

1 **Regulation by crop load of starch metabolism genes in leaves and roots**
2 **of Citrus.**

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12 **Running title:**

13 CROP LOAD REGULATES CARBOHYDRATE METABOLISM IN CITRUS

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26 ABSTRACT

27 The fruit is the main sink organ in *Citrus* and captures almost all available
28 photoassimilates during its development. Consequently, carbohydrate partitioning and
29 starch content depend on the crop load of *Citrus* trees. Nevertheless, little is known
30 about the mechanisms controlling the starch metabolism at the tree level in relation to
31 presence of fruit. The aim of this study was to find the relation between the seasonal
32 variation of expression and activity of the genes involved in carbon metabolism and the
33 partition and allocation of carbohydrates in ‘Salustiana’ sweet orange trees with
34 different crop loads. Metabolizable carbohydrates, and the expression and activity of the
35 enzymes involved in sucrose and starch metabolism, including sucrose transport, were
36 determined during the year in the roots and leaves of 40-year-old trees bearing heavy
37 crop loads (‘on’ trees) and trees with almost no fruits (‘off’ trees).

38 Fruit altered photoassimilate partitioning in trees. Sucrose content tended to be constant
39 in roots and leaves, and surplus fixed carbon is channeled to starch production.
40 Differences between ‘on’ and ‘off’ trees in starch content can be explained by
41 differences in ADP-glucose pyrophosphorylase (AGPP) expression/activity and α -
42 amylase activity which varies depending on crop load. The observed relation of AGPP
43 and UGPP is noteworthy and suggests a direct link between sucrose and starch
44 synthesis. Furthermore, different roles for SUT2 in leaves and roots have been
45 proposed. Variation in soluble sugars content cannot explain the differences in gene
46 expression between the ‘on’ and ‘off’ trees. A still unknown signal from fruit should be
47 responsible for this control.

48

49 **1. Introduction**

50 The amounts of carbon partitioned to different sink organs may be limited by both
51 source and sink ability to provide and utilize assimilates, respectively (Wareing and
52 Patrick, 1976). Limitations at the sink depend on organ genetic features and the
53 developmental stage, whereas source limitations may be affected by both whole plant
54 status and environmental conditions.

55 The major component of carbohydrate partitioning is the translocation of sugars from
56 photosynthetic sources to non-photosynthetic sink tissues (Slewinski and Braun, 2010).

57 In *Citrus*, and in most plants, sucrose is the main transported sugar (Zimmermann and
58 Ziegler, 1975). Diverse transport proteins and enzymes are involved in this process.

59 Phloem-localized sucrose transporters are essential for phloem loading, for maintenance
60 of phloem flux and for sucrose release in apoplastic unloaders (Sauer 2007). Other
61 enzymes, such as invertases or sucrose-phosphate synthase, allow the fine regulation of
62 sugar accumulation and distribution in the plant (Roitsch, 1999; Li et al., 2012).

63 Another component of carbohydrate partitioning is the mobilization of carbohydrate
64 reserves. Starch is the main reserve carbohydrate in plants and acts as a major integrator
65 in plant growth regulation. Marked regulatory properties have been found for ADP-
66 glucose pyrophosphorylase (AGPP), which are involved in starch biosynthesis and are
67 subjected to multilevel regulation (Geigenberger, 2011). Starch degradation occurs via a
68 network of reactions that includes amylases and debranching enzymes (Stitt and
69 Zeeman, 2012). The distribution of carbon units between starch and sucrose
70 biosynthetic pathways is tightly regulated to respond to carbon demands throughout the
71 day and night, and starch synthesis is a key process in the regulation of photoassimilate
72 partitioning and carbon allocation within the plant (Preiss, 1982; Zeeman et al., 2007).

73 In perennial plants, the carbohydrate reserves which accumulate during winter are
74 crucial for development as they supply the required energy and carbon skeletons to
75 sustain emergence and growth of new plant organs at the beginning of the growing
76 season (Naschitz et al., 2010). Under subtropical conditions, most *Citrus* trees
77 accumulate reserves during the winter rest and mobilize them during spring when the
78 main flush of bud sprouting occurs and vegetative sprouts and flowers are formed
79 (Goldschmidt and Koch, 1996). These reserves are stored mainly in roots, although high
80 concentrations can also be found in leaves and bark (Goldschmidt and Golomb, 1982).
81 After fruit set, most fixed carbon accumulates in the fruit. Both the accumulation and
82 mobilization of reserves and production of photoassimilates have been related to fruit
83 load in *Citrus* (Monerri et al., 2011).

84 Some citrus cultivars present an intense alternate bearing habit. Trees form a huge
85 number of flowers, resulting in a heavy crop load ('on' year), followed by a year with
86 very few flowers formed, or none at all ('off' year). Hormonal factors and changes in
87 carbohydrate and mineral status appear to participate in the regulation of these
88 processes (Monselise and Goldschmidt, 1982). In alternate bearing sweet orange
89 'Salustiana', the accumulation of reserves is inversely related to crop load (Monerri et
90 al., 2011), and changes in carbohydrate reserves during the year reflect variations in
91 supply and demand. Fruiting trees accumulate most fixed carbon in fruits, while no
92 accumulation is observed in roots before harvest. In the non-fruited trees, however,
93 most fixed carbon is transported to roots and utilized in growth processes, and after
94 December, stored as reserves. Reserve carbohydrate accumulation in leaves starts by
95 early December, and the levels in leaves are, until bud sprouting, the same in both the
96 'on' and 'off' trees. The heavy flower formation which follows an 'off' year causes the
97 rapid mobilization of the stored reserves, which are exhausted at full bloom.

98 Regulation of photosynthesis by fruit has been studied in *Citrus* (Iglesias et al., 2002;
99 Syvertsen et al., 2003; Nebauer et al., 2011). It is assumed that photoassimilate
100 production in leaves is modulated by the demand of sinks (Goldschmidt and Koch,
101 1996), but this effect is not always observable (Nebauer et al., 2011). It has been
102 described that the root system is a strong and unsaturable sink under cropping
103 conditions, and no enhanced photosynthetic rate by high sink strength related to fruiting
104 was found by Nebauer et al. (2013). The photosynthetic rate was similar in trees with
105 high and low crop loads in ‘Salustiana’ sweet orange (Monerri et al., 2011; Nebauer et
106 al., 2013) when differences in carbohydrate content were highest.

