergo cycles of heavy fruit load (ON-Crop) in one year, followed by low fruit load (OFF-Crop) the following year, a phenomenon known as alternate bearing (AB). The mechanism by which fruit load affects flowering induction during the following year (return bloom) is still unclear. Although not proven, it is commonly accepted that the fruit or an organ which senses fruit presence generates an inhibitory signal that moves into the bud and inhibits apical meristem transition. Indeed, fruit removal from ON-Crop trees (de-fruiting) induces return bloom. Identification of regulatory or metabolic processes modified in the bud in association with altered fruit load might shed light on the nature of the AB signalling process. The bud transcriptome of de-fruited citrus trees was compared with those of ON-and OFF-Crop trees. Fruit removal resulted in relatively rapid changes in global gene expression, including induction of photosynthetic genes and proteins. Altered regulatory mechanisms included abscisic acid (ABA) metabolism and auxin polar transport. Genes of ABA biosynthesis were induced; however, hormone analyses showed that the ABA level was reduced in OFF-Crop buds and in buds shortly following fruit removal. Additionally, genes associated with Ca<sup>2+</sup>-dependent auxin polar transport were remarkably induced in buds of OFF-Crop and de-fruited trees. Hormone analyses showed that auxin levels were reduced in these buds as compared with ON-Crop buds. In view of the auxin transport autoinhibition theory, the possibility that auxin distribution plays a role in determining bud fate is discussed.

Key words: Abscisic acid, alternate bearing, auxin, bud, citrus, flowering, fruit load.

#### Introduction

Fruit trees exhibit two major multiannual reproductive strategies (Goldschmidt, 2013). In the first, the amount of fruit produced allows a sufficient amount of vegetative growth to support production of an ample number of flowers during the following year (return bloom). Such trees, including fig and some orange and grapefruit cultivars, are defined as regular bearers. They are characterized by a relatively stable multiannual yield, and usually possess efficient mechanism(s) to control excess fruit production. A second strategy is also used by trees that bear a heavy fruit load (ON-Crop) in one year, which inhibits return bloom and vegetative growth the next year (Monselise and Goldschmidt, 1982). Thus, the second year is characterized by low yield (OFF-Crop) and high vegetative growth. Such trees, including olive, pistachio, mandarins, and many others, are defined as alternate or biannual bearers and they are usually characterized by low self-thinning ability (Goldschmidt, 2013). Alternate bearer cultivars present a serious economic problem to fruit growers. Therefore, chemical or manual fruit thinning are common practices in their cultivation (Dennis, 2000). In citrus culture, low temperatures

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during the autumn and winter are a major factor in inducing flowering (Valiente and Albrigo, 2004; Knauer *et al.*, 2011). Optimal flowering density is achieved only upon accretion of sufficient cool hours. It is assumed that a heavy fruit load prevents recognition of the low-temperature flowering inductive signal and/or blocks later stages of inflorescence, such as bud break (Albrigo and Galán-Saúco, 2004; Verreynne and Lovatt, 2009). As expected, fruit load affects the expression of flowering control genes, *FT*, *LFY*, *AP1*, *TFL*, and *miR156*regulated *SQUAMOSA PROMOTER BINDING* (*SPL5*) in leaves and buds of citrus (Muñoz-Fambuena *et al.*, 2011, 2012*b*; Shalom *et al.*, 2012) as well as in mango and apple (Kotoda *et al.*, 2010; Nakagawa *et al.*, 2012).

The mechanism by which heavy crop load affects return bloom is not fully understood. The developing fruit provides a strong sink for photoassimilates. It was therefore thought that depletion of photoassimilates, especially carbohydrates from the bud, prevents flowering induction, a hypothesis known as the nutritional theory (Goldschmidt et al., 1985; Goldschmidt, 1999). Sucrose was shown to play a regulatory role in Arabidopsis flowering control (Eriksson et al., 2006), but whether sugars indeed play a regulatory role in flowering induction under various fruit loads in fruit trees has been a controversial issue for many years (Hilgeman et al., 1967; Jones et al., 1970, 1974; Goldschmidt and Golomb, 1982; Li et al., 2003a, b). Recent work has shown that trehalose metabolism and its product trehalose-6-phosphate were involved in flowering control in Arabidopsis (van Dijken et al., 2004; Wahl et al., 2013). It was also shown that two genes encoding enzymes associated with trehalose metabolism were induced in OFF-Crop buds (Shalom et al., 2012). In addition to the nutritional control of alternate bearing (AB), it might well be that the fruit itself, or an organ which senses fruit presence, generates an inhibitory signal (AB signal) which moves into the bud and prevents flowering induction (Bower et al., 1990; Tálon et al., 1997). Fruit thinning or complete removal (de-fruiting) from ON-Crop trees induces return bloom (Monselise and Goldschmidt, 1981), thus providing support for this notion. Gibberellin (GA) is known to inhibit flowering in many perennials (Goldschmidt and Samach, 2004; Bangerth, 2009). However, while exogenous application of GA prevents flowering (Goldschmidt et al., 1997; Muñoz-Fambuena et al., 2012a; Goldberg-Moeller et al., 2013), the question of whether GA acts endogenously to inhibit flowering is still open. The involvement of abscisic acid (ABA) in the regulation of return bloom is even less clear (Jones et al., 1976; Goldschmidt, 1984; Koshita et al., 1999; Okuda, 2000). Polar auxin transport from a dominant sink was also suggested as a possible mobile signal affecting flowering (Caaejas and Bangerth, 1997; Smith and Samach, 2013).

Fruit load might act at various developmental stages such as flowering induction, transition of the shoot apical meristem, and subsequent stages of flower development and bud break (Verreynne and Lovatt, 2009). Regardless of the source of the AB signal and its nature, it must be recognized by its receptor in the bud which in turn must make the 'decision' of whether to proceed to inflorescence or not. In order to investigate metabolic and regulatory processes taking place in the bud and affected by fruit load, the transcriptome of buds from ON- and OFF-Crop trees was recently compared during three developmental stages. Changes in metabolic and regulatory pathways, including photosynthesis, and in flavonoid and trehalose metabolism were identified (Shalom et al., 2012). However, this work was biased due to the use of an Affymetrix Citrus Gene-Chip array that contained ~15 500 genes. In fact, with the exception of trehalose metabolism, no other regulatory pathways were identified. In the current work, a complementary approach was taken by comparing the transcriptome of buds of de-fruited trees with those of ON-Crop trees. The genomic analysis was non-biased, as it was based on RNA-deep sequencing. It was possible to identify an increase in ABA-metabolizing genes, accompanied by a decrease in ABA levels and those of its catabolites in buds of de-fruited trees. Moreover, a remarkable increase in the expression of genes encoding proteins associated with calcium-dependent auxin polar transport and a reduction in bud endogenous auxin levels following de-fruiting were identified. The results are discussed in light of the previously suggested auxin transport autoinhibition (ATA) theory (Bangerth, 1989) and its role in AB (Smith and Samach, 2013).

#### Materials and methods

#### Plant material and sample collection

Plant material was collected from a commercial orchard of 15-yearold Murcott mandarin (Citrus reticulate Blanco) trees grafted on sour orange (Citrus aurantium L.), located in the central coastal area of Israel, during the years 2011 (an ON year) and 2012 (an OFF year). Although most of the trees in the orchard bore similar yields in a given year, some were aberrant and showed an opposite AB trend. These and nearby trees with the opposite yield status were selected. Overall, nine triplets of trees were chosen, with each triplet (two ON trees and a nearby OFF tree) being considered one biological replicate. Fruits were completely removed (de-fruiting) on 22 August 2011 from one of the ON trees in each triplet, and this tree was labelled DEF. Samples were collected 1 d prior to de-fruiting (Time 0, from ON and OFF trees), 1 week following de-fruiting (Time 1, from ON and DEF trees), 2 weeks following de-fruiting (Time 2, from ON and DEF trees), and 4 weeks following de-fruiting (Time 4, from ON, OFF, and DEF trees).

