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Annual and residual urea nitrogen contribution to the nutrition of peach trees (Prunus persica L.) grown under subtropical climate

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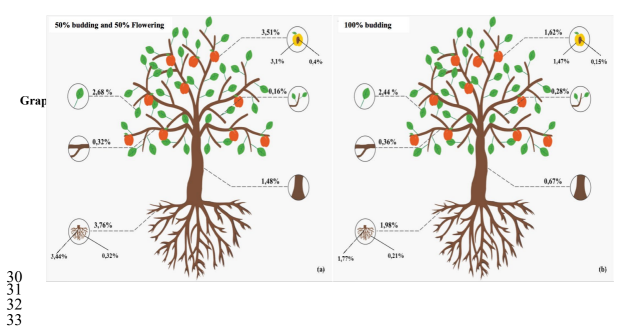
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1 2 3 4 Annual and residual urea nitrogen contribution to the nutrition of peach 5 trees (Prunus persica L.) grown under subtropical climate 6 7 Betania Vahl de Paula^{a*}, Beatriz Baticini Vitto^a, Paula Beatriz Sete^b, Talita 8 Trapp^b, Jovani Zalamena^c, George Wellington Bastos de Melo^d, Elena Baldi^e, Moreno Tosellie, Danilo Eduardo Rozanef, Gustavo Brunettoa 10 1 Department of Soil Science, Center of Rural Science, Federal University of Santa Maria, 11Av. Roraima 1000, Camobi, Santa Maria, RS, 97105-900, Brazil. E-mail: behde-12paula@hotmail.com; beabaticinivitto@gmail.com; brunetto.gustavo@gmail.com 13 Department of Rural Engineering, Center of Agricultural Sciences, Federal University of Santa Catarina, Rod. Admar Gonzaga 1346, Florianópolis, SC, 88034-000, Brazil. E-14mail: paulasete@gmail.com; talipptrali@yahoo.com.br $15_{\rm c}$ Federal Institute of Education, Science and Technology of Rio Grande do Sul, Rua 16Alberto Hoffmann 285, Restinga, Porto Alegre, RS, 91791-508, Brazil. E-mail: jovanizalamena@yahoo.com.br 17 d Embrapa Uva e Vinho, Rua Livramento, 515, Centro, Bento Gonçalves, RS, 95701-008, Brazil. E-mail: wellington.melo@embrapa.br 19 Department of Agricultural and Food Sciences, University of Bologna, Viale Fanin 44, 20Bologna, 40127, Italy. E-mail: elena.baldi7@unibo.it, moreno.toselli@unibo.it f São Paulo State University Julio de Mesquita Filho, Registro Campus, Rua Nelson Brihi 2 lBadur 430, Registro, SP, 13900-000, Brazil. E-mail: danilo.rozane@unesp.br 22 Corresponding author. E-mail: behdepaula@hotmail.com 23 Highlights 24 Residual urea nitrogen was found in the soil after two years. 25. Higher N deriving from fertilizer in leaves and fruits when the dose is split. 26 Most N found in trees is from sources other than nitrogen fertilizer. 27 28



34Recovery of N deriving from fertilizer (%) in peach organs subjected to the application of 40 kg N ha⁻¹ in 35ts urea form at budding (100% of the dose at budding - 100B) and split into two doses (50% of the dose at budding and 50% at flowering - 50B + 50F).

36 Abstract

37 Nitrogen (N) fertilizers such as urea are applied to peach orchards worldwide 38whenever the soil cannot meet plants' N demand. However, the actual nutritional contri-3 chution of different N supply modes applied to peach crops, or the contribution of residual N in the subsequent cycle, is yet to be fully known. The current study aims to assess the 40annual and residual urea N contribution to the nutrition of peach trees grown under sub-⁴Itropical climate. Forty kilograms of N per hectare supplied as enriched urea (3.0 at. % 42¹⁵N), were applied to peach in full production at a single rate (100% at budding) or split 43 in two rates (50% at budding and 50% at flowering). Total ¹⁵N and N concentrations in the application year and in the year after treatments were assessed in peach leaves 44 throughout the cycle, in fruits (pulp and stone) at harvest and in stratified soil samples. 45Total ¹⁵N, N concentrations and total dry mass were measured in annual and perennial 46 tree organs in the year after treatment application. Peach trees evidenced higher N deriving from fertilizer (NDFF) in leaves and fruits (pulp and stone) in the year N was applied 47 to the soil, as well as in the following year when N application was split into two rates. 48The highest NDFF amounts recorded in the year following N application were observed 40in leaves and fruits (annual organs) and thick roots (perennial organ), mainly when N was splitted. However, mainly in the year following its application, due to the small residual 50 N, the N found in trees derived from sources other than N fertilizer; this justifies annual 51

