Introduction

In sweet cherry (Prunus avium L.), bloom and fruit set heavily depend on carbon (C) and nitrogen (N) storage reserves accumulated during the previous season after fruit harvest (Neilsen et al., 1997; Lang, 2002; Ayala 2004; Lang, 2005). It has been reported that the remobilization of N reserves from storage organs (roots, wood, bark and buds) begins just after bud break and that N...
root absorption begins three weeks later, indicating a dependence on the N reserves early in the season (Grassi et al., 2002; Grassi et al., 2003; Ouzounis and Lang, 2011). N storage reserves represent approximately 50% of the N used for new growth during early spring. The remaining N comes from root absorption and soil uptake (Grassi et al., 2002; Grassi et al., 2003; Azarenko and Chozinski, 2005).

Dependence on N reserves is critical for sweet cherry dwarfing combinations because smaller trees develop reduced woody structures (Lang, 2002; Lang, 2005). Soil N applications after harvest improve the accumulation of storage reserves in aerial organs, such as buds, stems and roots (Azarenko and Chozinski, 2005), but the timing is critical. In apple (Malus domestica Borkh.) and pear (Pyrus communis L.) trees, N applications to the soil after harvest were less effective in terms of the absorption and recovery of the fertilizer than were foliar applications (Shim et al., 1972; Shim et al., 1973; Kang and Titus, 1980; Sanchez et al., 1990). In sweet cherry, Gil (2000) reported a higher N uptake from leaves compared to roots.

In Chile, sweet cherry nutrition programs recommend applying N to the soil as the main source, although postharvest foliar applications are becoming increasingly important for highly productive dwarfing combinations. However, the benefit of N foliar applications in sweet cherries is unknown, and most information comes from other tree fruit species (Shim et al., 1972; Shim et al., 1973; Sanchez et al., 1990; Curetti et al., 2013). Urea [CO(NH$_2$)$_2$; 46% N] is one of the sources used during foliar applications of N. Urea is water soluble and nonpolar, facilitating its absorption through the leaves (Gil, 2006).

Considering the lack of information on the effectiveness of urea foliar applications in sweet cherry dwarfing combinations, this study focused on foliar sprays of $^{15}$N-urea, simulating commercial applications at various times after fruit harvest.

The research hypothesis was that the timing of the urea foliar application might influence the differential uptake and distribution of N in the fruiting branches of the semi-dwarfing sweet cherry combination “Bing”/“GI® 6.”

### Materials and methods

The study was carried out in a 5-year-old “Bing”/“GI® 6” sweet cherry orchard, located in Itahue, Region VII, Chile (35° 09’ 53” S, 71° 20’ 43” W). The orchard was trained in a central axis pattern and planted at 5.0 x 3.0 m. The trees were under drip irrigation. The soil type was a sandy-clay.

A completely randomized block design was used for the experiment. From a population of three hundred trees, one-hundred and twenty 3-year-old branches (one per tree) of uniform length, spur number, shoot number, leaf area (LA) and basal diameter were chosen randomly (Figure 1). Each 3-year-old branch included fruiting spurs (growth 2006/2007), non-fruiting spurs (2007/2008) and terminal current season shoot (growth 2008/2009) (Figure 1) leaf populations, without lateral extension shoots.

Trees were sprayed with unlabeled urea at various dates after fruit harvest (TR$_1$=Jan, TR$_2$=Feb, TR$_3$=Mar and TR$_4$=Apr). Prior to the application, one 3-year-old branch was bagged to isolate it from the rest of the tree and then sprayed with a solution of $^{15}$N-urea (1.8% atom; at 2% conc.) Each application date was considered a treatment (TR): TR$_1$= Jan, TR$_2$= Feb, TR$_3$= Mar and TR$_4$= Apr. Thirty branches (replications), one per tree, were labeled with $^{15}$N-urea per TR. A total of 10 branches per tree were harvested 15 days after foliar urea application (DAA); 10 branches were harvested during dormancy (Jul, 2010); and 10 branches were harvested during early spring (stage I of fruit development, Sep, 2010) for each TR. Additional branches (36 units) were used to carry out natural abundance measurements.
The urea foliar applications (labeled and unlabeled) were made within a 24 h period but divided into two applications on two consecutive days, with one evening (day 1) and one morning (day 2) application. A 1 L manual knapsack sprayer was used to apply the $^{15}$N-urea solution to individual branches. Run-off from the trees and individual branches was recovered using paper sheets placed on a semi-cylindrical metal frame. $^{15}$N-enriched branches were removed and divided into leaves, buds, bark, wood, roots and fruit during each developmental stage of sampling. Organs/tissues were dried in an oven (model AD 810 L) at 70º C. Later, organs/tissues were ground and prepared for mass spectrometry analysis (GC-MS, model “20-20”).