107 As foregoing information clearly reveals, photoassimilate production and partitioning
108 are highly integrated processes, and understanding how they are controlled will
109 underpin many targets for plant biotechnologists (Halford, 2010).

110 There are no studies that analyze the effect of fruit on the seasonal expression of
111 carbohydrate metabolism-related genes. It has been shown that the seasonal expression
112 of flowering genes is regulated by fruit (Muñoz-Fambuena et al., 2011; Shalom et al.,
113 2012), although they do not provide enough information to understand the mechanism
114 by which fruit controls the flowering process.

115 Soluble sugars, like hormones, can act as primary messengers and regulate signals that
116 control the expression of different genes involved in plant growth and metabolism
117 (Rolland et al., 2006; Rosa et al., 2009)

118 The aim of this study was to analyze the influence of fruit load on the seasonal
119 expression and activity of the genes involved in carbon metabolism, and the possible
120 role of soluble sugars as signals controlling the starch metabolism gene expression in
121 citrus trees. The studied genes were selected from previous works which reported on the
122 relation between its expression and changes in carbohydrate levels provoked by girdling

123 (Li et al., 2003a,b,c; Nebauer et al., 2011). After taking into account that field studies
124 may reveal essential roles of genes which cannot otherwise be observed, this work has
125 been carried out in non-manipulated mature trees under cropping conditions during
126 periods when the tree physiology showed distinctive characteristics. Furthermore, in
127 order to assess the effect of fruit on the regulation of the activity of the studied genes,
128 this work was performed in a citrus cultivar that presents an intense alternate bearing
129 habit.

130

131 **2. Materials and methods**

132 *2.1. Plant material*

133 Experiments were performed on 40-year-old trees of the ‘Salustiana’ cultivar of sweet
134 orange (*Citrus sinensis* [L.] Osbeck) grafted onto a Troyer citrange (*C. sinensis* [L.]
135 Osb. × *Poncirus trifoliata* Raf.) rootstock. Trees were drip-irrigated, and mineral
136 elements were supplied in the irrigation water from February to September.

137 Trees present an alternate-year bearing habit, and flowering intensity depends on the
138 fruit load of the previous year. Trees alternated between years of abundant flowering
139 and fruit set (‘on’ year) and years of almost no flowering (‘off’ year). During each year,
140 the ‘on’ and ‘off’ trees were found in the same orchard. Mature fruits were harvested by
141 early February. The ‘on’ trees averaged 3,119 fruits per tree in the study orchard during
142 the previous season, whereas only 43 fruits per tree formed in the ‘off’ trees (Y.
143 Bordón, personal communication). At the beginning of the study (March), the ‘on’
144 trees, which entered an ‘off’ year, formed only 1.6 flowers per 100 nodes, unlike the
145 54.1 flowers formed in the ‘off’ trees that entered an ‘on’ year.

146 Sampling dates for determinations of carbohydrates, enzymatic activity and gene
147 expression were performed based on previous studies (Monerri et al., 2011): June, after

148 fruit abscission, when the maximum rate of accumulation by the fruit occurred;
149 September and December, in the middle and final period of fruit development,
150 respectively; January and February, just before and after fruit harvest, respectively; and
151 March, after the beginning of Spring bud sprouting. Plant material was sampled
152 between 10:00 h and 11:00 h on all six dates. The mature leaves (4th leaf from the apex)
153 from vegetative shoots formed last Spring and the fibrous roots (1.5-2.5 mm in
154 diameter) bearing new formed feeder roots were used in the study.

155

156 *2.2. Carbohydrate analysis*

157 The determination of total soluble sugars and starch (as mg per g of dry weight) was
158 performed as described by García-Luis et al. (2002). Three independent extracts, each
159 obtained from nine different trees (five leaves per tree and three trees per extract), were
160 assayed for each treatment in all the determinations. Sucrose was determined by HPLC,
161 as described by Iglesias et al. (2002).

162

163 *2.3. Gene expression analysis*

164 The expression of sucrose transporters SUT1 and SUT2 (Li et al., 2003c), sucrose
165 synthases SUS1 and SUSA, sucrose-phosphate synthase (SPS, EC 2.4.1.14), α -amylase
166 (AMY, EC 3.2.1.1) and ADP-glucose pyrophosphorylase (AGPP) genes (Li et al.
167 2003a), involved in carbohydrate metabolism, were studied (Table 1). Leaf tissue was
168 finely ground in liquid nitrogen and total RNA was extracted using the TRIzol reagent
169 (Invitrogen), purified using the RNEasy Mini Kit (Quiagen) and treated with RNase-
170 free DNase (Quiagen), according to the manufacturer's instructions. RNA was
171 quantified with a UV/VIS spectrophotometer, and first-strand cDNA was synthesized

172 from 1.2 µg of total RNA with the First Strand cDNA Synthesis Kit AMV (Roche) for
173 real-time PCR (RT-PCR).

174 The oligonucleotide primers used have been described in a previous work (Nebauer et
175 al. 2011). During the year, *Citrus sinensis* glyceraldehyde-3-phosphate dehydrogenase
176 (GAPDH)(Nebauer et al., 2011) exhibited a stable expression among the studied organs
177 and was used as the reference gene. The optimum concentration and amplification
178 efficiency were tested for all pairs of oligonucleotides (Livak and Schmittgen, 2001).

179 Diluted cDNA (2 µg) was used as a template for the semi-quantitative RT-PCR
180 amplification in the 20-µL reactions containing 0.3 µM of each primer (0.15 µM
181 GAPDH) and 10 µL of the SYBR Green PCR master mixture (Power SYBR[®]Green
182 PCR Master Mix; Applied Biosystems). The PCR mixtures were preheated at 50°C for
183 2 min and then at 95°C for 10 min, followed by 40 amplification cycles (95°C for 15 s;
184 60°C for 1 min). Amplification specificity was verified by a final dissociation (95°C for
185 15 s, 60°C for 20 s and 95°C for 15 s) of the PCR products. The levels of the PCR
186 products were monitored with an ABI PRISM 7000 sequence detection system and
187 were analyzed with the ABI PRISM 7000 SDS software (Applied Biosystems). At least
188 three independent biological replicates per sample and three technical replicates of each
189 biological replicate were used for the RT-PCR analysis. The relative expression levels
190 of the target genes were calculated by the $2^{-\Delta\Delta CT}$ method (Livak and Schmittgen, 2001).
191 For each gene and organ, the expression was related to the minimal value of the
192 measured dates.