The three most extreme conditions of spring flush flowering were compared (see fig. S2 in Shalom *et al.*, 2012): fruit-bearing flush of an ON tree, fruitless flush of an OFF tree, and de-fruited flush of a DEF tree. Branches of each of these conditions were collected from the southeast side of the trees, taken to the laboratory on ice, and buds were separated and immediately frozen in liquid nitrogen. The percentages of generative, mixed, and vegetative shoots were determined for all branches splitting off from one major 50–60 mm diameter branch located on the southeast side of the tree, during peak blossoming time (usually the first 10 d of April of the consecutive year). Selection of the branches to be sampled was done prior to bud break.

#### RNA extraction, quantification, and qPCR analyses

Total RNA was extracted, treated, and analysed from ~0.2g of frozen bud tissue, and cDNA was synthesized, as previously described (Shalom *et al.*, 2012). Primers for the genes *CiFT2*, *CsLFY*, *SPL5*, *RbcS*, *LHCB3*, *PRK*, *PSB28*, *PSAD*, *SHM*, *Fd*, *NCED3*, *CAX*, *PBP1-like1*, *PBP1-like2*, *NPH3*, *CA-binding EF hand1*, β-ACTIN, and a dual-labelled probe for *CiFT2* were designed based on genomic and expressed sequence tag (EST) sequences (Phytozome, http://www. phytozome.net/; HarvEST, http://harvest.ucr.edu/) using Primer 3 software (Supplementary Table S1 available at *JXB* online). Realtime PCR was carried out as described (Shalom *et al.*, 2012). For the *CiFT2* dual-labelled probe reaction, real-time PCR was carried out as described by Shalom *et al.* (2012). The mRNA levels of trehalose biosynthetic genes and flavonoid biosynthetic genes (Supplementary Table S2) were determined by nCounter analysis (Nanostring Technologies, Seattle, WA, USA) at the VIB MicroArrays Facility (Leuven, Belgium), as described by Shalom *et al.* (2012).

#### RNA deep sequencing

Extracted RNA integrity was determined by Agilent Bioanalyzer (Santa Clara, CA, USA) according to the manufacturer's instructions. A 2 µg aliquot of total RNA from each sample was prepared and used for cDNA library constructions using the TruSeq mRNA sample preparation kit according to the manufacturer's protocol (Illumina Inc., San Diego, CA, USA, REF 15025062). The libraries (10 pmol) were run on a single read 100 nucleotide run on the HiSeq 2000 (Illumina Inc.) on six lanes. Raw fastq files were quality checked using FastQC (http://www.bioinformatics.babraham.ac.uk/ projects/fastqc), and adaptor sequences were removed using fastqmcf (https://code.google.com/p/ea-utils/wiki/FastqMcf). They were then aligned to the orange (Citrus sinensis) genome database (Xu et al., 2013; http://citrus.hzau.edu.cn/orange/) using TopHat2 (Kim et al., 2013). FPKM (fragments per kilobase of transcript per million mapped reads) values, which normalize the read count by the length of the fragment and the total number of mapped reads, were calculated, and differential expression was checked using Cufflinks (Trapnell et al., 2010). A hierarchical clustering heatmap and a 2D principle component analysis (2D PCA) plot were generated by MATLAB (Mathworks, Cambridge, UK), using the log-FPKM values of each gene. For gene ontology (GO) analysis, sequences were blasted against the UniRef90 database (Suzek et al., 2007), by which a number of GO annotations were derived for each gene. Singular enrichment analysis (SEA), which lists enriched GO terms, was performed [false discovery rate (FDR)  $\leq 0.05$ ] for the five best GO terms of each gene using the AgriGo interface (http://bioinfo.cau.edu.cn/ agriGO/index.php). Differentially expressed genes were also functionally annotated via the automated Mercator pipeline (Lohse et al., 2013) (http://www.gabipd.org/biotools/mercator/) and displayed on diagrams of metabolic and other processes using MapMan (Usadel et al., 2009; http://www.gabipd.org/projects/MapMan/).

#### Hormonal content analysis

About 50 mg of bud tissue from each tested sample were lyophilized and homogenized to a fine powder in liquid nitrogen using a mortar and pestle. Quantification of ABA, ABA metabolites, and indole acetic acid (IAA) was conducted at the National Research Council of Canada (Saskatoon, Saskatchewan, Canada) according to published protocols (http://www.nrc-cnrc.gc.ca).

#### Protein extraction and western blot analysis

Proteins were extracted from buds, quantified, and analysed by western blot analysis using specific primary antibodies, raised against the following proteins: Rubisco complex large (RbcL) and small (RbcS) subunits, ferredoxin (Fd), chlorophyll *alb* protein [light-harvesting chlorophyll *alb* complex II (LHCII)], and D1 protein (PsbA), as described by Maayan *et al.* (2008).

#### Statistical analysis

The statistical analyses used for qPCR results, hormone analyses, and inflorescences numbers were one-way analysis of variance (ANOVA) with Tukey–Kramer multiple comparison tests as implemented in the software JMP version 10 (SAS Institute).

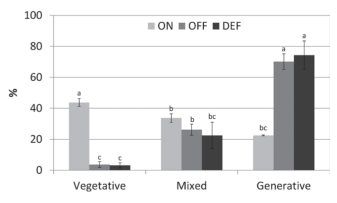
#### **Results**

#### Fruit removal (de-fruiting) induces back flowering

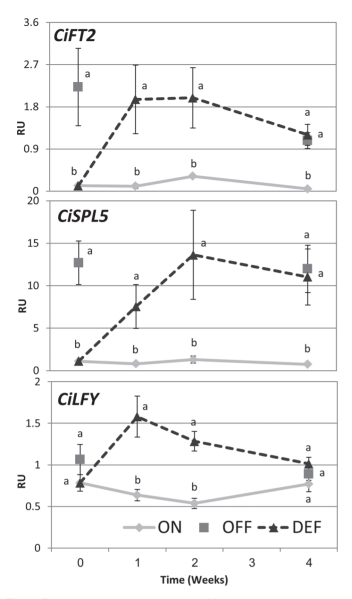
Normally, fruit load status is similar among most trees of the orchard-trees bear either a heavy crop (ON-Crop year) or a low crop (OFF-Crop year). A few trees, however, show the opposite trend, allowing the collection of samples from nearby trees bearing either high fruit load or low fruit load. In order to detect changes which might play a role in converting ON- to OFF-Crop buds, fruits were completely removed from ON-Crop trees in August and buds were collected 1, 2, and 4 weeks following de-fruiting. The effect of the treatment was verified the following spring by counting the number of inflorescences and vegetative shoots (Fig. 1). Citrus trees bear two major types of inflorescences: generative (leafless) and mixed (leafy; flowers and leaves in various ratios). ON-Crop trees had significantly less generative inflorescences compared with OFF-Crop trees and DEF trees (22% versus 70–75%) and more vegetative shoots (43%) versus 3-4%). As expected, no significant differences were detected in mixed-type inflorescences. Fruit counting during harvest time showed that ON-Crop trees yielded  $1635 \pm 98$ fruits per tree while OFF-Crop trees yielded  $36 \pm 12$  fruits per tree.

## Fruit removal alters expression of flowering control and other genes in the bud

In order to determine how quickly buds responded to defruiting, the mRNAs levels of several flowering control genes were quantified during the course of the experiment (Fig. 2). The *Citrus* genome contains three *FT* genes, but only the expression of *CiFT2* correlated well with tree flowering intensity (Nishikawa *et al.*, 2007; Shalom *et al.*, 2012). In buds of ON-Crop trees, *CiFT2* mRNA levels were relatively low, and remained unchanged. In contrast, *CiFT2* mRNA was ~15fold higher in buds of OFF-Crop trees. One week following de-fruiting, the expression of the gene in the buds of the DEF



**Fig. 1.** Fruit load affects flowering intensity the following year. Vegetative shoots, generative inflorescences containing only flower buds, and mixed inflorescences containing flower buds and leaves in various ratios were counted during flowering peak in trees which carried a heavy yield (ON) or light yield (OFF), and in de-fruited trees (DEF). The numbers are mean values of three independent biological replicates ±SE. Different letters represent a significant difference ( $P \le 0.05$ ).