The nitrogen derived from fertilizer (NDFF %), nitrogen derived from fertilizer in mg (NDFF mg) and nitrogen use efficiency (NUE %) were calculated according to widely used formulas (IAEA, 1976; Hauck and Bremner, 1976; Millard and Nielsen, 1989; Pino et al., 2006).

Statistical analysis was performed using the module PROC MIXED in SAS® 9.1 (SAS Institute Inc., Cary, NC, version 8.2).

Results and discussion

Sweet cherry spur and shoot leaves were able to take up $^{15}$N-urea. The highest NDFF % values from leaves were observed 0 days after urea foliar application, except for TR1. In general, the lowest NDFF% values were recorded at leaf fall (Figure 2). N uptake by the leaves and translocation has been reported for pear (Sánchez et al., 1990; Curetti et al., 2013), almond (P. dulcis (Mill.) D.A. Webb.; Youssefi et al., 2000), apple (Shim et al., 1972; Shim et al., 1973) and grape (Vitis vinifera L.) (Porro et al., 2006). Lower NDFF% values in sweet cherry leaf at leaf fall indicates the translocation of N compounds to other organs within the branch or out of it after foliar application. The translocation of N has been reported in sweet cherry (Azarenko et al., 2008; San Martino et al., 2010; Ouzounis and Lang, 2011).

The NUE % of labeled branches varied depending on the timing of the urea foliar application (TR) and the phenological stage of the plant when the destructive sampling of the branches was performed (15 days after urea foliar application, dormancy and stage I of fruit development).
The longer the delay between $^{15}$N-urea spray and harvest, the lower the NUE% observed at each sampling date. A lower NUE% may be related to reduced leaf metabolic activity and leaf age. Leaves were physiologically less senescent in Jan than in Apr, particularly current season shoot leaves (i.e., sylleptic shoots). In sweet cherry, fruiting and non-fruiting spur leaves become fully mature approximately 20 to 30 days after bloom, whereas extension shoots continue developing new leaves after harvest. As the growing season progressed, the leaves of spurs and shoots become older and less efficient. It is likely that the lower NUE% values observed late in the season were the result of leaf senescence (Table 1).

When labeled branches were removed 15 days after urea foliar application, significant differences were observed among TRs. At that time, TR$_1$ had the highest NUE% and TR$_4$ had the lowest value for this parameter. A similar trend was observed when labeled branches were removed during dormancy and stage I of fruit development.

In this experiment, NUE ranged between 6.5 to 25.0% depending on the TR and sampling date. NUE% was higher following $^{15}$N-urea foliar applications close to fruit harvest. In apple (Shim et al., 1972; Shim et al., 1973, Kang and Titus, 1980) and pear (Sánchez et al., 1990) trees, foliar N applications after fruit harvest were more efficient than were soil N applications. In peach trees ($P$. persica L.), Nario et al. (2003) reported a NUE of 12.7% for soil applications during the summer. However, Neto et al. (2006) reported a NUE of 6.3% after N soil applications in pear trees. These authors used $^{15}$N as a tracer, and the trees were analyzed during dormancy. The highest NUE% values (25%) were recorded in TR$_1$ 15 days after urea foliar application. In general, the NUE% decreased between Jan (soon after fruit harvest) and Apr (leaf fall). The higher NUE% values in Jan and Feb might be explained by more efficient

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**Table 1.** Nitrogen use efficiency (%) for four dates of $^{15}$N urea foliar applications to “Bing”/ “Gisela®6” (TR$_1$=January, TR$_2$=February, TR$_3$=March and TR$_4$=April) and three rounds of destructive branch removal (15 days after application, dormancy and stage I of fruit development). Columns with the same letter are not significantly different ($\alpha = 0.05$). DAA: days after foliar urea application.

<table>
<thead>
<tr>
<th>Month</th>
<th>15DAA</th>
<th>Winter (Jul)</th>
<th>Spring (Oct)</th>
</tr>
</thead>
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<tr>
<td>Jan</td>
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<td>20.4 a</td>
<td>17.1 a</td>
</tr>
<tr>
<td>Feb</td>
<td>19.2 ab</td>
<td>10.4 b</td>
<td>13.7 b</td>
</tr>
<tr>
<td>Mar</td>
<td>14.4 b</td>
<td>9.0 b</td>
<td>12.2 b</td>
</tr>
<tr>
<td>Apr</td>
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<td>7.9 c</td>
<td>6.5 c</td>
</tr>
<tr>
<td>p-value</td>
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<td>0.0012</td>
<td>0.0001</td>
</tr>
</tbody>
</table>

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**Figure 2.** Percentage of nitrogen derived from fertilizer (NDDF %) in leaves from the combination “Bing”/“Gisela®6,” sampled 0 or 5 days after foliar urea application (DAA) and leaf fall (Jun) for four timings of $^{15}$N-urea foliar applications (TR$_1$=Jan, TR$_2$=Feb, TR$_3$=Mar and TR$_4$=Apr). Bars with the same letter for the same month are not significantly different ($\alpha = 0.05$). DAA: days after foliar urea application.
N uptake by the current season shoot leaves and a higher metabolic activity of the leaves. In addition, other sinks become important soon after fruit harvest, such as the roots and reproductive buds. In the combination “Bing”/“GI®6,” a peak in root growth has been observed during Jan and Feb in the Central Valley of Chile, along with active flower bud differentiation (unpublished data Ayala, 2012).