193

194 2.4. Enzyme assays

195 One gram of frozen powder was resuspended at 4°C in 5 mL of 100 mM HEPES (pH
196 7.5), 2 mM EDTA and 5 mM dithiothreitol. The suspension was desalted (IVSS

197 Vivaspin 500, Sartorius Biolab, Germany) following the manufacturer's instructions
198 and assayed for enzymatic activity. The ADPG pyrophosphorylase (AGPP, EC
199 2.4.1.18), starch phosphorylase (SP, EC 3.6.1.1), UDPG pyrophosphorylase (UGPP, EC
200 2.7.7.9), sucrose synthase (SuSy, EC 2.4.1.13) and acid invertase (INV, EC 3.2.1.26)
201 activities were assayed (Table 1) as described by Baroja-Fernández et al. (2004) and
202 Muñoz et al. (2005). For the detection of the AGPP and UGPP activities, the production
203 of glucose-1-phosphate from ADP-glucose and UDP-glucose was determined,
204 respectively, in an NAD-linked glucose-6-phosphate dehydrogenase system (Müller-
205 Roeber et al., 1992). NAD reduction was measured spectrophotometrically at 340 nm.
206 Starch phosphorylase activity was assayed by measuring the glucose-1-phosphate
207 released from glycogen in a similar assay. The sucrose synthase and invertase activities
208 were measured in the sucrose breakdown direction. Fructose content was determined
209 spectrophotometrically at 340 nm by the NAD-linked
210 hexokinase/phosphoglucoisomerase/glucose-6-phosphate dehydrogenase coupling
211 method. All the enzymatic reactions were performed at 37°C. One unit (U) is defined as
212 the amount of enzyme that catalyzes the production of 1 µmol of product per min.

213

214 2.5. *Statistical analysis*

215 Treatment comparison analyses were performed by ANOVA (Statgraphics Plus 5.1 for
216 Windows, Statistical Graphics Corp.). Mean separations were made with the Tukey
217 multiple range test. A linear regression analysis was used to evaluate the relationships
218 between parameters.

219

220 **3. Results**

221 3.1. *Carbohydrate content in leaves and roots*

222 The carbohydrate content in leaves from the vegetative sprouts formed during spring in
223 study year 1 and in roots were examined during the fruit development period, from June
224 to January, which ended in March just after the beginning of the spring flush of study
225 year 2 (Fig. 1).

226 Starch content was significantly higher in the leaves of vegetative sprouts in the ‘off’
227 trees than in the ‘on’ trees (Fig. 1A). Differences were maximal in June. Afterwards,
228 this content decreased gradually to a common minimum level in both the ‘on’ and ‘off’
229 trees during December. From this time point, starch accumulated until the beginning of
230 bud sprouting in the two tree types to lower again in the ‘on’ trees by June.

231 Almost no differences in leaves between the ‘on’ and ‘off’ trees were observed in either
232 sucrose content or total soluble sugars (Fig. 1C and 1E), which remained nearly
233 constant during the study period. However, a significant increase in total soluble sugars
234 and sucrose occurred in January.

235 No differences in the starch content of the roots between the ‘on’ and ‘off’ trees were
236 observed until November (Fig. 1B). Afterwards, starch accumulated in the roots until
237 the beginning of bud sprouting. The accumulation rate was higher in the ‘off’ trees. The
238 soluble sugar and sucrose content in roots showed a similar behaviour as in leaves (Fig.
239 1C and 1D).

240

241 *3.2. Effect of crop load on gene expression*

242 The expression pattern of the genes involved in starch metabolism, sucrose transport
243 and sucrose metabolism in the ‘on’ and ‘off’ trees is shown in Figures 2, 3 and 4,
244 respectively.

245 In leaves, the expression of AGPP decreased from June to November in the ‘off’ trees,
246 but rose from December to February (Fig. 2A). After spring flush had started, the

247 expression levels fell again. In the ‘on’ trees, AGPP showed a similar behaviour,
248 although the starting level in June was significantly lower, while the winter raise was
249 observed in November, earlier than in the ‘off’ trees (Fig. 2A). In addition, almost no
250 changes were observed in the expression of the AMY gene from June to January in the
251 ‘off’ leaves (Fig. 2C). From that time onwards, a sharp increase occurred until March. A
252 similar trend was observed in the ‘on’ trees despite the higher expression value in June.
253 Very few or no differences were observed between the ‘on’ and ‘off’ trees in the
254 expressions of AGPP and AMY in roots (Fig. 2B and D). The expression of AGPP
255 remained low and nearly constant until December, and a slight increase was observed
256 afterwards. The AMY expression progressively decreased from June to December,
257 followed by a slight increase from February. This increase was more pronounced in the
258 ‘off’ trees (these being the ‘on’ trees in the previous year) than in the ‘on’ ones (Fig.
259 2D).

260 Sucrose transporters SUT1 and SUT2 showed different expression profiles during the
261 year (Fig. 3). The expression of SUT1 fell from June to September in leaves (Fig. 3A).
262 From then onwards, it remained virtually unchanged in the ‘off’ trees, although a slight
263 increase was observed from March. In the ‘on’ trees, a transient increase was observed
264 in January. The SUT2 expression in leaves was significantly higher in the ‘off’ trees in
265 June. Both these levels in the ‘on’ and ‘off’ trees decreased to a minimum in September,
266 and no changes were observed until January, when an increase took place (Fig. 3C).

267 In roots, the SUT1 expression differed between both tree types. Practically no changes
268 were seen in the expression of this gene from June to December in the ‘off’ trees, which
269 fell from this time onwards. However, its expression was lower in the ‘on’ trees during
270 September. The SUT2 levels did not change until December, and a slight increase was
271 observed in both the ‘on’ and ‘off’ trees from January onwards (Fig. 3D).

272 The expression of the SUS1 gene in leaves oscillated during fruit development with
273 differences found between the tree types (Fig. 4A). These changes were more
274 pronounced in the ‘off’ trees, with a higher expression in early summer and January. In
275 contrast, almost no changes were noted in SUSA (Fig. 4C). Despite being higher in the
276 ‘on’ leaves until September, the SPS expression in both the ‘on’ and ‘off’ trees
277 decreased until January to rise afterwards at the same level in both trees (Fig. 4E).
278 The SUS1 expression in roots did not change in the ‘on’ and ‘off’ trees during the study
279 period (Fig. 4B). Practically no changes were observed in the SUSA expression until
280 February, when it increased in both the ‘on’ and ‘off’ trees (Fig. 4D). The SPS
281 expression in roots fell in November, recovered in January, and decreased after the fruit
282 harvest in February (Fig. 4F). Except for June, no differences were observed between
283 the ‘on’ and ‘off’ trees.