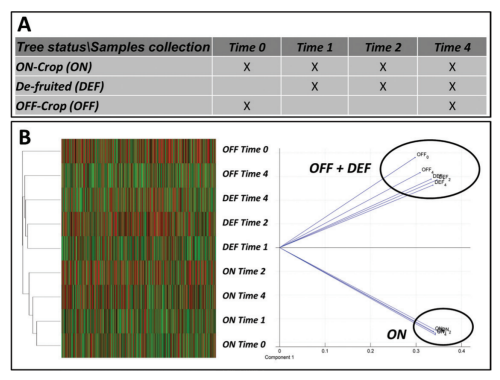
**Fig. 2.** Fruit removal alters the expression of flowering control genes in buds. The mRNA levels (RU, relative units) of the indicated genes were determined in ON-Crop (ON), OFF-Crop (OFF), and de-fruited (DEF) trees at the indicated weeks following de-fruiting. The numbers are mean values of three independent biological replicates  $\pm$ SE. Different letters represent a significant difference (*P*≤0.05) between the states at the same time point.

trees was similar to that of OFF-Crop buds, and it remained at this level during the entire test period. The mRNA levels of *LFY* were similar in buds of ON- and OFF-Crop trees, but it increased 3-fold within 1 week in DEF buds and returned to its basal level after 4 weeks. It was previously shown that *miR156*-regulated *SPL5* displayed elevated mRNA levels in OFF-Crop buds; thus, it may act as a positive inducer of flowering in *Citrus* trees (Shalom *et al.*, 2012). As expected, *SPL5* mRNA levels were ~14-fold higher in buds of OFF-Crop trees as compared with ON-Crop buds. DEF buds displayed increased mRNA as compared with OFF-Crop buds within 2 weeks of de-fruiting. The expression of genes associated with trehalose and flavonoid metabolism was elevated in ON- and OFF-Crop buds (Shalom *et al.*, 2012). As expected, the mRNA levels of trehalose phosphate synthase and trehalose phosphate phosphatase were reduced by 2- and 4-fold, respectively following de-fruiting (Supplementary Fig. S1 at *JXB* online). The mRNA levels of flavonoid biosynthesis genes, *UF3GT*, *4CL*, *CHS*, and *CHI*, were induced by the treatment. De-fruiting resulted in a reduction of *C4H* mRNA levels in the ON-Crop trees.

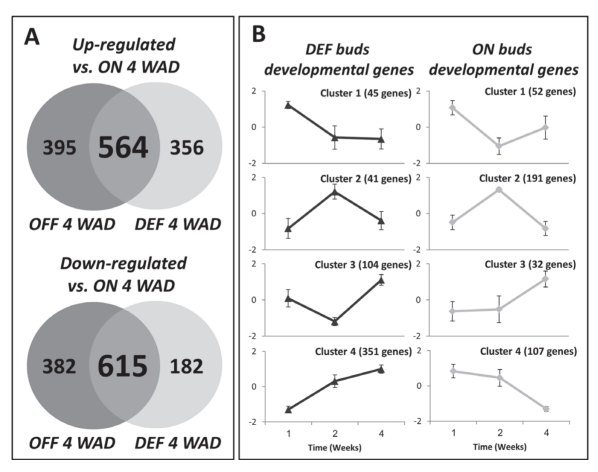
# Fruit removal induces rapid changes in the bud global gene expression

The results indicated that the transition of the bud from an ON to an OFF state took place relatively quickly. In order to analyse the metabolic and regulatory pathways playing a role in this transition, changes in global gene expression were analysed in buds before de-fruiting (Time 0) and 1 (Time 1), 2 (Time 2), and 4 (Time 4) weeks after de-fruiting (Fig. 3A). In addition, buds of OFF-Crop trees were also analysed at Time 0 and Time 4. The minimum number of reads per sample was ~15 million and the maximum was ~40 million, indicating a deep and satisfactory coverage of the existing transcripts (Supplementary Table S3 at JXB online). Overall, the number of transcripts in all libraries was ~14 400. Hierarchical cluster analysis (Fig. 3B, left) and 2D PCA (Fig. 3B, right) were performed based on the log-FPKM values of each sample. Results of both analyses showed that during all time points, including Time 1, the transcript profiles of DEF buds were more closely related to those of OFF-Crop buds than to those of ON-Crop buds, thus supporting the notion that the transition from an ON bud to an OFF bud was relatively quick following de-fruiting. In order to analyse the metabolic and regulatory pathways mediating the ON bud to OFF bud transition, two major comparisons were made (the ratios for all possible comparisons in the experiment are presented in Supplementary File 2 at JXB online). First, Time 4 included the three fruit load states, ON, OFF, and DEF. Therefore, the genes that were up- or downregulated ( $P \le 0.05$ ) in OFF or DEF buds in comparison with ON-Crop buds were identified (Fig. 4A; Supplementary File 2). Overall, 997 genes in OFF-Crop buds and 797 genes in DEF buds were down-regulated relative to ON-Crop buds at Time 4, with 615 genes common to the two groups (OFF and DEF). Overall, 959 genes in OFF-Crop buds and 920 genes in DEF buds were up-regulated relative to ON-Crop buds at Time 4, with 564 genes common to the two groups (OFF and DEF) (Fig. 4A; Supplementary File 2). The second comparison aimed at identifying genes which showed alternation in their expression during the course of the experiment and also common or different pathways altered developmentally. For that, genes were clustered according to their expression patterns relative to Time 0. For ON-Crop and DEF buds, four clusters were identified (Fig. 4B; Supplementary File 2). For OFF-Crop buds, only two time points were analysed, and altered genes were either reduced or repressed (Supplementary File 2)

In order to identify common and unique metabolic and regulatory pathways in DEF and OFF-Crop buds which were altered in comparison with ON-Crop buds at Time 4, up-regulated genes (564+395 and 564+356, Fig. 4A) and



**Fig. 3.** Global gene expression in buds of de-fruited (DEF) trees is similar to that of buds of OFF-Crop (OFF) trees. The experimental design and sample collection (A). Hierarchical clustering heatmap (left panel) and 2D principle component analysis (right panel) plots were generated using the log-*FPKM* value of each gene for all samples, as indicated in A (B). (This figure is available in colour at *JXB* online.)



**Fig. 4.** Venn diagrams of differentially expressed genes and clustering analysis of developmentally altered genes. The number of genes in buds of OFF-Crop (OFF) and de-fruited (DEF) trees which were significantly ( $P \le 0.05$ ) up-regulated (left) or down-regulated (right) compared with ON-Crop buds at Time 4 (A). Clustering analysis of genes which were significantly ( $P \le 0.05$ ) altered developmentally during the course of the experiment in comparison with Time 0 as generated by Expender (http://acgt.cs.tau.ac.il/expander/overview.html) using the Click Algorithm (B). All genes are listed in Supplementary File 2 at *JXB* online.

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down-regulated genes (615+382 and 615+182, Fig. 4A) were GO annotated and further analysed using SEA (AgriGO). Table 1 presents these common processes. However, in a few cases, unique GO terms were included under the same general process either because of redundancy or because they were part of one process. The analysis indicated that the great majority of common processes which were enriched in DEF/OFF-Crop buds were related to light sensing (such as response to light stimulus/intensity and response to far red/ blue and red light), photosynthesis (such as dark and light reactions and photosynthetic electron transport chain), chloroplast reorganization (such as plastid localization, chloroplast relocation and organization), response to carbohydrate stimulus (i.e. sucrose and disaccharide), ion homeostasis (such as cation homeostasis and transport, proton transport), and the pentose-phosphate cycle. Fewer genes were down-regulated in DEF/OFF-Crop buds and belonged to several secondary metabolic pathways such as the terpenoid and phenylpropanoid metabolic process, and oxidative reduction. Major unique processes induced in OFF-Crop buds included starch biosynthesis, carbohydrate metabolism, glycoside and glycosinolate metabolism, and water homeostasis (Supplementary Table S4 at JXB online). Considerably more unique processes were induced in DEF buds as compared with OFF-Crop buds (Supplementary Table S4). They included regulation of peptidase activity, regulation of dephosphorylation, and salicylic acid metabolism. Downregulated unique processes in DEF buds included amino acid/amine metabolism and aromatic compound metabolism (Supplementary Table S4). Responses to biotic stress were either induced or repressed in DEF buds. Unique genes down-regulated in OFF-Crop buds could not be GO annotated, due to their low number.