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59N applications, whenever necessary. The cultivation of cover crops and the preservation of organic matter could help N peach nutrition that seems to take advantage more of residual N in soil than on fertilizers.

61Keywords: Fruit farming, ¹⁵N isotope, N distribution, N fertilization, orchard.

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1. Introduction 63

Nitrogen (N) deriving from the mineralization of organic matter (OM) and de-64 composition of plant residues is not always able to satisfy the N demand of peach trees 65 (*Prunus persica* L.). To avoid N deficiency, which can cause a decrease in crop yield and 66 negatively affect fruit quality parameters (Damour et al., 2014; Jannoyer et al., 2011), N fertilizers are often applied to the soil surface. To establish the correct N rates to supply, 67 soil OM concentration and plant nutritional status, as well as growth and yield parameters, 68 should be evaluated (Brunetto et al., 2016b).

Urea is the N fertilizer most used in orchards thanks to its high N concentration and low cost per nutrient unit (Brunetto et al., 2016b). However, urea in the soil is rapidly 70 hydrolyzed by extracellular ureases enzymes produced by microorganisms such as bac-7 Iteria and fungi, and produce ammonium carbonate (NH₄⁺)₂CO₃, which is not stable in the 72soil. Urea decomposes into bicarbonate ion (HCO3-), hydroxide ion (OH-) and ammonium 73 ion (NH₄⁺) when it gets in contact with water. NH₄⁺ can react to OH⁻ and stimulate ammonia (NH₃) loss due to volatilization; however, part of NH₄⁺ in the soil can be trans- $74_{\rm formed}$ into nitrite (NO₂-) due to biological oxidation and, later, into nitrate (NO₃-) (Bru-75 netto et al., 2016a), which is often the prevalent form of N found in drained soils. How-76 ever, NO₃ in the soil is extremely mobile; therefore, it can be easily leached and contaminate subsurface water in orchards (Baram et al., 2016; Nevison et al., 2016). Therefore, 77_{it} is recommended to split N supply during the vegetative season, according to the phe-78 nological stage and plant needs focusing on the period of maximum requirement. Nitro-70gen supply at budding and spring can be able to increase root emission and longevity in the soil since this highly active organ accounts for the absorption of larger volumes of 80 water and nutrients such as N (Jordan, 2015). In addition, intense cell division and elon-8 Igation in shoot organs can be mainly observed just after flowering, a fact that leads to dry 82 matter increase and increases plant demand for N (Ventura et al., 2010). However, peach trees planted in soil presenting clayey texture and average organic matter contents, which was subjected to the application of single or split low N doses such as 40 kg N ha⁻¹ 84(CQFS-RS/SC, 2016), may absorb similar N amounts from the fertilizer. This outcome

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93 would be the desirable one since it would decrease costs with split nitrogen fertilizer applications, as well as potential water-contamination rates and increase the amount of N (from the fertilizer) absorbed by peach trees.

Part of the N absorbed by roots is preferably transported to growing organs such 96as leaves, annual shoots and fruits (El-Jendoubi et al., 2013; Jordan, 2015). Moreover, 97 part of the N accumulated in annual organs is redistributed, after harvesting, to storage organs such as stem, shoots older than one year and, mainly to the roots (Rivera et al., 982016; Zhang et al., 2012). If large amounts of N from the fertilizer accumulate inside 99beach trees in the fertilizer application year, it may not be necessary applying large N 100 rates to the soil in the next crop to reduce soil dependence on the fertilizer (Jordan et al., 2012) or even on N, since this element often derives from organic matter mineralization 101 and the decomposition of plant residues (Sabahi et al., 2016; TerAvest et al., 2010).