284

285 *3.3. Effect of crop load on enzyme activity*

286 The activity of enzymes related to starch and sucrose metabolism are presented in
287 Figures 5 and 6. AGPP activity was higher in ‘off’ tree leaves than in the ‘on’ trees until
288 September (Fig. 5A), after which time it decreased until February, but recovered in
289 March. SP activity was also higher in the leaves of ‘off’ trees in June, but similar from
290 September to March in both tree types (Fig. 5C).

291 In roots, AGPP activity was very low during the study period, although a slight increase
292 occurred from November (Fig. 5B). SP activity increased slowly and progressively in
293 the ‘on’ trees (Fig. 5D). This increase was delayed until September in the ‘off’ trees,
294 although higher levels were reached from November as compared to the ‘on’ trees.

295 A similar trend of UGPP activity was observed in the leaves of both the ‘on’ and ‘off’
296 trees (Fig. 6A), which was initially higher in June in the ‘off’ trees, and equalled as

297 from September, decreased until February and increased afterwards. Susy activity in
298 leaves was very low during the study period (Fig. 6C). Nevertheless, a transient strong
299 increase was observed in February in the ‘off’ trees. INV activity increased from
300 September to January, and then progressively decreased in both the ‘on’ and ‘off’ trees
301 (Fig. 6E).

302 In roots, UGPP activity increased at the beginning of the study period (Fig. 6B), and
303 decreased from January in the ‘on’ trees and from February in the ‘off’ trees. Susy
304 activity progressively increased with time to peak in February (Fig. 6D). In the ‘off’
305 trees, a transient decrease was observed in September. INV activity remained nearly
306 constant and at low rates (Fig. 6F), despite a transient maximum in recorded September
307 in the ‘off’ trees.

308

309 *3.4. Relations among carbohydrate content, enzyme activities and gene expression*

310 The relations between carbohydrate contents in leaves and roots and the expression and
311 activity of related enzymes and transporters were studied. The main significant relations
312 are schematically illustrated in Figure 7. The higher starch levels in leaves during
313 summer and in roots during winter observed in the ‘off’ trees (Fig. 1A and 1B)
314 correlated with a higher AGPP expression ($r^2 = 0.80$; $P = 0.01$) and greater activity ($r^2 =$
315 0.62 , $P = 0.03$). Furthermore, AGPP and UGPP activities were highly related in leaves
316 (Fig. 7A). The high correlation between starch content and the expression of both
317 sucrose transporters SUT1 ($r^2 = 0.84$; $P = 0.04$) and SUT2 ($r^2 = 0.89$; $P = 0.02$) is
318 noteworthy. Leaf INV activity related negatively to starch ($r^2 = - 0.81$; $P = 0.04$), but
319 positively to soluble sugar ($r^2 = 0.67$; $P = 0.02$) content. Soluble sugar content related
320 negatively to the SPS ($r^2 = - 0.77$; $P = 0.01$) and SUSA ($r^2 = - 0.60$; $P = 0.04$)
321 expression levels (Fig. 7A), due mainly to sugars other than sucrose (Fig. 1C and 1E).

322 In roots, similar relations were observed between starch and the AGPP expression and
323 activity (Fig. 7B). Changes in the AGPP expression also related to changes in the SUT2
324 ($r^2 = 0.97$; $P = 0.00$) and SUSA ($r^2 = 0.70$; $P = 0.02$) expression levels. Sucrose
325 synthase and invertase activities related to AGPP activity ($r^2 = 0.58$; $P = 0.05$, and $r^2 = -$
326 0.59 ; $P = 0.05$, respectively) (Fig. 7). Soluble sugars related positively to the SUT2
327 expression ($r^2 = 0.64$; $P = 0.03$) and negatively to AMY expression ($r^2 = - 0.57$; $P =$
328 0.05).

329

330 **4. Discussion**

331 Crop load is known to affect carbohydrate production and partitioning in several trees,
332 such as apple (Naschitz et al., 2010), olive (Bustan et al., 2011) and citrus (Goldschmidt
333 and Golomb, 1982; Monerri et al., 2011). During its development, citrus fruit is the
334 main sink organ (Monerri et al., 2011), and it captures almost all available
335 photoassimilates. Accordingly, differences in carbohydrate content and related enzyme
336 activities throughout seasons between the ‘on’ and ‘off’ trees are reported in our study.
337 This different behaviour was observed mainly from May to September in leaves, and
338 from December to March in roots, when higher starch levels were found in non-fruiting
339 trees. This finding suggests a role of fruit in the regulation of the genes relating to the
340 metabolism of this reserve carbohydrate.

341 The higher starch level noted in leaves from June to September in the ‘off’ trees can be
342 explained by a higher gene expression, greater AGPP activity, and a lower expression of
343 the α -amylase and sucrose phosphate synthase genes. Furthermore, the increased leaf
344 starch content correlates with not only AGPP activity, but also with the expression of
345 sucrose transporters. These results, as previously reported (Li et al., 2003c), suggest
346 different physiological roles for these transporters.

347 SUT1 has been described to drive sucrose loading in sources. Accordingly, the
348 expression of this transporter is enhanced under the high photoassimilate availability
349 and demand conditions of June. The use of dry matter by fruit in the 'on' trees and by
350 vegetative growth, mainly the root system, in the 'off' trees in June (Goldschmidt and
351 Golomb, 1982; Monerri et al., 2011) could explain this result. However, the less
352 demand in the 'off' trees during winter and, to a lesser extent in the 'on' trees, provoked
353 increased starch synthesis. Starch content and AGPP expression correlated highly with
354 the SUT2 expression in leaves. These results support the hypothesis that the SUT2
355 protein may act as a sugar sensor (Barker et al., 2000).

356 In 'Salustiana' sweet orange, no differences were observed in the photosynthetic rate
357 between the fruiting and non-fruiting 'Salustiana' trees throughout the year (Monerri et
358 al., 2011; Nebauer et al., 2013). Therefore, similar photoassimilate production at the
359 tree level has to be assigned to the 'on' and 'off' trees as similar total leaf area and
360 photosynthetic capacity have been estimated in both tree types (Monerri et al., 2011).
361 Although photoassimilate synthesis is similar between trees, but with differing demand,
362 our data reveal that sucrose content tends to be maintained more or less constant in
363 leaves in the 'off' trees by channeling the surplus fixed carbon to starch production, and
364 to fruit in the 'on' trees. In line with this, a high correlation is seen between AGPP and
365 UGPP activities in leaves, suggesting the connection via hexoses as proposed by Muñoz
366 et al. (2006).