GO annotation and SEA were also performed to developmentally altered genes compared with Time 0 in ON-Crop, OFF-Crop, and DEF buds (Fig. 4B). Significantly altered biological processes (FDR  $\leq 0.05$ ) could be identified for genes of clusters 3 and 4 (DEF buds), cluster 2 (ON-Crop buds), and up/down regulated genes of OFF-Crop buds. No significant biological processes could be identified even under FDR ≤0.1 in genes of the other clusters. Cluster 4 of DEF buds included processes involved in light responses and photosynthesis (Supplementary Table S5 at JXB online). A considerable induction was also detected in genes of respiratory burst and in those involved in responses to abiotic stresses. Genes up-regulated in OFF-Crop buds from Time 0 to Time 4 were annotated into light responses, photosynthesis, and response to carbohydrate stimulus (Supplementary Table S6). Down-regulated genes in OFF-Crop buds from Time 0 to Time 4 included responses to jasmonic acid (JA) and ethylene (Supplementary Table S7). Other processes included amino acid and amine metabolism, and responses to abiotic stresses. Cluster 3 of DEF buds included 104 genes that could be annotated into different biological processes, a few of them related to development and morphogenesis, such as trichoblast and root differentiation, floral organ development, and shoot morphogenesis (Supplementary Table S8). Cluster 2 of ON-Crop buds was enriched in genes associated with lipid transport (Supplementary Table S9).

An additional analysis for developmentally regulated genes was aimed at identifying common genes in clusters showing a similar expression pattern in OFF-Crop and DEF buds and opposite patterns in ON-Crop buds. The first comparison included cluster 4 of DEF buds, up-regulated genes of OFF-Crop buds, and cluster 4 of ON-Crop buds (Supplementary Fig. S2A at JXB online). However, only one unknown gene was common among the three states. The second comparison included cluster 1 of DEF buds, down-regulated genes of OFF-Crop buds, and cluster 3 of ON-Crop buds (Supplementary Fig. S2B). Only the expression of three genes was common; two of them were homologous to terpene (nerolidol) syntheses (*Cs2g07240* and *Cs2g07250*).

## Photosynthetic genes and proteins are up-regulated in response to fruit removal

Consistent with a previous report (Shalom et al., 2012), the above-described results and MapMan analysis of differentially expressed genes in DEF versus ON-Crop buds at Time 4 showed that genes associated with light reactions, the Calvin-Benson cycle, and to a lesser extent photorespiration were induced (Supplementary Fig. S3 at JXB online). Fold changes of 41 photosynthetic genes in DEF and OFF-Crop buds relative to ON-Crop buds are shown in Fig. 5A. Out of 41 differentially expressed genes, 38 were up-regulated while only three were down-regulated at at least one time point throughout the experiment. Validation of the above results by qPCR analyses was performed for seven genes, Ribulose bisphosphate carboxylase-small subunit (RbcS), Light-harvesting chlorophyll B-binding protein (LHCB3), Phosphoribulokinase (PRK), Photosystem II reaction centre PSB28 (PSB28), Photosystem I subunit D (PSAD), Serine hvdroxymethyltransferase (SHM), and Ferredoxin (Fd) (Fig. 5B). The mRNA levels of all these genes were significantly higher in buds of OFF trees as compared with ON trees at Time 4. As expected, gene expression increased in DEF buds by 2.6- to 5.2-fold within 1 week of fruit removal. Furthermore, western blot analyses using specific antibodies raised against RbcS, Fd, LHCII, and PsbA showed that their protein level was higher in OFF-Crop buds relative to ON-Crop buds, and increased in DEF buds (at Time 4) following fruit removal (Fig. 5C). However, RbcL protein levels remained unchanged in DEF and OFF-Crop buds relative to ON-Crop buds.

# Fruit removal-induced changes in ABA-metabolizing genes

Results of genomic analysis showed that genes homologous to 9-cis-epoxycarotenoid dioxygenase (NCED), coding for this rate-limiting enzyme of ABA biosynthesis (Supplementary Fig. S5 at JXB online), were higher in buds of OFF-Crop buds relative to ON-Crop buds (Fig. 6A). In Arabidopsis, the NCED gene family comprises nine members, and the roles of NCED2, 3, 5, and 6 in ABA biosynthesis were demonstrated (Tan et al., 2003). The Citrus genome contained nine highly homologous genes, and changes in three of them were detected in genomic analysis: Cs5g14370.1, homologous to NCED3, Cs5g14370.1, homologous to NCED1, and

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General process	GO term	Description	Ref	OFF				DEF			
			item	Query item	<i>P</i> -value	FDR	Fold enrichment <sup>a</sup>	Query item	<i>P</i> -value	FDR	Fold enrichment <sup>a</sup>
Up-regulated											
Response to light	GO:0010218	Response to far red light	25	11	2.80E-07	0.00057	6.48	14	7.20E-11	9.40E-08	8.51
	GO:0010114	Response to red light	47	14	1.90E-06	0.00089	4.39	17	3.80E-09	3.00E-06	5.50
	GO:0009639	Response to red or far red light	-	22	0.00023	0.024	2.31	22	0.00015	0.014	2.39
	GO:0009416	Response to light stimulus	380	44	0.00055	0.047	1.70	59	2.20E-09	2.20E-06	2.36
	GO:0009637	Response to blue light	27	7	8.20E-05	0.049	6.39	11	5.20E-07	0.00015	6.19
	GO:0009644	Response to high light intensity	20					20	1.80E-08	8.90E-06	4.34
	GO:0009642	Response to light intensity	88					23	1.10E-08	7.00E-06	3.97
	GO:0009314	Response to radiation	404					59	2.10E-08	9.30E-06	2.22
Photosynthesis and carbon fixation	GO:0015977	Carbon fixation	7	9	6.60E-07	0.00068	12.62	Q	2.40E-05	0.0031	10.85
	GO:0009773	Photosynthetic electron transport in	12	7	4.00E-06	0.0013	8.59	Ŋ	0.00067	0.038	6.33
		photosystem I									
	GO:0015979	Photosynthesis	128	33	2.90E-11	1.20E-07	3.80	47	9.80E-23	3.90E-19	5.58
	GO:0009767	Photosynthetic electron transport chain	31	10	2.60E-05	0.0059	4.75	11	2.60E-06	0.00055	5.39
	GO:0019684	Photosynthesis, light reaction	85	19	3.90E-06	0.0013	3.29	31	1.60E-15	3.10E-12	5.54
	GO:0019685	Photosynthesis, dark reaction	Ð	Ð	1.50E-06	0.00087	14.72				
	GO:0010207	Photosystem II assembly	36					13	2.60E-07	8.50E-05	5.49
	GO:0009765	Photosynthesis, light harvesting	13					Ø	3.50E-07	0.00011	9.35
Chloroplast	GO:0051667	Establishment of plastid localization	28	6	6.70E-05	0.011	4.73	Ø	0.00034	0.023	4.34
	GO:0051644	Plastid localization	28	6	6.70E-05	0.011	4.73	Ø	0.00034	0.023	4.34
	GO:0009902	Chloroplast relocation	28	0	6.70E-05	0.011	4.73	Ø	0.00034	0.023	4.34
	GO:0051656	Establishment of organelle localization	з <del>1</del>	0	0.00016	0.02	4.27	œ	0.00073	0.04	3.92
	GO:0009658	Chloroplast organization	81	18	7.60E-06	0.002	3.27	14	0.00083	0.044	2.63
	GO:0009657	Plastid organization	123	21	9.50E-05	0.014	2.51	20	0.00018	0.015	2.47
Response to carbohydrate stimulus	GO:0009744	Response to sucrose stimulus	47	12	5.60E-05	0.011	3.76	14	1.30E-06	0.00033	4.53
	GO:0034285	Response to disaccharide stimulus	48	12	7.10E-05	0.011	3.68	14	1.80E-06	0.00039	4.43
	GO:0009743	Response to carbohydrate stimulus	178	31	1.70E-06	0.00088	2.56	33	9.30E-08	3.30E-05	2.82
Cellular homeostasis	GO:0071704	Organic substance metabolic process	7	9	6.60E-07	0.00068	12.62	Ŋ	2.40E-05	0.0031	10.85
	GO:0019725	Cellular homeostasis	129	23	2.30E-05	0.0055	2.63	21	0.00012	0.012	2.47
	GO:0055082	Cellular chemical homeostasis	57	12	2.90E-06	0.0037	5.19	16	5.90E-07	0.00016	4.26
Ion homeostasis	GO:0006873	Cellular ion homeostasis	52	12	0.00016	0.02	3.40	14	5.00E-06	0.00095	4.09
	GO:0030003	Cellular cation homeostasis	50	11	0.00046	0.043	3.24	13	1.60E-05	0.0023	3.95
	GO:0055082	Cellular chemical homeostasis	57	15	4.70E-06	0.0014	3.87	41	0.00013	0.012	1.87
	GO:0006812	Cation transport	333	43	6.00E-05	0.011	1.90	14	0.00037	0.023	2.84
	GO:0006811	lon transport	470	55	0.0001	0.014	1.72	52	0.00028	0.021	1.68
	GO:0010155	Regulation of proton transport	1-	9	3.40E-05	0.0075	8.03	7	1.40E-06	0.00033	9.67
	GO:0055080	Cation homeostasis	64					13	0.00025	0.019	3.09
Pentose cycle	GO:0019253	Reductive pentose-phosphate cycle	Ð	Ð	1.50E-06	0.00087	14.72				
	GO:0006098	Pentose-phosphate shunt	65					13	0.0003	0.022	3.04
Response to biotic stress	GO:0010200	Response to chitin	06	16	0.00038	0.038	2.62	17	8.10E-05	0.0089	2.87
	GO:0050832	Defence response to fungus	123	20	0.00027	0.028	2.39	20	0.00018	0.015	2.47
Abiotic stress	GO:0009611	Response to wounding	128	20	0.00046	0.043	2.30	20	0.00031	0.022	2.37