102 It is not clear whether the peach tree absorbs a greater amount of N, when the 103 fertilizer rate is applied in a fractional way and if the highest concentration of N present in the plant is from the origin of the fertilizer or other sources. However, the N amount accumulated in plant organs in the fertilizer application year, and in the subsequent year 105 when N different application modes (single or split N rates) are adopted are yet to be fully 106 known and this information could be more reliable when 15 N is used as a tracer (Brunetto et al., 2014). The hypothesis of the study is that peach trees absorb and accumulate greater amounts of N of the fertilizer when the application is split and that the greater amount of 108 N absorbed is derived from other N sources than not of the fertilizer.

109 The current study aimed to evaluate the annual and residual contribution of N 110^{derived} from the urea at a single rate or split for the nutritional status of peach trees.

111₂. Materials and methods

112.1. Experimental Site and Treatments

The experiment was conducted in July 2016, in Bento Gonçalves County, Rio Grande do Sul State, Southern Brazil (latitude 29°9'54.50"S; longitude 51°32′3.87"W) 114 on a peach orchard of the cultivar 'Chimarrita' grafted on 'Capdeboscq' rootstock . Trees, 115 trained as in "Epsilons" system were planted in 2009 at density of 1,666 plants ha $^{-1}$ (1.5 116m between plants and 4 m between rows). The orchard was planted in Cambisol Humic (Sibcs, 2013) and Typic Hapludalf soil (Soil Survey Staff, 2014) and presented, before planting, in the 0-0.2 m layer, the following characteristics: clay (310 g kg $^{-1}$), silt (468 g 118kg $^{-1}$) and sand (280 g kg $^{-1}$); OM (26.5 g kg $^{-1}$); pH in water 5.7 (1:1 ratio); exchangeable

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 $127^{Al~(0.0~cmol_c~dm^{-3})}, Ca~(7.4~cmol_c~dm^{-3})~and~Mg~(2.3~cmol_c~dm^{-3})~extracted~through~1mol~128^{KC1~L^{-1}}; available~P~(8.6~mg~dm^{-3})~and~K~(207~mg~dm^{-3})~-both~extracted~through~Mehlich-1;~and~total~N~(2.60~g~kg^{-1}).~Climate~in~the~region~is~subtropical~(type~Cfa);~the~mean~129_{annual~rainfall~is~1,736~mm~(Table~S1)}.$

The following N application strategies were compared as in completely random131 ized block design with three replicates: 100% N rate supplied at budding (100B) and N splitted as 50% of total rate and at budding + 50% of N at the end of flowering (50B + 132 50F). Each replicate comprised five plants; the three central plants were subjected to N 133 application and evaluated. Nitrogen was applied at a rate of 40 kg N ha⁻¹, which is equiv134 alent to 54.5 g N plant⁻¹; this N rate is the quantity recommended for soils presenting 2.6% to 5.0% OM (CQFS-RS/SC, 2016) and was supplied as enriched urea (3.0 at. % 135 15 N) in 2016. The fertilizer was applied on the soil surface of a 1-m⁻² area considering the 136 free stem in the center of the area. Ground cover plants found in the urea application area 137 were manually removed at treatment application time. Cover plants found in the treatment application region were desiccated with non-residual herbicide (glyphosate was the active ingredient) every 30 days, throughout the peach tree cycle.

1402.2. Assessment and analyses

Ten full expanded leaves per plant were collected at budding, flowering, fruit growth, harvest and senescence in 2016 and 2017. Leaves were then washed, dried, 142weighted, milled and analyzed for N and ¹⁵N concentration determined with mass spec-143rometry (Finnigan MAT mass spectrometer, Delta Plus model), according to Brunetto 144⁽²⁰¹⁴⁾.