367 No differences were observed in the soluble sugar content between the 'on and 'off'
368 trees, although an increase took place in January. The highest content of soluble sugars
369 in leaves correlates with the lowest starch accumulation, which is due mainly to an
370 increase in hexoses (data not shown). The higher sink strength of leaves during this
371 period coincides with higher invertase and diminished Susy activity. It has been stated

372 that their relative activities determine how much carbon enters the storage pathways for
373 starch biosynthesis, and how much enters the glycolytic pathway (Halford, 2010). Some
374 studies have demonstrated that Susy activity is closely related to starch accumulation
375 and invertase is associated with glucose and fructose production, principally for flux
376 into glycolysis (Trethewey et al., 1998). However, the increase in soluble sugars
377 towards mid-winter in *Citrus* was observed long before (Jones and Steinacker, 1951;
378 Toritaka et al., 1974) and has been related to the role of soluble sugars as an osmotic,
379 cryoprotective strategy against cold injury.

380 The rise in soluble sugars, other than sucrose mainly, is also observed in roots. Unlike
381 leaves however, this higher content correlates with increased starch content. The
382 accumulation of reserves in roots occurs from December onwards in both the 'on' and
383 'off' trees, which coincides with the lower sink strength of the 'on' trees fruit.
384 Nevertheless, starch content is higher in the roots of the 'off' trees and correlates with
385 root AGPP activity. The soluble sugar level correlates with both sucrose synthase
386 activity and the SUT2 expression (Fig. 1B, 3D and 6D). The role of SUT2 as a
387 transporter in sink organs has been previously described in *Citrus* (Li et al., 2003b,c).

388 A significant correlation between the expression levels of a member of a gene family
389 and total activity has been proposed to be related to the transcriptional regulation of the
390 enzyme activity (Li et al., 2012). However, the fact that these correlations are lacking
391 suggests that the post-translational regulation of the protein might regulate its activity or
392 that another family member may play a predominant role in total activity.

393 The AGPP expression in leaves, which explains the differences in starch accumulation
394 between the 'on' and 'off' trees, is well-related to AGPP activity, thus indicating its
395 mainly transcriptional regulation. Besides, the differences in root starch content
396 correlate with the activities of those enzymes involved in starch synthesis. Nonetheless,

397 the AGPP expression shows no differences between the ‘on’ and ‘off’ trees, suggesting
398 additional levels of regulation.

399 It has been hypothesized that soluble sugars modulate the expression of those genes
400 involved in starch synthesis (Koch, 1996). However, we observed no differences in
401 soluble sugars between the ‘on’ and ‘off’ trees, and sucrose content remained nearly
402 constant throughout the study period. Apparently the absolute levels of sugars do not
403 drive the regulation of the differential gene expression between the ‘on’ and ‘off’ trees.
404 However, this control may also be exerted by different phytohormones produced by
405 fruit, whose participation in the regulation of many carbon metabolism-related activities
406 is well-known (Albacete et al., 2008). GAs enhance sucrose formation, activates SPS
407 activity, phloem loading and unloading, and increases sink strength through activating
408 invertase activity (Iqbal et al., 2011). It has been reported that GAs interacts with other
409 phytohormones, such as ABA or salicylic acid, to regulate carbon allocation and
410 distribution (Moreno et al., 2011). Furthermore, Peng et al. (2011) described that ABA
411 regulates SUT1 activity in apple by stimulating sugar accumulation in fruit. A previous
412 work (Nebauer et al., 2011) found significant differences in the expression of the
413 enzymes analyzed in this manuscript in ‘Salustiana’ sweet orange between the shoots
414 bearing fruit and those without, thus confirming that the signals generated by fruit may
415 regulate the carbohydrate metabolism in trees. It has been recently reported that fruit
416 inhibits flowering by repressing the expression of flowering genes in leaves of alternate
417 bearing *Citrus* (Muñoz-Fambuena et al., 2011). The specific role of phytohormones in
418 all these regulations has to be further studied.

419 There are no differences in the soluble sugar content between the ‘off’ and ‘on’ trees
420 that explain the differences observed in the starch-metabolism gene expression.
421 Nevertheless, there is a strong relation between variation in the soluble sugar content

422 throughout the year and the activity of these genes. The changes in soluble sugar
423 content and the AGPP and SUT2 expressions correlate highly in roots, suggesting that
424 the expression of these genes may be modulated by hexoses, as hypothesized by Koch
425 (1996). However, these carbon metabolism-related activities are under complex spatial
426 and temporal regulation (Kleczkowski et al., 2010), and nothing is known about
427 whether there being a common mechanism responsible for differential sugar regulation
428 (Rosa et al. 2009). In fact, distinct relations between gene expressions in accordance
429 with tissues, stress conditions and light rhythms have been reported (Kleczkowski et al.,
430 2009). Accordingly, a negative correlation is found between soluble sugar content and
431 the expression of SUT2 and AGPP in leaves.

432 Although the expression of the carbon metabolism-related genes has been previously
433 studied in relation to crop load and carbon status in *Citrus* (Komatsu et al., 2002; Li et
434 al., 2003a,b,c), these works were neither carried out under natural field conditions nor
435 throughout the year to cover all developmental stages of a tree. One important factor is
436 that growing plants in greenhouses or growth chambers may not represent an optimal
437 environment for functional studies (Kleczkowski et al., 2010). The evaluation of the
438 roles of each gene/isozyme should include field trials conducted under natural
439 conditions, as is the case in this work. In addition, the used techniques have allowed the
440 study into the relation between the expression patterns of carbon metabolism genes with
441 variation in carbohydrate content along the year.

