Preliminary Study on Effect of Early Defoliation on Dry Matter Accumulation and Storage of Reserves on ‘Abbé Fétel’ Pear Trees

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Additional index words. crop-load, dormancy stage, Pyrus communis, soluble carbohydrates, starch

Abstract. Annual accumulation of starch is affected by carbon reserves stored in the organs during the growing season and is controlled mainly by sink strength gradients within the tree. However, unfavorable environmental conditions (e.g., hail events) or application of management practices (e.g., defoliation to enhance overcolor in bicolor apple) could influence the allocation of storage carbohydrates. This preliminary research was conducted to determine the effects of early defoliation on the dry matter, starch, and soluble carbohydrate dynamics in woody organs, roots, and mixed buds classified by age and two levels of crop-load for one growing season in ‘Abbé Fétel’ pear trees (Oct. 2012 to mid-Jan. 2013 in the northern hemisphere). Regardless of the organs evaluated (woody organs, roots, and mixed buds), an increase of soluble carbohydrate concentration was observed in these organs in the period between after harvest (October) and January (dormancy period). Among all organs, woody short-old spurs showed the highest increase (+93.5%) in soluble sugars. With respect to starch, woody organs showed a clear trend of decreasing in concentration between October and January. In this case, short-old spurs showed the smallest decline in starch concentrations, only 6.5%, whereas in other tree organs starch decreased by 34.5%. After harvest (October), leaves showed substantially higher starch and soluble sugar concentrations in trees with lower crop-loads. These results confirm that in the period between October and January, dynamic interconversions between starch and soluble carbohydrates occur at varying magnitudes among organs in pear trees.

Annual accumulation, mobilization, and allocation of carbohydrates in fruit trees among individual organs is greatly affected by the availability of stored carbon reserves, assimilates from photosynthesis, and crop-load within a tree (Keller and Loescher, 1989; Monerri et al., 2011). Ultimately, the final accumulation of carbohydrates in tree organs is determined by their sink strength (Ho, 1988). Storage carbohydrates sustain the early stages of growth in spring, during budbreak and leaf growth until leaves have developed the photosynthetic capacity to independently support net carbon assimilation (Flores and Layne, 1999; Regier et al., 2010; Whiley et al., 1996a). During the dormant period (fall-winter), hydrolysis of starch and storage proteins occurs (Grocchowska, 1973; Titus and Kang, 1982), supporting bud development (Keller and Loescher, 1989; Loescher et al., 1990) in the form of nucleic acid and protein synthesis (Zimmerman et al., 1970).
Alternate bearing is a phenomenon that affects both deciduous and evergreen trees. In avocado (Scholefield et al., 1985; Whiley et al., 1996a, 1996b), pistachio (Nizama et al., 1997), and citrus (Goldschmidt and Golomb, 1982), the likelihood of showing alternate bearing tendency appears to be closely related to the amount of carbohydrates stored between the postharvest period and before the dormancy break. It has been suggested that unfavorable growing conditions, such as severe water stress (Loescher et al., 1990; Lopez et al., 2007), or management practices, such as a high crop-load in the previous season (Park, 2011), could affect the accumulation of carbohydrate reserves in roots, and thereby affect the yield in the following season as a result of a decrease in available assimilates. Whiley et al. (1996b) stated that the accumulation of carbohydrate reserves in avocado trees would be larger in a year of low crop-load in comparison with a high crop-load year, and as a consequence high yields would be obtained because of greater availability of assimilates accumulated during the previous year.

The performance of future cropping is affected mainly by alteration of assimilate reserves in the tree caused by abiotic and biotic stress at the end of the season as well as management practices that increase light penetration (Loescher et al., 1990; Kozlowski, 1992). Palmer (1999), Robinson and Lakslo (1991), and Wünsche et al. (1996) stated that there is a positive relationship between dry matter production and the total amount of solar radiation intercepted by the orchard. Moreover, Robinson and Lakslo (1991) pointed out that these two variables depend on factors such as canopy characteristics and their spacing. Rom (1991) described the importance of light on different physiological and morphological aspects, as well as its influence on management practices, such as pruning. Chalmers and Van den Ende (1975) suggested that the age or size at which the tree optimizes dry matter partitioning to fruit may change with orchard design or management. Cruz-Castillo et al. (2010) demonstrated in kiwifruit that by reducing vigor and exposing leaves and fruits to sunlight by early defoliation 5 days after full bloom and throughout the season can cause a poor return bloom the following year because of the rapid depletion of nonstructural carbohydrates in the previous summer. Adverse environmental factors (e.g., hailstorm or high temperature and humidity often in tropical and subtropical regions) could produce bacterial and fungal disease resulting in early natural defoliation as Lloyd and Firth (1990) reported for peach trees. Early defoliation (3–4 weeks before harvest) is also becoming more spread in biologic apple genotypes to improve the percentage of overcolored surface of the fruits. Specialized orchard equipment made in Europe can perform tree defoliation in a fast way using an air compressor jet stream. Loescher et al. (1990) noted that treatments like defoliation or pruning performed in the orchard affect mainly roots in terms of depletion of stored carbohydrates. In one of the first studies on carbohydrate reserve dynamics in pear trees, Cameron (1923) indicated that nonpruned trees began to store starch earlier than the pruned trees because of a major presence of young spurs. Further, summer pruning practice may redirect the distribution of carbohydrates to fruit as a consequence of the removal of vegetative sinks (Loescher et al., 1990). In this way, all orchard practices aiming to maintain an optimal leaf area and delay leaf senescence could allow a greater carbohydrate assimilation after harvest, resulting in a sufficient/optimal accumulation of carbohydrate reserves to support initial growth and development in the following season (Kozlowski, 1992; Tustin et al., 1997).