General process	GO term	Description	Ref	OFF				DEF			
			item	Query item	P-value	FDR	Fold enrichment <sup>a</sup>	Query item	<i>P</i> -value	FDR	Fold enrichment <sup>a</sup>
Down-regulated											
Secondary metabolism	GO:0019748	GO:0019748 Secondary metabolic process	467	58	5.80E-05	0.042	1.73	55	1.20E-06	0.0011	2.04
	GO:0006721	Terpenoid metabolic process	118	23	1.20E-05	0.016	2.72	18	0.00017	0.028	2.64
	GO:0006720	Isoprenoid metabolic process	186	29	8.10E-05	0.049	2.17	00	0.00031	0.044	4.46
	GO:0009699	Phenylpropanoid biosynthetic process	143					23	1.00E-05	0.0035	2.78
	GO:000698	Phenylpropanoid metabolic process	184					27	1.10E-05	0.0035	2.54
Various	GO:0055114	Oxidation reduction	1344	166	2.00E-10	7.40E-07	1.72	137	2.60E-09	7.50E-06	1.76
	GO:0042221	Response to chemical stimulus	1345	142	1.30E-05	0.016	1.47	120	1.00E-05	0.0035	1.54
	GO:0010035	GO:0010035 Response to inorganic substance	462	58	4.30E-05	0.039	1.75				
<sup>a</sup> Fold enrichment as calculated based o respectively) in the list of genes (959, 920, 711715 in the momented list (111700, 2000)	lated based on GC es (959, 920, 997,	<sup>a</sup> Fold enrichment as calculated based on GO-annotated genes (777, 753, 821, 662 in OFF-Crop up-regulated, DEF up-regulated, OFF-Crop down-regulated, DEF down-regulated, not concert and the list of genes (959, 920, 997, 797 in OFF-Crop up-regulated, DEF-Up down-regulated, DEF down-regulated, DEF down-regulated, not concert and the list of genes (959, 920, 997, 797 in OFF-Crop up-regulated, DEF-Up down-regulated, DEF down-regulated, not concert and the list of genes (1100, 2000, 997, 797 in OFF-Crop up-regulated, DEF-Up down-regulated, DEF-Up down-regulated, not concert and the list of genes (1100, 2000, 997, 797 in OFF-Crop up-regulated, DEF-Up down-regulated, DEF-Up down-regulated, respectively) per GO-annotated genes	2 in OFF- p-regula	-Crop up- ted, OFF-	-regulated, [ -Crop down	DEF up-regu -regulated, I	lated, OFF-Crop DEF down-regul	o down-re ated, resp	igulated, DE bectively) per	F down-reg r GO-annot	ulated, ated genes

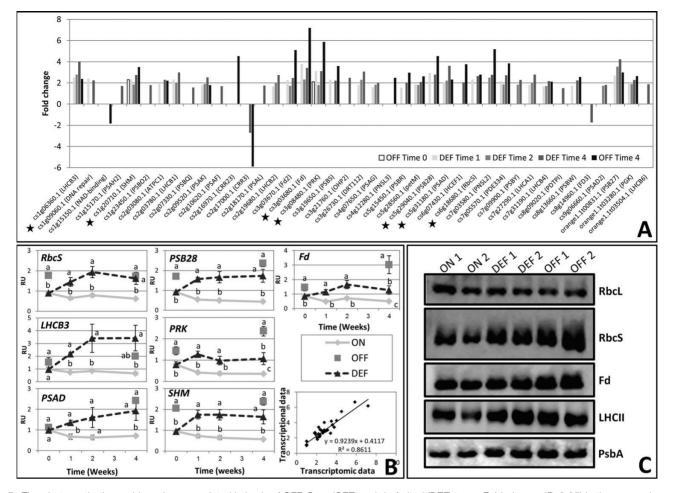
Table 1. Continued

Cs7g14820.1, homologous to NCED4. Among these three genes, NCED3-like induction was increased by ~4-fold in OFF-Crop buds relative to ON-Crop buds at Time 0, and was ~2-fold higher at Time 4. Fruit removal induced a 3-fold increase at Time 1 and its mRNA level remained higher relative to ON-Crop buds throughout the experiment. Induction of NCED1-like was seen only in OFF-Crop buds at Time 4, while NCED4-like was induced in DEF buds at Time 1 and 4 and in OFF-Crop buds at Time 4. Among the three genes, NCED3 is considered the major enzyme catalysing the ratelimiting step in ABA biosynthesis, and the mRNA levels of its Citrus counterpart were higher in OFF and DEF buds throughout the experiment. qPCR validation showed that Cs5g14370.1 mRNA levels in OFF-Crop buds were significantly higher relative to ON-Crop buds, by ~4.5-fold at Time 0 and by 2.1-fold at Time 4 (Fig. 6B). The mRNA levels in DEF buds increased relative to ON-Crop buds by 3.2-fold at Time 1 and remained higher throughout the experiment. The transcriptomic data showed that the expression of a gene homologous to PYR1, a component of the ABA receptor, was reduced (Supplementary Fig. S6).

The levels of ABA and its catabolites were analysed at three time points following de-fruiting (Fig. 7). In addition to ABA and its isomer, trans-ABA (t-ABA), four catabolites were detected in the buds, 7'OH-ABA, ABA glucose ester (ABAGE), phaseic acid (PA), and dihydrophaseic acid (DPA), with t-ABA, PA, and ABAGE showing relatively high levels. ABA levels in ON-Crop buds were significantly higher relative to OFF-Crop buds, by 2.4-fold at Time 0 and by 3.3-fold at Time 4. In DEF buds, ABA levels decreased relative to ON-Crop buds by 1.3-fold at Time 1 and by 1.8-fold at Time 4. In addition, the levels of PA and ABAGE were significantly higher in ON-Crop buds relative to OFF-Crop buds by 4.5fold and 4.6-fold at Time 1 and by 7.7-fold and 2.9-fold at Time 4, respectively. In DEF buds, PA and ABAGE levels decreased by 3.4-fold and 1.8-fold at Time 1 and by 4.1-fold and 3.5-fold at Time 4, respectively. The levels of DPA and t-ABA were generally higher in ON-Crop buds relative to OFF and DEF buds, especially at Time 4, but the differences were not significant.