442

443 **5. Conclusion**

444 Our data indicate the complexity of the carbohydrate metabolism network in *Citrus* by
445 integrating source-sink interactions and environmental conditions, mediated by sugar
446 signals, and probably by hormones as well. Differences in the starch content between

447 the 'on' and 'off' trees can be explained by the differential expression/activity of AGPP
448 and α -AMY. Different regulation (transcriptional and posttranscriptional) levels for
449 leaves and roots are revealed for AGPP. Significant linear correlations are found
450 between the AGPP expression or activity and other starch metabolism-related genes.
451 The relation with UDPG is of special interest as it links sucrose and starch synthesis,
452 while the relation with SUT2 transporter suggests that SUT2 may act as a sugar sensor
453 in leaves and as a sucrose transporter to sink organs in roots. The control exerted by
454 fruit of the genes related to starch metabolism is not mediated through changes in the
455 content of soluble sugars as primary messengers, and a hormonal signal should be
456 responsible for this regulation. Nevertheless, a strong relation exists between variation
457 in soluble sugar content throughout the year and the AGPP expression. In addition,
458 differences between sources and sinks are observed. In roots, the soluble sugars
459 variation pattern runs in parallel with the AGPP and SUT2 expressions. However, a
460 negative correlation is found between AGPP activity and the SUT2 expression in
461 leaves.

462

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470

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477

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616 **Legends for Figures**

617

618 **Fig. 1.** Seasonal pattern of starch (A,B), soluble sugars (C,D) and sucrose (E,F) content
619 in the leaves (A,C,E) and roots (B,D,F) in the ‘on’(●) and ‘off’(○) Salustiana trees.
620 Values are mean (\pm SE) of three determinations in nine different trees performed from
621 June (Jn) to March (Ma). Significant differences ($P<0.05$) between trees for each date
622 are indicated by an asterisk.

623

624 **Fig. 2.** Changes in the relative expression of ADP-glucose pyrophosphorylase (A,B)
625 and α -amylase genes (C,D) in the leaves (A,C) and roots (B,D) of the ‘on’ (●) and ‘off’
626 (○) Salustiana trees. Values are mean (\pm SE) of three determinations in nine different
627 trees performed from June (Jn) to March (Ma). Significant differences ($P<0.05$)
628 between trees for each date are indicated by an asterisk.

629

630 **Fig. 3.** Changes in the relative expression of SUT1 (A,B) and SUT2 (C,D) sucrose
631 transporter genes in the leaves (A,C) and roots (B,D) of the ‘on’ (●) and ‘off’ (○)
632 Salustiana trees. Values are mean (\pm SE) of three determinations in nine different trees
633 performed from June (Jn) to March (Ma). Significant differences ($P<0.05$) between
634 trees for each date are indicated by an asterisk.

635

636 **Fig. 4.** Changes in the relative expression of sucrose synthase 1 (A,B), sucrose synthase
637 A (C,D) and sucrose phosphate synthase (E,F) genes in the leaves (A,C,E) and roots
638 (B,D,F) of the ‘on’ (●) and ‘off’ (○) Salustiana trees. Values are mean (\pm SE) of three
639 determinations in nine different trees performed from June (Jn) to March (Ma).

640 Significant differences ($P<0.05$) between trees for each date are indicated by an
641 asterisk. nd: not determined.

642

643 **Fig. 5.** Changes in the ADPG pyrophosphorylase (A,B) and starch phosphorylase (C,D)
644 activities in the leaves (A,C) and roots (B,D) of the ‘on’ (●) and ‘off’ (○) Salustiana
645 trees. Values are mean (\pm SE) of three determinations in nine different trees performed
646 from June (Jn) to March (Ma). Significant differences ($P<0.05$) between trees for each
647 date are indicated by an asterisk.

648

649 **Fig. 6.** Changes in the UDPG pyrophosphorylase (A,B), sucrose synthase (C,D) and
650 invertase (E,F) activities in the leaves (A,C,E) and roots (B,D,F) of the ‘on’ (●) and
651 ‘off’ (○) Salustiana trees. Values are mean (\pm SE) of three determinations in nine
652 different trees performed from June (Jn) to March (Ma). Significant differences
653 ($P<0.05$) between trees for each date are indicated by an asterisk.

654

655 **Fig. 7.** Main significant relations ($P<0.05$) among carbohydrates and related enzyme
656 expression and activities in Salustiana leaves (A) and roots (B). +: positive correlations,
657 -; negative correlations

658

659

Table 1. Nomenclature and reactions catalyzed by the studied enzymes.

Starch metabolism		
AGPP	ADP-glucose pyrophosphorylase	$\text{glucose-1-P} + \text{ATP} \rightarrow \text{ADP-glucose} + \text{PPi}$
AMY	α -amylase	$[\text{glucose}]_n \rightarrow [\text{glucose}]_{n-m} + [\text{glucose}]_m$
SP	starch phosphorylase	$[\text{glucose}]_n + \text{Pi} \leftrightarrow \text{glucose-1-P} + [\text{glucose}]_{n-1}$
Sucrose metabolism		
UGPP	UDP-glucose pyrophosphorylase	$\text{glucose-1-P} + \text{UTP} \rightarrow \text{UDP-glucose} + \text{PPi}$
SUS/SuSy	Sucrose synthase	$\text{sucrose} + \text{ADP} \leftrightarrow \text{ADP-glucose} + \text{fructose}$
INV	Invertase	$\text{sucrose} \rightarrow \text{glucose} + \text{fructose}$
SPS	Sucrose-phosphate synthase	$\text{UDP-glucose} + \text{fructose-6-P} \rightarrow \text{UDP} + \text{sucrose-6-P}$
SUT	Sucrose transporter	$\text{H}^+/\text{sucrose}$ symporter

Abbreviations: fructose-6-P: fructose-6-phosphate; glucose-1-P: glucose-1-phosphate; Pi: phosphate; PPi: pyrophosphate; sucrose-6-P: sucrose-6-phosphate