In the present work, we evaluated starch and soluble sugar dynamics in relation to crop-load in ‘Abbé Fétel’ pear trees. In addition, we manipulated the reserve accumulation by an early defoliation (October) of the tree after harvest to investigate the effect of this practice on dry matter, starch, and soluble carbohydrates in woody organs, roots, and mixed buds. This research provides preliminary results to understand the effect of leaf removal on carbohydrate reserve accumulation across woody organs.

Materials and Methods

Plant material

This study was conducted during one growing season (from Oct. 2012 to Jan. 2013) in an orchard established in 1997 of ‘Abbé Fétel’ (Pyrus communis L.) grafted on Quince MC (Cydonia oblonga Mill.) rootstock and trained as spindle at a planting density of 3.6 m × 0.7 m. Quince MC is one of the most dwarfing rootstocks available to control vigor in high-density pear orchards (Musacchi, 2011; Sansavini et al., 2008). The orchard (north-south oriented) was located at the Experimental Station of University of Bologna, Faculty of Agriculture, in Cadrozo, Italy (lat. 44°54′ 88.53″ S, long. 11°38′ 59.30″ W). Cultural practices were standardized according to those used in commercial orchards of ‘Abbé Fétel’ grafted on quince rootstocks.

Experimental design and sampling. A total of 12 homogeneous trees in size and vigor (average TCSA 60.2 ± 1.5 cm²) were selected during July 2012 within the orchard for this experimental trial. According to crop-load evaluated at harvest on 17 Sept. 2012, two levels were present for further tree categorization corresponding to medium crop-load (MCL) with an average of 25 pears per tree and 0.41 fruits/cm² TCSA, and low crop-load (LCL) with an average of 10 pears per tree and 0.16 fruits/cm² TCSA (Supplemental Table 1). A high crop-load level was not selected because it was absent in the orchard that year. In general, the orchard presented a quite low level of crop due to unfavorable weather conditions during bloom. For each level of crop-load, six trees were destructively evaluated, three of them after harvest of the 2012 season (9 Oct. 2012) with an average of 0.27 ± 0.06 fruits/cm² TCSA, and the others in January (14 Jan. 2013) with an average of 0.31 ± 0.09 fruits/cm² TCSA.

In each crop-load level (medium and low) three of the six trees were defoliated by hand before natural abscission (October) and three followed a natural defoliation during fall-winter (26 Nov. 2012).

In the 12 trees under investigation, we determined the fresh weight (FW) and dry weight (DW), starch, and carbohydrate concentrations of the different organs: wood, mixed buds, roots, and leaves. We also considered the growing degree days and chilling unit accumulation (Anderson et al., 1986; Richardson et al., 1974) (Table 1).

Trees were carefully dug out beneath the main root system (100-cm depth) using a trencher-equipped tractor at each time of sampling. Aboveground tree organs were fractioned into the following components: leaves (when present), 1-year-old branches (brindle-type shoots, mixed and vegetative buds, and vegetative shoots as 1-year shoots “mixed shoots”), 2-year-old branches, 3-year-old branches, 4+-year-old branches, short-old spurs, trunk (aboveground stem starting at the grafting point), and mixed buds. The mixed buds (buds containing both flower and leaf primordia are present) were sampled from each structure when they were present, and these were removed without including the portion of the spur to which they belonged. Vegetative buds at the time of destruction were considered as part of the holding wood structure where they belonged (no separation was made). Below-ground organs (below the grafting point) were divided into root stump, coarse roots (thickness >2 mm), and fine roots (thickness < 2 mm) (Silver and Miya, 2001; Zhang and Wang, 2015). Ages of the tissues...
refer to their physiological ages at the beginning of each season (e.g., 1-year shoots developed in the season 2011–12).

**Dry matter partitioning.** Immediately after tree destruction and organ partitioning, FW was measured for all components. Roots were rinsed with tap water to remove soil debris and left in the open air for 40 min before FW determinations. Randomly selected representative tissue subsamples were weighed then dried in a forced-air oven at 60 °C until constant mass to determine DW. Dry matter (%) was calculated as (DW/FW) × 100 (Palmer, 1988, 1992).