#### Induction of calcium-related genes associated with auxin transport in DEF and OFF-Crop buds

Among the genes which showed relatively high levels of expression in OFF-Crop and DEF buds relative to ON-Crop buds were calcium-related genes (Fig. 8A; Supplementary File 2 at JXB online). The expression of most of these genes was induced 3- to 40-fold at all time points of the experiment. Many of these genes encode proteins containing an EF-hand domain. PINOID (PID)-binding protein (PBPs) are a subgroup of EF-hand proteins. PBP1 has been shown to play a role in auxin polar transport in response to changes in calcium levels (Benjamins et al., 2003). Four citrus PBP1like genes (out of the five found in the citrus genome) were induced in DEF and OFF-Crop buds relative to ON-Crop buds at all time points tested (Fig. 8A). Phylogenic analysis of these genes showed remarkable homology with PBP1 and its closely related gene in Arabidopsis (Supplementary Fig. S8A).

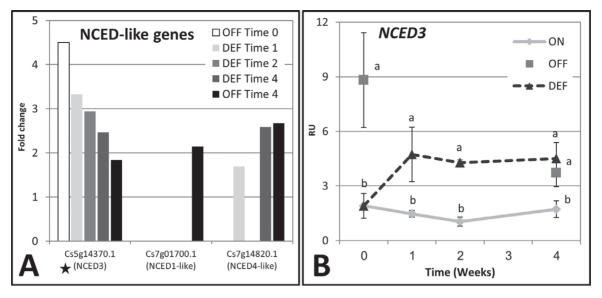


**Fig. 5.** The photosynthetic machinery is up-regulated in buds of OFF-Crop (OFF) and de-fruited (DEF) trees. Fold-change ( $P \le 0.05$ ) in the expression of photosynthetic genes (determined by MapMan analysis, see Supplementary Fig. S2 at *JXB* online) in buds of OFF and DEF trees relative to buds of ON-Crop (ON) trees at the indicated time points. Asterisks mark genes selected for validation by qPCR analyses, and specific genes are listed in Supplementary File 2 (A). Expression analysis of selected genes at the indicated weeks following de-fruiting, as determined by qPCR analyses (RU, relative units). The numbers are mean values of three independent biological replicates ±SE. Different letters represent a significant difference ( $P \le 0.05$ ) between the states at the same time point. The lower right graph shows the linear regression between the transcriptomic and transcriptional (qPCR) data (B). Immunoblot analyses in two independent replicates of photosynthetic proteins (RbcL, Rubisco large subunit; RbcS, Rubisco small subunit; Fd, ferredoxin; LHCII, light-harvesting complex II; Psba, D1 protein) extracted from two replicates of buds of ON-Crop, (ON1 and ON2), OFF-Crop (OFF1 and OFF2), and de-fruited (DEF1 and DEF2) trees at Time 4 (C). The quantification of the protein signals, generated using ImageJ software, is presented in Supplementary Fig. S4.

Two other induced genes, Cs9g20300.1 and Cs8g20150.1, showed high homology (80% and 70%, respectively) to a calcium-dependent protein kinase (At1g08650.1), and to a Ca<sup>2+</sup>/ H<sup>+</sup> antiporter CAX3 (At3g51860), respectively. Another gene, Cs1g21460.1, which showed homology to members of the NPH3 gene family from Arabidopsis also plays a role in auxin polar transport (Furutani et al., 2011; Knauer et al., 2011; Li et al., 2011; Wan et al., 2012). This family contains 33 genes in the Arabidopsis genome, and the Citrus genome comprises 25 homologous genes with relatively close taxonomic relationships (Supplementary Fig. S8B). The NPH3-like gene was induced 25- to 40-fold in OFF-Crop and DEF buds relative to ON-Crop buds at all time points tested (Fig. 8A). The plausible mechanistic relationships between calcium, PBP1, NPH3, and the polar subcellular localization of PIN-FORMED (PIN) auxin efflux carriers are schematically presented in Fig. 8B. The expression levels of CAX-like, NPH3-like, two *PBP1-like* genes, and one Ca<sup>2+</sup>-binding EF-hand gene were validated by qPCR analyses (Fig. 8C). The mRNA levels of all these genes were relatively low in ON-Crop buds and remained low throughout the experimental period. In OFF-Crop buds, they were significantly higher at Time 0 and Time 4 (by factors of 5–19 and 10–28, respectively). One week after de-fruiting, the expression of all these genes was significantly increased, and within 2 weeks they attained their maximal levels. While the mRNA levels of CAX-like, NPH3-like, and  $Ca^{2+}$ -binding EF-Hand1-like remained high 4 weeks after defruiting, those of PBP1-like1 and PBP1-like2 were reduced by ~2-fold and 1.5-fold, respectively, during the same period.

#### Auxin levels are significantly higher in ON-Crop buds and decrease following fruit removal

Next, levels of endogenous IAA were examined in the buds (Fig. 9). IAA levels were significantly higher in ON-Crop buds relative to OFF-Crop buds by 2.9-fold at Time 0 and by



**Fig. 6.** *NCED*-like genes are induced in buds of de-fruited (DEF) trees. Fold change ( $P \le 0.05$ ) in the expression of three *NCED*-like genes in buds of OFF-Crop (OFF) and DEF trees relative to buds of ON-Crop (ON) trees (an asterisk marks the *NCED3*-like gene which was validated by qPCR) (A). Expression of the *NCED3*-like gene in buds of ON, OFF, and DEF trees as determined by qPCR analyses at the indicated weeks following de-fruiting (RU, relative units). The numbers are mean values of three independent biological replicates  $\pm$ SE (B). Different letters represent a significant difference ( $P \le 0.05$ ) between the states at the same time point.

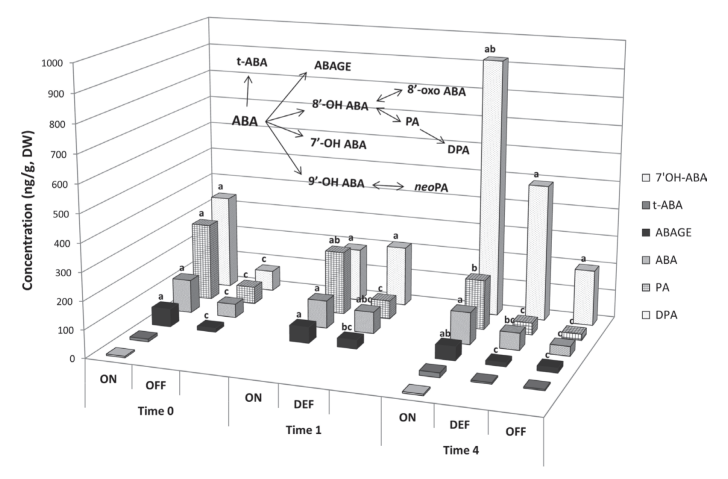
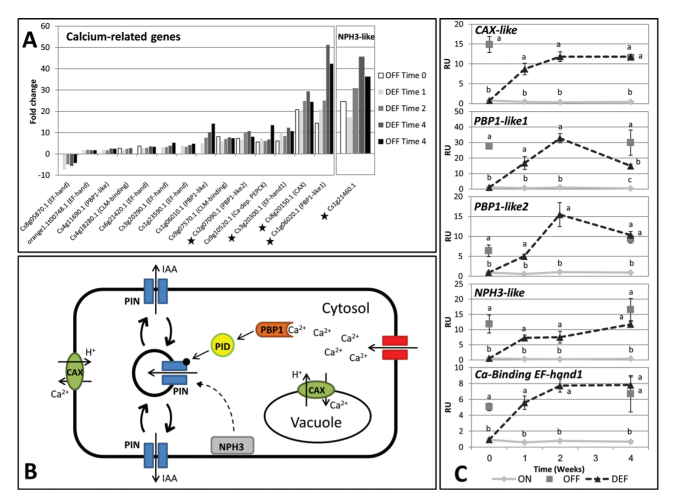
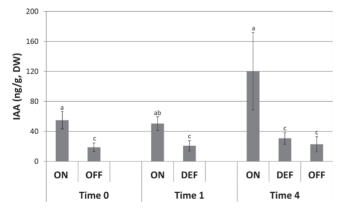


Fig. 7. Fruit load affects the levels ABA and its catabolites in the bud. ABA and ABA catabolites (7'OH ABA, t-ABA, trans-ABA; ABAGE, ABA glucose ester; PA, phaseic acid; DPA, di-hydrophaseic acid) whose metabolic relationships are schematically represented in the inner scheme were determined at the indicated time points in buds of ON-Crop (ON), OFF-Crop (OFF), and de-fruited (DEF) trees. Means followed by different letters are significantly different (*P*>0.05) according to Tukey–Kramer multiple comparisons tests.