Figure 1

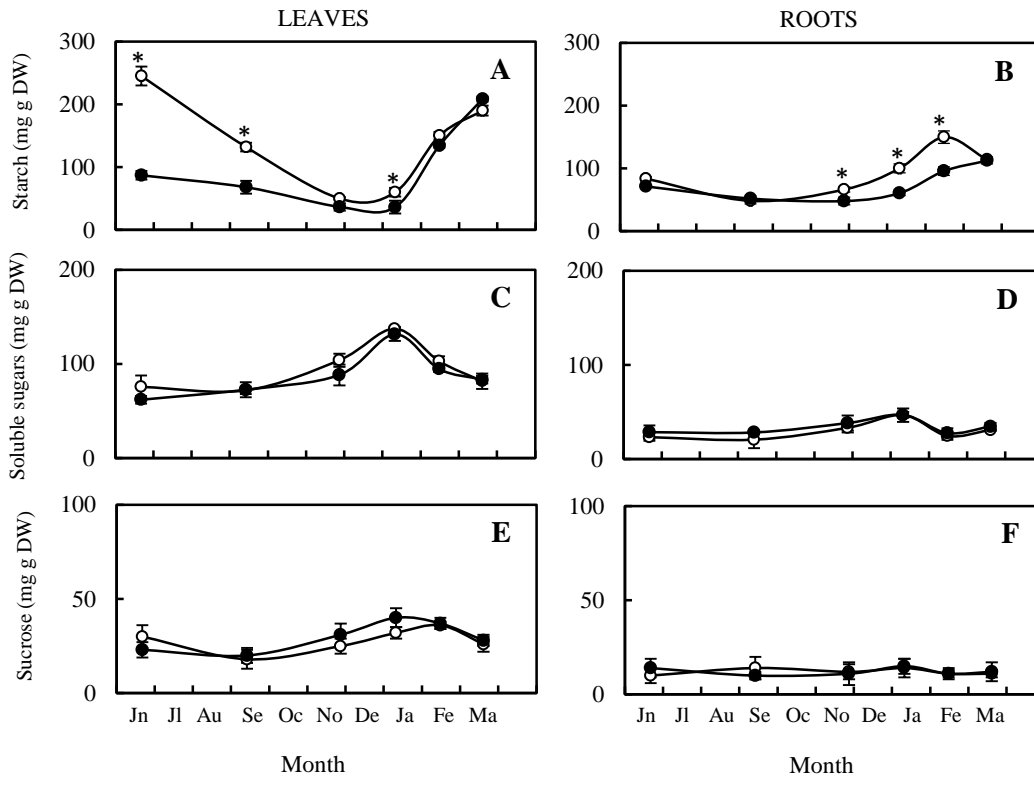


Figure 2

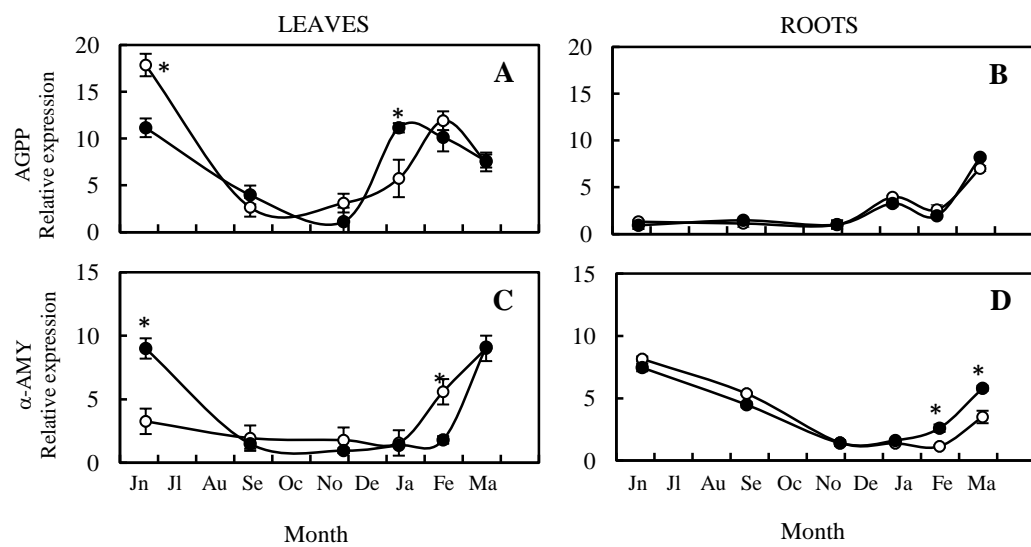


Figure 3

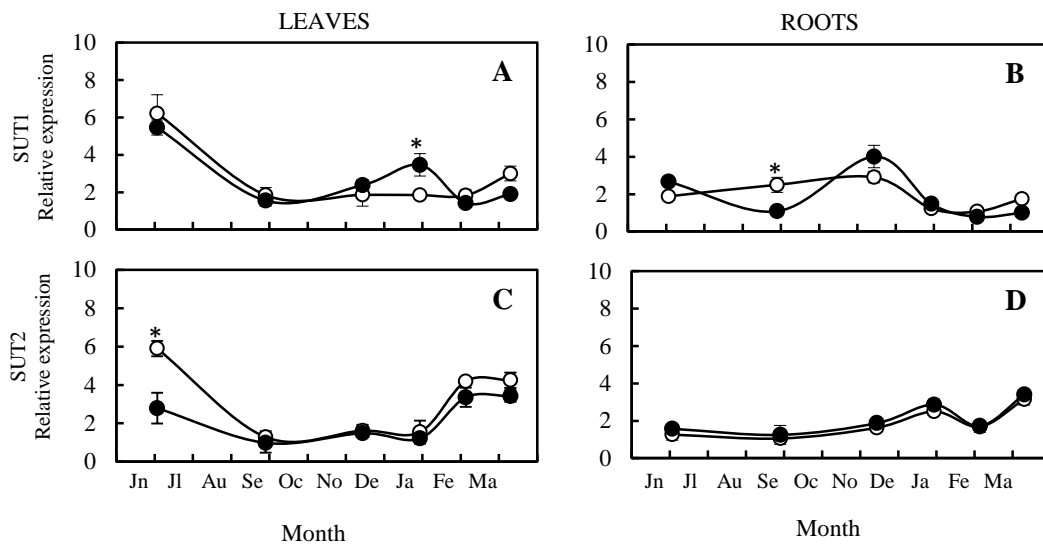