**Leaf measurements.** Leaf measurements were carried out at two different times of the experiment: the first sampling took place in October (22 d after harvest), and the second during natural leaf abscission (late November, Table 1). The total number of leaves was counted for each of three LCL and three MCL trees. At the October sampling, the total number of leaves was obtained by manually removing and counting all leaves in the canopy. Meanwhile, chosen trees for the late November leaves sampling were enclosed in a white net bag (material similar to nets used for hail protection) to collect leaves that otherwise would have fallen on the ground and dispersed. Leaves were harvested at four intervals from each net between 19 Oct. 2012 and natural leaf fall. The remaining leaves still attached to the trees (average of 5%) were manually harvested at the last sampling (end of November). At each partial sampling date, the number and total weight of leaves collected from the net were determined. Leaf area (LA, in centimeters squared) was determined with a leaf area meter (LI-3100; LI-COR, Lincoln, NE) on a subsample of leaves from each tree corresponding to 20% of total leaf weight/tree/sampling time. Total LA was then estimated by multiplying the number of leaves by the average LA obtained from the 20% subsample. Leaf Area Index (LAI) was determined with a leaf area meter (LI-3100; LI-COR, Germany) on a subsample of leaves and natural leaf fall. The remaining leaves still attached to the trees (average of 5%) were manually harvested at the last sampling (end of November). At each partial sampling date, the number and total weight of leaves collected from the net were determined. Leaf area (LA, in centimeters squared) was determined with a leaf area meter (LI-3100; LI-COR, Lincoln, NE) on a subsample of leaves from each tree corresponding to 20% of total leaf weight/tree/sampling time. Total LA was then estimated by multiplying the number of leaves by the average LA obtained from the 20% subsample. Leaf Area Index (LAI) was calculated as total LA of the tree divided by 20% subsample. Leaf Area Index (LAI) was determined with a leaf area meter (LI-3100; LI-COR, Germany) on a subsample of leaves.

**Spectrophotometer analysis.** Spectrophotometer analysis was performed as previously described by Mesa et al. (2016). The subsample was composed of 1 mL aliquot of the sample plus 10 mL of anthrone reagent. Subsequently, samples were boiled in water and then cooled to room temperature. The analysis was performed by reading the absorbance by spectrophotometer (VIS-ultraviolet Varian model Cary 1E; Varian, Inc., Palo Alto, CA) at 620 nm. Nonstructural carbohydrate concentrations were reported as glucose equivalents (milligrams per gram of DW of tissue) and corrected for the appropriate dilutions. Starch concentration was also multiplied by 0.9 to account for the mass of glucose theoretically hydrolyzed from a unit mass of starch (Seager and Haslemore, 1993; Smith and Zeeman, 2006).

**Statistical analysis.** The experiment was arranged as a completely randomized design with 12 trees as experimental units and two factors with two levels each (2 × 2): two crop-loads and two dates of sampling for leaf measurements. Three replicates were used for each date of sampling and crop-load level (2 dates × 2 crop-load levels × 3 trees/crop-load/date = 12 trees total). In the case of woody organs, roots, and mixed buds, only the date of sampling was considered as factor (not the crop-load), and for each treatment six replicates were used. All data collected were statistically analyzed with R software version 3.4.3 (R Core Team, 2017) using analysis of variance followed by Student-Newman-Keuls means separation. Differences in pairwise comparisons were considered significant at P ≤ 0.05.

**Results**

**FW and DW analysis.**

MCL trees of ‘Abbé Fétel’ pear trees evaluated in 2012 showed values for yield ≈2.5 times higher than those in LCL trees (Supplemental Table 1).

**Woody organs, roots, and mixed buds.** FW measurements for the different tissues showed no significant differences between sampling dates for the woody organs, mixed buds, and roots (Table 2). Mixed buds and 2-year-old branches exhibit a tendency of greater FW in January, although mean values were not statistically significant. Mixed bud FW corresponded to a significant 20% increase (Fig. 1). October treatment registered the highest number of mixed buds (619), although it did not differ statistically from the January treatment.

At the sampling date in October, the two levels of crop-load established (MCL and LCL) showed significant differences in terms of DW (g) only for the structure of 2-year-old (wood) branches (data not shown). The 2-year-old branches sampled from LCL trees obtained a 66.3% higher value for DW (g) in comparison with MCL trees. For all other sampled organs, no statistically significant differences were found (data not shown).

**Comparison between dates of sampling for DW (g) showed significant differences only for coarse roots (Supplemental Table 2).**

Mixed buds, 3-year-old branches, 4+-year-old branches, short-old spurs, trunk, and coarse and fine roots presented the same trend
the highest values.

where organs sampled in October showed
observed by FW (Supplemental Table 2),

Fig. 1. Average fresh weight (gray bars) and number (black squares) of mixed buds (MB) at different
sampling dates for average fresh weight. n.s., *** indicate no significance or significance at
respectively; Student-Newman-Keuls test was used for mean comparisons.

Table 2. Total fresh weight of each organ sampled per tree: woody organs, roots, and mixed buds evaluated
at different dates.

| Organs        | Unit | October | January | Significance |
|---------------|------|---------|---------|--------------|
| Wood          |      |         |         |              |
| 1-year        | g    | 534.0   | 528.7   | ns           |
| 2-year        | g    | 357.8   | 415.4   | ns           |
| 3-year        | g    | 365.2   | 282.8   | ns           |
| 4-year        | kg   | 3.4     | 3.0     | ns           |
| Spurs         | g    | 305.3   | 197.1   | ns           |
| Trunk         | kg   | 8.0     | 8.9     | ns           |
| Mixed buds    | g    | 48.0    | 60.0    | ns           |
| Total aboveground | kg   | 13.0    | 13.4    | ns           |
| Roots         |      |         |         |              |
| Coarse (>2 mm)| kg   | 2.2     | 1.7     | ns           |
| Fine (<2 mm)  | kg   | 111.0   | 107.1   | ns           |
| Root stump    | kg   | 1.7     | 1.8     | ns           |
| Total below ground | kg   | 4.0     | 3.7     | ns           |
| Whole tree    | kg   | 17.0    | 17.1    | ns           |