**Fig. 8.** Fruit load affects the expression of  $Ca^{2+}$ -related auxin polar transport and *NPH3-like* genes. Fold change (*P*≤0.05) in the expression of  $Ca^{2+}$ -related and *NPH3-like* genes in buds of OFF-Crop (OFF) and de-fruited (DEF) trees relative to buds of ON-Crop (ON) trees (asterisks mark genes selected for validation by qPCR analyses; all genes are listed in Supplementary File 2 at *JXB* online) (A). Expression of the indicated genes in buds of ON, OFFm and DEF trees as validated by qPCR at the indicated time points following de-fruiting (RU, relative units). The numbers represent mean values of three independent biological replicates ±SE. Different letters represent a significant difference (*P*≤0.05) between the states at the same time point. (B) Linear regression between transcriptomic data and transcriptional data (qPCR analyses) is presented in Supplementary Fig. S7. Schematic model representing Ca<sup>2+</sup> and NPH3 regulation of PIN cellular localization (C). (This figure is available in colour at *JXB* online.)



**Fig. 9.** Fruit load affects auxin level in buds. Auxin was determined at the indicated time points in buds of ON-Crop (ON), OFF-Crop (OFF), and de-fruited (DEF) trees. The numbers are mean values of three independent biological replicates. Means followed by different letters are significantly different (*P*>0.05) according to Tukey–Kramer multiple comparisons tests. (This figure is available in colour at *JXB* online.)

5.2-fold at Time 4. As expected, IAA levels decreased in DEF buds relative to ON-Crop buds, by a factor of 2.4 at Time 1, and remained at this level 4 weeks after de-fruiting.

#### Discussion

Intensity of return bloom is affected by de-fruiting, which induces relatively rapid changes in expression of flowering control genes and in the transcriptome

Fruit removal has been reported to be effective in inducing return bloom (Monselise and Goldschmidt, 1981). However, since annual variation and cultivar-dependent divergence may affect its effectiveness (Verreynne and Lovatt, 2009; Martinez-Fuentes *et al.*, 2010; Muñoz-Fambuena *et al.*, 2011), fruit removal was carried out as early as the last 10 d of August. Indeed, the number of inflorescences and vegetative shoots, counted during the following spring (Fig. 1), showed that

those of DEF trees were similar to those of OFF-Crop trees, thus demonstrating the effectiveness of the treatment. Although it could be expected that the number of generative inflorescences would be higher in DEF than OFF-Crop trees (Verreynne and Lovatt, 2009), the actual number was identical, probably due to the low number of fruits on OFF-Crop trees (~45-fold lower than in ON-Crop trees). De-fruiting resulted in relatively rapid changes in the expression of genes controlling and genes associated with trehalose and flavonoid biosynthesis, allowing determination of the time frame of the genomic analysis.

As a rule, the expression of flowering control genes is induced in leaves, buds, and stems in association with the onset of the flowering induction period (November-December) in regular bearer cultivars, and in AB cultivars during the OFF-Crop year, while GA treatment reduced their expression (Nishikawa et al., 2007; Muñoz-Fambuena et al., 2011, 2012b; Shalom et al., 2012; Goldberg-Moeller et al., 2013). CiSPL5 is an exception to this rule, probably due to its highly regulatory role, and it exhibited higher expression in buds from May until September (Shalom et al., 2012). CiFT2 expression was usually earlier than the onset of the flowering induction period (Shalom et al., 2012). Taking into account these expression patterns and the expected year to year alternation, the mRNA levels of CiFT2 and CiSPL5 were higher in OFF- than in ON-Crop buds. Therefore, the increase in the expression of CiFT2 and CiSPL5 in buds of DEF trees to levels similar to those in buds of OFF-Crop trees can be expected, and reinforces their role in return bloom. It is not surprising that during the time of the experiment, LFY did not show any difference in its expression between buds of ON- and OFF-Crop trees. Nevertheless, de-fruiting resulted in a 2-fold increase in its mRNA level, which returned to its basal level 4 weeks after treatment. Whether this temporary response has any relationship to return bloom requires further research.

The differences between bud populations-those with a 55% chance to flower (ON-Crop) and those with a 96% chance (OFF-Crop and DEF buds)-did not seem to be very high. However, genomic analysis resulted in numerous differentially expressed genes (DEGs), allowing the partial identification of mechanisms that convert ON into OFF buds. Previously it was shown that the number of DEGs between ON- and OFF-Crop buds was considerably lower in September than in May (Shalom et al., 2012). However, the present work showed that the number of these DEGs was quite high. This difference could be explained by year to year alternations, and differences in the methodologies used. Based on a cut-off of 50% coverage between sequences on the microarray and the currently identified sequences, and on at least 75% identity, it is estimated that only  $\sim$ 30–35% of the current sequences are present on the microarray, supporting this notion. Only 40% of the auxin transport-related genes (Fig. 8A) were found on the microarray. Below, three identified mechanisms, common to OFF and DEF buds, which are altered during the conversion of DEF buds into OFF buds, and might play a role in the signalling mechanism of fruit load are discussed.

# Induction of photosynthetic gene expression and protein levels in the bud following de-fruiting

In agreement with Shalom et al. (2012), this study demonstrates that de-fruiting induces expression of photosynthetic genes in the bud. Although a recent proteomic analysis did not show an increase in photosynthetic proteins in OFF-Crop trees (Muñoz-Fambuena et al., 2013), here the induced gene expression resulted in increased protein levels of four major genes. According to the C/N theory, the proteins of photosynthetic machinery represent the majority of leaf nitrogen which is directly related to photosynthetic capacity (Evans, 1989); thus, the induced levels of photosynthesis proteins would suggest the induction of photosynthesis in OFF buds, although direct evidence is missing. Although bud photosynthesis was never measured in fruit trees, leaf photosynthesis in relation to fruit load has been measured in previous studies. While some workers found no change in photosynthesis between leaves of ON- and OFF-Crop trees (Roper et al., 1988; Monerri et al., 2011; Nebauer et al., 2013), others reported increased photosynthetic and  $CO_2$ assimilation rates in fruit-bearing as compared with nonfruit-bearing trees (Fujii and Kennedy, 1985; Dejong, 1986; Gucci et al., 1995; Palmer et al., 1997; Iglesias et al., 2002; Syvertsen et al., 2003; Urban et al., 2004). Vegetative growth is induced in buds of OFF-Crop and DEF trees (Monselise and Goldschmidt, 1982), suggesting that increased photosynthesis may mark the initiation of vegetative growth. That is, due to fruit absence, the OFF-Crop and DEF trees are heavily loaded with photoassimilates, suggesting that by induction of its photosynthetic machinery, the bud signals to stop translocation of photoassimilates. The possibility that the flow of photoassimilates into the bud is reduced due to lower leaf photosynthesis in OFF-Crop trees, resulting in increased synthesis of photosynthesis proteins and higher CO<sub>2</sub> assimilation, cannot be excluded.