Figure 4

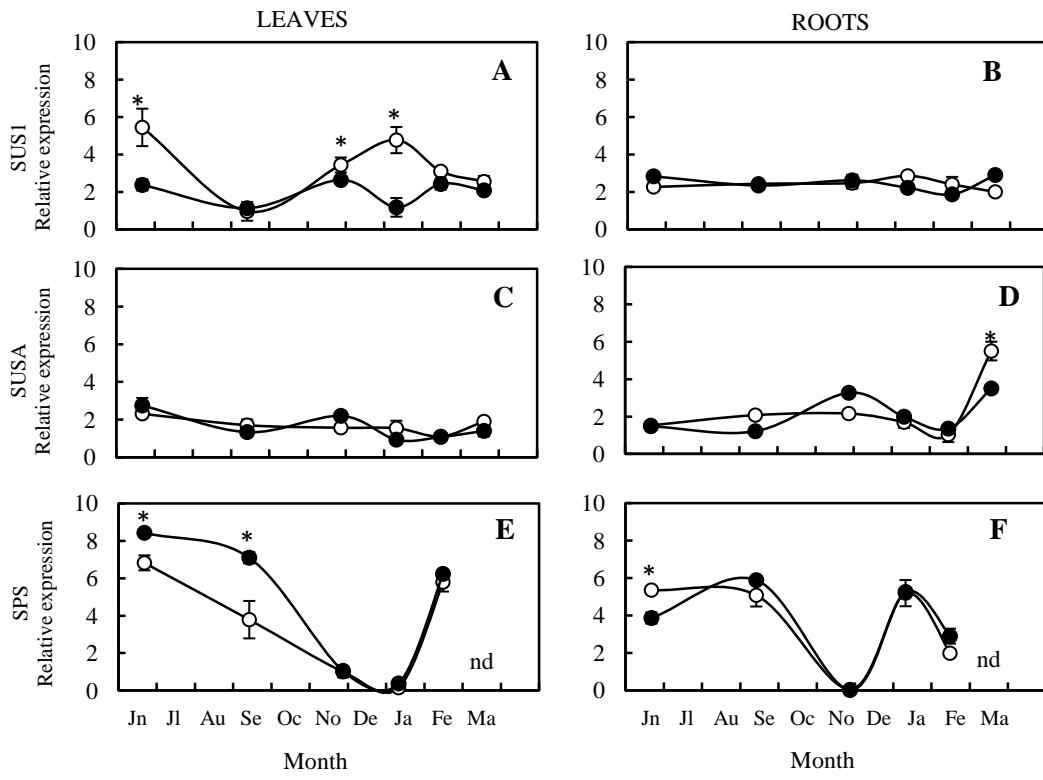


Figure 5

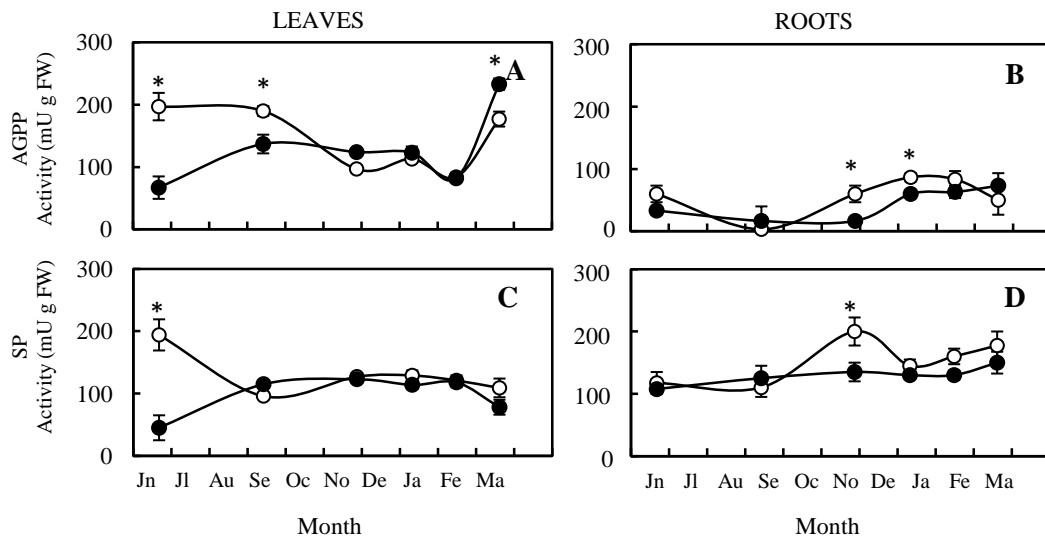


Figure 6

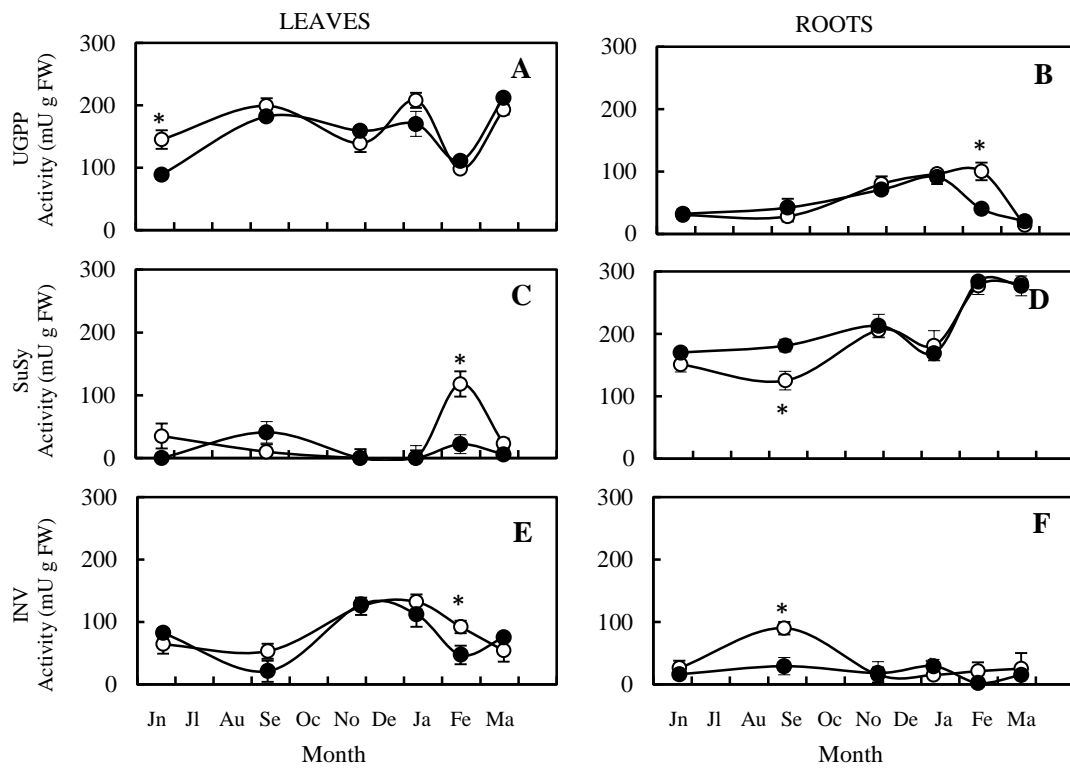


Figure 7