These values (means of six replicates) are obtained from the complete destruction of the trees. Lower case
letters indicate significant differences among means along rows.
*ns indicates no significance; Student-Newman-Keuls test was used for mean comparisons.

observed by FW (Supplemental Table 2),

Leaf measurements. Crop-loads showed an
effect on total leaf number, total leaf weight,
LA, and LAI, where MCL presented the highest
values for these parameters (Table 3). Regarding
the comparison between dates of sampling,
October vs. natural leaf abscission (late No-
vember), the parameters of total leaf weight,
average leaf weight, dry matter percentage (%
DM), and percentage of leaf water content (%
WC) were significantly different (Table 3). For
total leaf weight, the value recorded in October
was higher (1.67 kg/tree) than in late November
(1.12 kg/tree). The % WC and % DM showed
significant differences both for the level of
crop-load and sampling date, but not for the
interaction between them (Table 3). In this
way, leaves sampled from trees with an LCL
obtained a % DM of 2.7% higher compared
with MCL trees; whereas leaves sampled at
natural leaf fall showed DM equal to 70.5%;
23.1% higher with respect to leaves sampled
after harvest (Table 3). However, the interac-
tion between crop-load and sampling date was
meaningful only for the average leaf weight, as
shown in Table 3. Leaves collected in October
from trees with an LCL obtained the highest
value (356 ± 15 mg), whereas leaves at natural
leaf fall exhibited lower and similar values
regardless of the level of crop-load (Supple-
mental Fig. 1).

Nonstructural carbohydrate
determination
Nonstructural carbohydrate concentrations
of leaves. Between crop-load levels, the highest
values of soluble carbohydrate concentration in
leaves were obtained in LCL trees [49.2 mg
glucose (Glc)/g DW] (Table 4). Soluble carbo-
hydrates in leaves did not present a significant
interaction between crop-load level and date of
sampling, unlike what was observed for starch
concentration (Table 4). Regarding the interac-
tion between factors evaluated for the starch
concentrations, differences were found between
trees with different crop-load levels sampled in
October (Fig. 2), where leaves from trees with
an MCL showed the lowest concentration
(12.7 mg starch/g DW), being 53.6% lower
than leaves from trees with an LCL (27.4 mg
starch/g DW) (Fig. 2). At natural leaf abscission
(November), the starch concentrations corre-
spond to 28.3 ± 0.6 mg starch/g DW and 29.2 ±
1.2 mg starch/g DW for leaves with an MCL
and LCL, respectively, without significant sta-
tistical differences between them (Fig. 2).

Starch and soluble carbohydrates of
woody organs, roots, and mixed buds. The
two levels of crop-load identified at harvest
(MCL and LCL) did not present significant
differences in terms of starch and soluble
carbohydrate concentration for the roots and
mixed buds evaluated in October (Supple-
mental Tables 3 and 4). Regarding the dif-
ferent woody organs, starch concentration
was significantly different between MCL
and LCL only in spurs (higher in MCL) and
for soluble carbohydrates in 1- and 3-year-
old branches (higher in MCL).

Mixed buds (Fig. 3). roots (Supplemental
Fig. 2), or woody organs (Fig. 4) showed a
consistent pattern in soluble carbohydrate
concentration through the growing season.
The concentrations of soluble carbohydrates
for the woody structures increased from
October to January (Table 5). Among the
aboveground woody organs (Table 5), short-
old spurs obtained the largest increases be-
tween sampling dates, which corresponded to
an increase of 93.5% in mg Glc/g DW. How-
ever, the structures of 2-year-old branches
reported only a minor increase between dates
(51.4%) (Table 5). With respect to mixed buds,
nearly a 2-fold increase was recorded from
October to January (Fig. 3). This trend was
similar for the fine roots, in which changes
in carbohydrate concentration differed signi-
ficantly with an increase of 10.9% be-
tween sampled dates. The coarse roots did
not show significant statistical differences
between sampling dates (Supplemental
Fig. 2).

Regarding starch concentrations, only
the woody organs showed a clear trend through-
out the growing season. Namely, a decrease
occurred between the intervals of each sam-
pling date (Table 6, Fig. 4). Starch concen-
tration in short-old spurs exhibited the
smallest decline (6.5%) between October
and January, whereas the other aboveground
structures showed an average decrease of
34.5% (± 2.0% se) for the same period. The
starch concentrations in mixed buds and fine
roots increased by 21.8% and 18.0%, re-
spectively, between the two sampling dates
(Fig. 3, Supplemental Fig. 2).

With respect to trends among the
woody tissues, the greatest variation reg-
istered for starch concentration within the
The short-old spurs presented the highest values concentration values in October (Fig. 4). The significant results observed for DW found in 2-year-old branches in October between the two levels of crop-load (data not shown), may be explained by their yield performance (Supplemental Table 1). During the growing season, there is competition among sink organs by assimilates (Flore and Layne, 1999); therefore, in a year of LCL it could be reasonably expected that there may be less contention among fruits and vegetative sinks, in contrast to a year with a high crop-load where competition would be expected low growth (Naschitz et al., 2010), which increases with the age of the plant, as noted by Chalmers and Van den Ende (1975). In fact, our results showed that the 1-year-old branches obtained the highest values for the DW (active growth) (data not shown). Park (2011) and Monerri et al. (2011) agreed that during the growing season until harvest for persimmon (Park, 2011), and during postharvest for sweet orange (Monerri et al., 2011), the fruit FW and DW analysis