# Bud ABA is reduced in OFF-Crop trees and following de-fruiting compared with ON-Crop trees

Increased expression of three NCED-like genes, in buds of DEF and OFF-Crop trees compared with buds of ON-Crop trees, suggests the induction of ABA biosynthesis. However, direct measurements of ABA and its catabolites showed the opposite trend, namely reduced levels in buds of OFF-Crop and DEF trees. Direct biochemical evidence demonstrated that NCED3 (Cs5g14370) cleaved 9-cis-violaxanthin to form xanthoxin, a precursor of ABA (Kato et al., 2006); its expression paralleled ABA levels in the peel and during cycles of drought and re-watering of leaves and fruit (Rodrigo et al., 2006; Agustí et al., 2007). Therefore, one would expect higher ABA levels in OFF-Crop and DEF buds than in ON-Crop buds. A possible explanation of these apparently contradictory results is that the source of ABA in the ON-Crop bud is not within the bud itself, but external to it, and dependent on the presence of fruit. In OFF-Crop trees or following de-fruiting, the translocation of ABA from this source into the bud is blocked, at least partially, reducing the bud's

ABA contents, and inducing NCDE3 expression in order to increase endogenous ABA production. Nevertheless, the possibility cannot be excluded that induced expression of NCED genes is futile, and has no physiological role. A closer look at ABA-responsive genes in the transcriptomic data (overall, eight differentially expressed genes between the states) did not solve this contradiction, as no common trend in their response was evident (data not shown). Regardless, ABA levels (and the expression of ABA receptor component, PYR1-like) in OFF-Crop buds and following de-fruiting were reduced, raising the question of its possible involvement in AB control. Consistent with the present results, buds of ON-Crop trees have been shown to contain higher levels of ABA or its isomer, t-ABA, than those of OFF-Crop trees (Jones et al., 1976; Goldschmidt, 1984). It has been suggested that elevated levels of ABA in ON organs may reflect a stress imposed by the fruit overload. Moreover, ABA might serve as an inhibitor of return bloom, since the local application of ABA to Citrus unshiu buds in late December inhibited bud sprouting and intensive flowering (Garcialuis et al., 1986). Alternatively, the possibility that flowering promotes ABA activity has been suggested, since increased ABA levels were detected in leaves of OFF-Crop trees and following de-fruiting of ON-Crop trees in association with flowering induction (Koshita et al., 1999; Okuda, 2000). Whether ABA plays a role in AB control, or in other processes, such as maintaining the bud in an inactive state (Little and Edit, 1968; Horvath et al., 2003; Shalom et al., 2012), requires further investigation.

# De-fruiting induces genes of calcium-dependent auxin polar transport

The results showed an increase in the expression of calciumrelated genes together with significant reduction in auxin levels in OFF-Crop buds and in buds following de-fruiting as compared with ON-Crop buds. Changes in the concentration of cytosolic free Ca<sup>2+</sup> ([Ca<sup>2+</sup>]<sub>cvt</sub>), mediated by ion channels, Ca<sup>2+</sup>-ATPases, and Ca<sup>2+</sup>/H<sup>+</sup> antiporters, form the basis of the Ca<sup>2+</sup> signalling mechanism. The CAX-type antiporters are a family of cytosolic low-affinity Ca2+/H+ antiporters, which in Arabidopsis comprises six members. In Citrus there are four highly CAX homologous genes, and the expression of a CAX3 homologue was highly induced following de-fruiting. Transduction of Ca<sup>2+</sup> signals is carried out by specific calcium-binding proteins, containing a common structural motif called the 'EF-hand', a helix-loop-helix structure that binds a single  $Ca^{2+}$  ion (Day *et al.*, 2002). The present results showed a significant up-regulation of a few genes encoding EF-hand proteins in OFF and DEF buds compared with their level in ON buds. Overall, these results might suggest that  $[Ca^{2+}]_{cvt}$  is affected by fruit load, although at this stage a plausible scenario as to the nature of the change and its cellular signature cannot be suggested. How are these changes related to auxin polar transport? Four of the up-regulated EF-hand genes show remarkable homology to the genes encoding PBP1 in Arabidopsis. PBP1 interacts physically with PID protein kinase, regulating its activity in response to changes in calcium levels (Benjamins et al., 2003). PID

regulates the polarity of PIN proteins (Friml *et al.*, 2004), which are known to direct auxin flow (Wisniewska *et al.*, 2006). NPH3-like proteins have recently been shown to affect PIN localization (Furutani *et al.*, 2011; Wan *et al.*, 2012). As shown here, *NPH3-like* genes are part of a relatively large gene family. Divergence of its different members occurred before the divergence of *Arabidopsis* and *Citrus*. The citrus *NPH3-like* gene induced in OFF and DEF buds compared with ON buds showed very close homology to an *Arabidopsis* gene which has not yet been subjected to detailed analysis.

Taken together, the present results lead to the suggestion that higher levels of IAA in ON buds reflect their inability to distribute IAA efficiently via the Ca2+-dependent PIN-based polar auxin transport mechanism. In addition, efficient auxin removal from the bud appears to be a key component in transforming the ON bud into an OFF bud. The involvement of auxin in flowering inhibition following an ON-Crop year was recently suggested (Smith and Samach, 2013), and is based on the ATA hypothesis proposed by Bangerth (Bangerth, 1989, 2006; Caaejas and Bangerth, 1997). The application of auxin polar transport inhibitors resulted in flowering induction in a number of fruit trees (Bukovac, 1968; Ben-Tal and Lavee, 1985; Ito et al., 2001; Blaikie et al., 2004; Bangerth, 2006). The strong polar transport of auxin from the dominant sinks (i.e. the fruit or the seed), as suggested by the ATA hypothesis, preventing auxin export from the bud, would explain why auxin levels in OFF buds and in buds following de-fruiting are lower than in ON buds.

The parallel reduction in ABA and IAA levels in the bud would suggest cross-talk between the ABA and IAA signalling pathways. Such cross-talk interactions were suggested in *Arabidopsis* embryo axis elongation and root development (Belin *et al.*, 2009; Shkolnik-Inbar and Bar-Zvi, 2010; Wang *et al.*, 2011), but not in flowering control processes.

#### Supplementary data

Supplementary data are available at JXB online.

Supplementary File 1

Figure S1. De-fruiting alters the expression of trehaloseand flavonoid-metabolizing genes.

Figure S2. Venn diagrams of developmentally regulated genes.

Figure S3. Photosynthetic genes are induced following defruiting in the bud.

Figure S4. Quantification of protein blot results (Fig. 5C).

Figure S5. Schematic representation of the cleavage of 9-*cis* xanthophylls to xanthoxin by 9-*cis*-epoxycarotenoid dioxygenase (NCED), a key regulated step in the biosynthesis of ABA in plants.

Figure S6. Changes in the expression of the *PYR1-like* gene.

Figure S7. Linear regression between transcriptomic and transcriptional (qPCR) data of Ca-related and *NPH3-like* genes, presented in Fig. 8A and B.

Figure S8. Genes encoding PINOID (PID)-binding protein 1 (PBP1) and Non-Phototropic Hypocotyl 3-like (NPH3) show homology in *Arabidopsis* and *Citrus*. **3042** | Shalom *et al*.

Table S1. List of primers used in this study.

Table S2. List of genes used to design probes for nCounter analysis (Shalom *et al.*, 2012).

Table S3. Statistical data of deep sequencing analysis.

Table S4. GO categorization of unique genes up- or downregulated in buds of OFF-Crop and de-fruited (DEF) trees relative to buds of ON-Crop trees during Time 4.

Table S5. GO categorization of genes of cluster 4 of DEF buds.

Table S6. GO categorization of up-regulated genes of OFF-Crop buds.

Table S7. GO categorization of down-regulated genes of OFF-Crop buds.

Table S8. GO categorization of genes of cluster 3 of DEF buds.

Table S9. GO categorization of genes of cluster 2 of ON-Crop buds.

Supplementary File 2

The file includes: (i) all possible comparisons between the treatments; (ii) the accession numbers of genes presented in the Venn diagrams (Fig. 4A, B; Supplementary S2A, B); (iii) photosynthesis genes (Fig. 5A); and (iv) calcium-related genes (Fig. 8A).

#### Acknowledgements

We thank Oren Ostersetzer for the contribution of antibodies, and Giora Ben-Ari for his help in the statistical analyses. This work was supported by fund number 203–0870 of the Chief Scientist of the Ministry of Agriculture and Rural Development.

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