N translocation was evident during all foliar applications dates. The \(^{15}\)N contents of the leaves declined until leaf fall (Figure 2). This event coincided with a rise in the levels of \(^{15}\)N found in the bark, wood and buds, indicating the translocation of \(^{15}\)N compounds to these organs from the leaves after harvest (Figure 3 and 4). Because the recovery of N (expressed as NDDF mg) in the whole branch was less than the amount initially applied (data not shown), part of the taken-up N was translocated to other storage organs outside the branch, such as the trunk, other branches and root (Figure 4). In this experiment, roots of various sizes were sampled at each branch removal date, but no \(^{15}\)N enrichment was found in them (data not shown). The roots have been reported as a major

![Figure 3](image)

Figure 3. Distribution of nitrogen derived from fertilizer (%) in 3-year-old branches of “Bing”/“Gisela®6” sweet cherry from three growing sections: 2006/2007 (fruiting spur section); 2007/2008 (non-fruiting spur section); and 2008/2009 (current season shoot section), as well as after 4 dates of urea foliar application (Jan, Feb, Mar, Apr) and two times (A.- dormancy and B.- stage I of fruit development) with destructive branch removal. Bars with the same letter within the same month are not significantly different (\(\alpha = 0.05\)).
storage tissue for sweet cherry by other authors (Loescher et al., 1990; Grassi et al., 2003; Ayala, 2004; Gil, 2006). It is also likely that some of the urea applied was lost due to volatilization (Silva and Rodriguez, 1995).

A differential N distribution within labeled branches was observed depending on the timing of the foliar spray (Jan, Feb, Mar and Apr), the growth period (2006/2007; 2007/2008, and 2008/2009) and the phenological stage (15 days after application, dormancy and stage I of fruit development) at the time of branch removal (Fig 3). During dormancy, significant differences in N distribution among growth periods were observed for all TRs. Most of the $^{15}$N was recovered from the fruiting spur section (2006/2007 growth), whereas the lowest N levels were observed in the terminal current season shoots (2008/2009 sylleptic growth). Additionally, significant differences were detected in the N levels among organs during dormancy (bark, wood and buds) in all TRs. Bark exhibited the highest N levels (62 to 68% of the total recovery), followed by wood (27 to 35%) and buds (3 to 8%). Most of the N recovered from the fruiting spur section (2006/2007 growth) was in the bark. A similar pattern was observed in branches that were destructively removed during stage I of fruit development. Most of the N was recovered from the fruiting spur section. However, the N distribution among organ/tissues was different from the pattern observed during dormancy. In early spring, bark and wood showed a decrease in N levels following translocation to developing fruit and shoots.
These results are interesting because it is possible to conclude that in branches of the sweet cherry combination “Bing”/“GI®6,” reproductive spurs are a priority sink for N partitioning. Most of the N was translocated and accumulated in the fruiting spurs section to be remobilized into flowers, immature fruits and young leaves early in the spring, indicating that the N derived from foliar applications is not only taken up by the leaves and translocated to other organs but is also remobilized during the spring to promote budbreak, bloom and initial fruit development.

Therefore, the leaves of 3-year-old sweet cherry branches were able to take up foliar N supplied as 15N-urea after fruit harvest. N was translocated basipetally to various tissues/organs of the branch. The buds and bark registered the highest NDFF%, indicating a higher sink activity of these organs compared with wood. The highest N levels (expressed as NDFF mg) were found in the bark of the fruiting spur section for all dates evaluated. These results indicate that the fruiting spur section of the branch has the highest sink strength for N after harvest.

N foliar applications provided soon after fruit harvest (Jan) resulted in the highest NUE%. This high NUE % might be explained by the need for N during flower bud differentiation and/or a more efficient foliar N uptake by the current season shoot leaves, which are younger and thicker than the spur leaves and have a greater capacity to take up N.

In all TRs, the stored N was recovered in immature fruits and young leaves the following spring. Accordingly, urea sprays after fruit harvest might constitute an alternative to complement the N supply from the soil in sweet cherry combination “Bing”/“GI®6.” Further studies are needed to evaluate whether postharvest foliar urea should be applied multiple times and whether a similar pattern of N partitioning can be observed in other highly productive scion/rootstock combinations.

Resumen


Palabras clave: Corteza, dardos, dosel, eficiencia de uso del nitrógeno, frutos, hojas, madera de cerezo dulce, nitrógeno de almacenaje, yemas.
References