In the mixed buds, the significantly increased average FW between October and January (Fig. 1) could be explained by the corresponding increase in soluble carbohydrate concentration (Fig. 3). This observation aligns with Marquat et al. (1999), who noted the important absorption potential of sucrose and sorbitol in the stems and buds of peach trees (10-year-old) in October, just before the temperature fell during the dormancy period. Soluble carbohydrate transport increases during dormancy release, and buds become strong sink organs at that phenological stage. Indeed, budburst is triggered by soluble carbohydrate metabolism (Bonhomme et al., 2010; Ito et al., 2012; Lacointe et al., 1993; Marafon et al., 2011; Marquat et al., 1999). In our study, it is possible to observe a trend in the increase of soluble carbohydrates from October to January (Fig. 3), which probably can continue once budbreak begins, although we evaluated the concentrations of soluble carbohydrates and starch before dormancy stage and during dormancy but not up to dormancy release.

The significant results observed for DW found in 2-year-old branches in October between the two levels of crop-load (data not shown), may be explained by their yield performance (Supplemental Table 1). During the growing season, there is competition among sink organs by assimilates (Flore and Layne, 1999); therefore, in a year of LCL it could be reasonably expected that there may be less contention among fruits and vegetative sinks, in contrast to a year with a high crop-load where competition would be greater. Further, according to results reported by Gagliardi et al. (2014) in ‘Abbé Fétel’ pear trees, branches 3 years old and older held the highest production, followed by 2-year-old branches. The focus on these types of structures is because perennial branches form the skeleton (architecture) of the tree with minimal demand for assimilates for expected low growth (Naschitz et al., 2010), which increases with the age of the plant, as noted by Chalmers and Van den Ende (1975). In fact, our results showed that the 1-year-old branches obtained the highest values for the DW (active growth) (data not shown). Park (2011) and Monerri et al. (2011) agreed that during the growing season until harvest for persimmon (Park, 2011), and during postharvest for sweet orange (Monerri et al., 2011), the fruit FW and DW analysis

In the mixed buds, the significantly increased average FW between October and

**Table 3. Leaf measurements: number/leaves per tree, total weight per tree, average leaf weight, leaf area/tree, leaf area index (LAI), dry matter percentage (% DM) and water content percentage (% WC) regarding the levels of crop-load (CL) in the 2012 season: medium crop-load (MCL) and low crop-load (LCL), and dates of sampling (SD): October and natural leaf fall (November).**

| Sampling date (SD) | Leaf number/tree (n) | Total leaf wt/tree (kg) | Avg leaf wt (mg) | Leaf area/tree (m²) | LAI | % DM | % WC |
|-------------------|----------------------|------------------------|------------------|---------------------|-----|------|------|
| October           | 5,065                | 1.67 a                 | 333.0 a          | 6.48                | 2.57| 47.4 b| 52.6 a|
| November          | 5,289                | 1.12 b                 | 209.2 b          | 6.76                | 2.68| 70.5 a| 29.5 b|
| Significance      | NS                   | ***                    | NS               | NS                  | NS  | NS   | NS   |
| Crop-load (CL)    |                      |                        |                  |                     |     |      |      |
| MCL               | 5,959 a              | 1.59 a                 | 267.2            | 7.81 a              | 3.10| 57.6 b| 42.4 a|
| LCL               | 4,395 b              | 1.20 b                 | 275.0            | 5.44 b              | 2.16| 60.3 a| 39.7 b|
| Significance      | **                   | ***                    | NS               | ***                 | NS  | NS   | NS   |
| Interaction CL × SD| NS                   | NS                     | **               | ***                 | NS  | NS   | NS   |

Values are means of six replicates for each factor. The factorial analysis (interaction CL × SD) considered three replicates. Lower case letters indicate significant differences among means along rows. NS, *, **, *** indicate no significance or significance at P < 0.05, 0.01, or 0.001, respectively; Student-Newman-Keuls test was used for mean comparisons.

**Table 4. Leaf soluble carbohydrates [mg Glucose (Glc)/g dry weight (DW)] and starch concentrations (mg starch/g (DW)) for the two factors evaluated in the 2012 season, sampling dates (SD): after harvest (October) and natural leaf fall, and crop-load (CL): medium (MCL) and low (LCL) level.**

| SD                | Leaf soluble carbohydrates (mg Glc/g DW) | Leaf starch (mg starch/g DW) |
|-------------------|-----------------------------------------|-----------------------------|
| October           | 76.5 ± 1.9 a                           | 20.0 ± 2.1 b               |
| November          | 15.6 ± 1.0 b                           | 28.8 ± 0.7 a               |
| Significance      | VS                                      | ***                         |
| CL                |                                        |                             |
| MCL               | 42.8 ± 7.2 b                           | 20.5 ± 1.9 b               |
| LCL               | 49.2 ± 7.8 a                           | 28.3 ± 1.1 a               |
| Significance      | ***                                     | ***                         |
| Interaction CL × SD| NS                                     | NS                          |

Lower case letters along columns indicate significant difference in means. NS, **, *** indicate no significance or significance at P < 0.001, respectively; Student-Newman-Keuls test was used for mean comparisons.

same evaluation date, ranging from 50.2 to 102.0 mg g⁻¹ DW (October) for old short spurs and 2-year-old branches, respectively (Table 6). One-year-old shoots and 2-year-old branches presented the highest starch concentration values in October (Fig. 4). The short-old spurs presented the highest values of soluble carbohydrate concentrations, 35.6 mg g⁻¹ DW in October and 68.9 mg g⁻¹ DW in January, respectively (Fig. 4).
detachment allowed the remaining organs to become stronger sinks.

Starch and soluble carbohydrate analysis

Starch and soluble carbohydrates in leaves. During the fruit development stage up to 107 to 115 d after full bloom, fruits also accumulated starch (Mesa et al., 2016), behaving as storage organs to support metabolic activities. Mesa et al. (2016) also pointed out a decrease of starch accumulation in leaves during the first 100 d after full bloom, perhaps attributable to the high demand for assimilates by fruit (as sinks). Several studies have explained in detail the process of starch degradation in leaves (Zeeman et al., 2004) using Arabidopsis as a model (Smith et al., 2005; Streb and Zeeman, 2012; Zeeman et al., 2007), but little information is available about this process during the endormancy stage in nonphotosynthetic organs.

When the fruits reach their maturation on the tree, they are no longer considered sinks for assimilates because of the onset of starch degradation (Mesa et al., 2016). The hydrolysis of starch is correlated with a rise in soluble sugars content, an indicator of the incipient fruit maturity stage (Berüter, 2004; Mesa et al., 2016). This behavior is in contrast to other organs, such as shoots and roots (depending on their sink strength), as they require assimilates to accumulate reserves during fall before the tree enters into the dormancy period. This experiment showed the importance of leaf retention after harvest; leaves sampled in October showed a significantly higher amount of soluble carbohydrates with respect to leaves sampled at natural leaf fall (late November, Table 4).

Even if leaves entered in a senescence period as confirmed by their weight loss (Supplemental Fig. 1, Table 3) and % WC decrease in late November (Table 3), they still presented a significant concentration of soluble carbohydrates (15.6 mg Glc/g DW, Table 4). The occurrence of an adverse climate event or damage by diseases or pests would cause a source-sink imbalance (even in fall), due to the decrease of active photosynthetic foliage (Flore and Layne, 1999). McCamant (1988), cited by Loescher et al. (1990), noted that sweet cherry trees defoliated in August have the lowest amount of starch in all tissues compared with trees defoliated at later dates. Hudina and Stampar (2002) studied the effect of the reduction of LA on the quality of pear fruits cv. Williams during their development and concluded that a 30% reduction in LA induced lower soluble solids and sugars in fruits because of a decrease in production of assimilates. Other research on ‘Hayward’ kiwifruit reports that a 75% reduction of foliage at 5 d after full bloom decreases the rate of starch accumulation in summer on current shoots and trunk bark but does not affect the starch concentrations in roots (Cruz-Castillo et al., 2010). Kwack et al. (2014), unlike the other results reported on defoliation treatment, observed an increase of starch concentrations in roots of kiwifruit cv. Goldrush in October,
Table 5. Soluble carbohydrates concentration [mg Glucose (Glc)/g dry weight (DW)] in different types of branches: 1-year-old branches, 2-year-old branches, 3-year-old branches, 4+-year-old branches, and short-old spur.

| Organs          | October | January | Significance | Δ(January–October) | % Increase |
|-----------------|---------|---------|--------------|--------------------|------------|
| Wood            |         |         |              |                    |            |
| 1-year          | 32.7 b  | 51.1 a  | ***          | 18.4               | 56.3       |
| 2-year          | 31.7 b  | 48.0 a  | ***          | 16.3               | 51.4       |
| 3-year          | 29.7 b  | 52.0 a  | ***          | 22.3               | 75.1       |
| 4-year          | 32.1 b  | 49.9 a  | ***          | 17.8               | 55.5       |
| Spurs           | 35.6 b  | 68.9 a  | ***          | 33.3               | 93.5       |

Values are means of six replicates. Lower case letters along rows indicate significant difference in means between sampling dates (horizontally). *** indicates significance at P < 0.001; Student-Newman-Keuls test was used for mean comparisons.

Table 6. Starch concentration [mg Glucose (Glc)/g dry weight (DW)] in different types of branches: 1-year-old branches, 2-year-old branches, 3-year-old branches, 4+-year-old branches, and short-old spur.

| Organs          | October | January | Significance | Δ(January–October) | % Decrease |
|-----------------|---------|---------|--------------|--------------------|------------|
| Wood            |         |         |              |                    |            |
| 1-year          | 101.4 a | 66.7 b  | ***          | 34.7               | 34.2       |
| 2-year          | 102.0 a | 64.3 b  | ***          | 37.7               | 36.9       |
| 3-year          | 90.4 a  | 59.0 b  | ***          | 31.4               | 34.7       |
| 4-year          | 78.0 a  | 53.0 b  | ***          | 25.0               | 32.0       |
| Spurs           | 50.2 a  | 46.9 a  | ***          | 3.3                | 6.5        |

Values are means of six replicates. Lower case letters along rows indicate significant difference in means between sampling dates (horizontally). *** indicate no significance at P < 0.001, respectively; a Student-Newman-Keuls test was used for mean comparisons.

with a reduction of 75% and 100% LA between August and September. Moreover, the significant loss of LA on summer-pruned trees may lead to a reduction in the carbohydrate and nutrient element concentrations in the remaining tissues, ultimately limiting tree growth (Ikinci, 2014). Therefore, a defoliation treatment conducted between harvest and natural leaf fall could more intensely affect some downstream growth stages, causing impacts on crop phenology and fruit development (Tustin et al., 1997).

Naschitz et al. (2010) reported that high carbon demands induce soluble sugar synthesis, whereas low carbon demand enhances starch accumulation first in leaves and then in roots and woody tissues. Moreover, Monerri et al. (2011) observed that the levels of carbohydrates in the leaves of low-crop-loaded trees are double those in the high-crop-load trees. This appears to match our findings, in which LCL leaves registered 115.9% more starch concentration than in MCL 1 month after harvest (Fig. 2). However, similar results obtained by Wünsche et al. (2005) on ‘Braeburn’ apple trees showed no significant differences in starch concentration between two levels of cropping at the end of the season (natural leaf fall).

Starch and soluble carbohydrates of woody organs, roots, and mixed buds. Studies on other species are largely consistent with our findings. Primarily, the starch stored in woody tissues reaches its maximum level in October. This result appears initially in agreement with observations made in young walnut trees (Lacointe et al., 1993), in which a decrease of starch concentrations between postharvest and dormancy was mainly due to conversion to soluble sugars in the aerial organs and finest roots. It has been reported in previous works that starch accumulated in woody tissues is almost entirely hydrolyzed during the dormancy period for the further availability to new growth, budbreak, and fruit set of the following season (Table 6, Fig. 4). Short-old spurs showed the smallest decline in starch concentration in the transition between October and January (6.5%; Table 6, Fig. 4), but the highest increase in soluble carbohydrates (Table 5, Fig. 4). According to Naschitz et al. (2010) and Goldschmidt and Golomb (1982), the carbohydrate reserves in the current-year shoots and roots can be used directly during the growing season. The starch found in the tree architecture (old wood) can be considered an available reserve to support and maintain the growth of other tree organs. This might explain a higher soluble carbohydrate concentration obtained in January in short-old spurs (Table 5, Fig. 4).

Regarding soluble carbohydrates in woody organs, a reverse pattern of behavior was observed clearly in aboveground organs and in fine roots. Indeed, in January (dormancy period) an increase in soluble carbohydrate concentration was observed, confirming a process of interconversion between starch and soluble sugars (Oliveira and Priestly, 1988; Tables 5 and 6, Supplemental Fig. 2, Fig. 4). Sauter (1988) reported that low temperatures induced starch conversion to soluble sugars to improve cold acclimation and support the transition to ecodormancy. This could be explained by mobilization of soluble sugars from storage organs (roots and branches) toward utilization sinks (such as mixed buds before budbreak) in accordance with Marafon et al. (2011). Their results showed responses to cold temperature treatments (natural conditions, continuous artificial chilling, alternating temperatures, and lack of chilling) during the winter period. Marafon et al. (2011) pointed out that the increase of soluble sugars in branches of 16-year-old Japanese pear trees cultivar Housui could be the result of low temperatures during the dormancy period. This tendency also was observed in our experiment, where sugar accumulation in buds was reported in January (Fig. 4, Table 5).

Regarding the root systems, unlike what the other studies have reported (Goldschmidt and Golomb, 1982; Monerri et al., 2011), our experiment showed great storage of starch both in coarse and fine roots (Supplemental Fig. 2). It would be expected that after the harvest of an LCL year (“year off”), a greater accumulation of reserves in roots would occur considering the roots as the most active organ, as it has been reported for other species (Goldschmidt and Golomb, 1982; Lopez et al., 2007; Monerri et al., 2011); however, this was not observed in this experiment (Supplemental Fig. 2). One reason that at least partially explains the contradictory results could be that a small subportion had been selected as a representative subsample of the whole root system for the quantification. Therefore, there is a possible risk (larger the risk the larger the organ) of obtaining a value not entirely representative of the whole organ (Lopez et al., 2013). Despite this differing outcome, it was observed that the coarse roots showed greater storage capacity than fine roots where the starch concentration was lower. This agreed with the results described by Regier et al. (2010) for poplar trees. Fine roots, also known as “feeder roots,” are primarily essential for the absorption of water and mineral nutrients. Therefore, they are relatively short-lived and remain unaffected by the growing conditions of the season, as also observed by Monerri et al. (2011).

Conclusions

The results of this work confirm that in pear trees cv. Abbé Fétel, the interconversion between starch and soluble carbohydrates during the dormancy period was organ-specific, as reported for mixed buds, woody organs, and roots. The level of crop-load during the season had a major influence on the concentrations of starch compared with soluble carbohydrates after harvest (October); however, during the dormancy period (January) these differences were diminished. Among the woody organs, short-old spurs obtained the highest increase in soluble carbohydrates during dormancy, whereas for starch concentration, a significant decrease was observed in young structures. This increase in soluble carbohydrate concentrations observed during dormancy explains the need for soluble sugars to support the metabolic process and budbreak in early spring. On the other hand, the importance of the retention of active LA for a longer period showed that leaves in late November continued to produce assimilates, which are transported to the “sinking” organs in that period. Depending on time and intensity, the defoliation practice can influence the concentrations of nonstructural carbohydrates, which ultimately could affect bud quality and fruit set development for the following season. Our data
proved that a very heavy defoliation can reduce reserve in mixed buds and other organs.

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Supplemental Table 1. Yield performance of ‘Abbé Fétel’ pear trees evaluated in 2012, with two levels of crop-load.

| Crop-load 2012 | TSCA average (cm²) | Fruit (no./tree) | Yield (kg/tree) | Average fruit weight (g) |
|----------------|--------------------|------------------|-----------------|--------------------------|
| LCL            | 59.0               | 10 b             | 2.4 b           | 249                      |
| MCL            | 61.3               | 25 a             | 6.3 a           | 252                      |
| Significance   | NS                 | ***              | ***             | NS                       |

Lower case letters along columns indicate significant differences. NS, *, **, *** indicate no significance or significance at $P < 0.05$, 0.01, or 0.001, respectively; a Student–Newman–Keuls (SNK) test was used for mean comparisons. Values are mean of six replicates.

Supplemental Table 2. Dry weight of woody organs (above and underground) and mixed buds per pear trees cv. ‘Abbé Fétel’ evaluated at different dates.

| Organs      | unit | October | January | Significance |
|-------------|------|---------|---------|--------------|
| Wood        |      |         |         |              |
| 1-year      | g    | 255.1   | 263.6   | NS           |
| 2-year      | g    | 177.3   | 215.6   | NS           |
| 3-year      | g    | 188.2   | 149.5   | NS           |
| 4-year      | g    | 1,798.8 | 1,619.8 | NS           |
| Spurs       | g    | 152.9   | 99.6    | NS           |
| Trunk       | g    | 4,077.2 | 4,568.6 | NS           |
| Mixed buds  | g    | 20.0    | 26.6    | NS           |
| Roots       |      |         |         |              |
| Coarse (>2 mm) | g | 988.8   | a       | 709.5 b      | *            |
| Fine (<2 mm) | g  | 60.4    | 49.9    | NS           |

Values are means of six replicates. Lower case letters indicate significant differences among means along rows. NS, *, **, *** indicate no significance or significance at $P < 0.05$, 0.01, or 0.001, respectively; a Student–Newman–Keuls (SNK) test was used for mean comparisons.
Supplemental Table 3. Starch concentrations (mg·g⁻¹ DW) in different woody organs (above and underground) and mixed buds of pear trees cv. Abbé Fétel evaluated after harvest 2012 (October) according to the level of crop load: medium crop load (MCL) and low crop load (LCL).

| Organs/Crop load | Starch (mg starch/g DW) | MCL  | LCL  | Significance |
|------------------|-------------------------|------|------|--------------|
| Wood             |                         |      |      |              |
| 1-year           | 94.6                    | 108.2| NS   |              |
| 2-year           | 112.2                   | 91.7 | NS   |              |
| 3-year           | 95.6                    | 85.1 | NS   |              |
| 4-year           | 87.7                    | 68.3 | NS   |              |
| Spurs            | 62.6 a                  | 37.8 b| **   |              |
| Mixed buds       | 22.9                    | 24.6 | NS   |              |
| Roots            |                         |      |      |              |
| Coarse (>2 mm)   | 44.2                    | 40.1 | NS   |              |
| Fine (<2 mm)     | 17.3                    | 15.6 | NS   |              |

Lower case letters indicate significant differences along rows. NS, *, **, *** indicate no significance or significance at P < 0.05, 0.01, or 0.001, respectively; a Student–Newman–Keuls (SNK) test was used for mean comparisons. Values are mean of six replicates.

Supplemental Table 4. Soluble carbohydrate concentrations (mg·g⁻¹ DW) in different woody organs (above and underground) and mixed buds of pear trees cv. Abbé Fétel evaluated after harvest 2012 (October) according to the level of crop load: medium crop load (MCL) and low crop load (LCL).

| Organs/Crop load | Soluble Carbohydrates (mg glucose/g DW) | MCL  | LCL  | Significance |
|------------------|-----------------------------------------|------|------|--------------|
| Wood             |                                        |      |      |              |
| 1-year           | 34.4 a                                  | 31.1 b| *    |              |
| 2-year           | 31.9                                    | 31.6 | NS   |              |
| 3-year           | 31.8 a                                  | 27.6 b| *    |              |
| 4-year           | 32.1                                    | 32.1 | NS   |              |
| Spurs            | 33.8                                    | 37.3 | NS   |              |
| Mixed buds       | 23.4                                    | 21.9 | NS   |              |
| Roots            |                                        |      |      |              |
| Coarse (>2 mm)   | 50.5                                    | 50.1 | NS   |              |
| Fine (<2 mm)     | 35.0                                    | 34.4 | NS   |              |

Lower case letters indicate significant differences along rows. NS, *, **, *** indicate no significance or significance at P < 0.05, 0.01, or 0.001, respectively; a Student–Newman–Keuls (SNK) test was used for mean comparisons. Values are mean of six replicates.
Supplemental Fig. 2. Starch (white bars) and soluble carbohydrate (gray bars) concentrations (mg g⁻¹ DW) in coarse (A) and fine roots (B), for each of the two-sample date during the trial. Values means ± SE (n = 6). Capital letters indicate significant differences in means among dates for soluble carbohydrates, and lower case and underlined letters for starch. ns, *, **, *** indicate no significance or significance at P < 0.05, 0.01, or 0.001, respectively; Student–Newman–Keuls test was used for mean comparisons.