At harvest of both years, the yield was recorded and fruits were counted; in addi145tion, a sample of 10 fruits was collected, fruit pulp was manually separated from the stone;
146organs were dried, weighted and analyzed as described for leaves. In 2017, plants were
147uprooted with the aid of a tractor and separated into leaves, annual shoots, branches older
than one year and stem. Roots were manually separated from the soil and divided into
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thin (diameter \leq 2 mm) and thick (diameter \geq 2 mm) roots (Hendrick and Pregitzer, 1992).
149They were washed with running water and, subsequently, with distilled water. All organs
150were weighted to determine fresh weight; a subsample of each organ was then collected
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and fresh and dry weight was determined. All organs were then ground and analyzed as
described before. Soon after fruit harvest in 2016 and 2017, soil samples were collected
152at the depth of 0.0-0.025, 0.026-0.05, 0.051-0.10 and 0.11-0.20 m in the crown projection

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161 area where treatments were applied. The soil was air-dried, subjected to 2-mm-mesh sieve
and stored for total N and ^{15}N analysis. 162
163_{2.3.} Calculations and statistical analysis
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                   Excess <sup>15</sup>N atoms was calculated according to the following equation
165(Equation 1):
Excess ^{15}N atoms in the sample (%) = % ^{15}N atoms in the sample – 0.3663% 166
                                                                                      Equation 1
                   The percentage of excess <sup>15</sup>N atoms in the sample, total N amount and the
167 percentage of <sup>15</sup>N in the fertilizer applied to the soil were used to calculate N deriving
168 from the fertilizer (NDFF) (Equations 2 and 3):
169NDFF (%)=(% excess ^{15}N atoms in the sample/% excess ^{15}N atoms in the fertilizer) x100
NDFF (g)=Total N in the sample (g)x(% excess ^{15}N atoms in the sample/% excess ^{15}N atoms in the fertilizer)
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                   Afterwards, results of the aforementioned equations were used to calculate
172N deriving from soil (NDFS) (Equation 4):
173 NDFS (%) = 100-NDFF (%)
                                                                                      Equation 4
                   Recovery of N deriving from the fertilizer by plants (R) was calculated
174according equation 5:
175R (%) = NDFF / Amount N fertilizer applied to the soil (mg) x 100
                                                                                     Equation 5
           Total N, NDFF and NDFS content in each organ for calculated by multiplying N
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    concentration for organ dry weight.
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           Results were subjected to the D'Agostino-Pearson normality test. Data of total N,
178 excess <sup>15</sup>N and NDFF in 2016 and 2017 in leaves, were analyzed as in a factorial experi-
179mental design with application mode (2 levels: 100B and 50B+50F) and sampling time
    (4 levels: budding, end of flowering, fruit growth, fruit harvest) as main factors. Data of
180^{\circ}_{total\ N,\ excess\ ^{15}N\ NDFF} in 2016 in fruits, were analyzed as in a factorial experimental
18 Idesign with application mode (2 levels: 100B and 50B+50F) and organ (2 levels: pulp
182 and stone) as main factors. Total N and 15N in soil were analyzed as in a factorial exper-
    imental design with application mode (2 levels: 100B and 50B+50F) and sampling depth
183 (4 levels: 0-0.025, 0.026-0.05, 0.051-0.01, 0.011-0.02) as main factors. When analysis of
184 variance showed a statistical effect of treatments (P \leq 0.05), means were separated by
185Student Newman Keuls test. When the interaction between factors was significant, 2
times standard error of means (2SEM) was used as the minimum difference between two 186
    means statistically different for P \le 0.05.
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1963. Results

Interaction between treatment and sampling time was not significant for total N, excess \$^{15}N\$ and NDFF in leaves in 2016 and 2017, consequently Tables 1 and 2 report the \$198_{\rm effects}\$ of main factors. Total N was higher in 100B than in 50B+50F in 2016 (Table 1) 199and 2017 (Table 2); excess \$^{15}N\$ and NDFF were higher in 50B+50F than in 100B both in \$200^2016\$ (Table 1) and 2017 (Table 2). The highest total N concentrations were observed in leaves collected at budding and end of flowering, followed by those sampled at fruit har-201_vest, senescence and fruit growth (Table 1). Excess \$^{15}N\$ atoms and NDFF rates were 202nigher in leaves collected at end of flowering and fruit harvest than those sampled at fruit 203_growth, senescence and budding (Table 1). Leaves sampled at fruit growth showed higher \$^{15}N\$ excess and NDFF than those at budding and senescence that showed similar values 204_(Table 1).

In 2017, total N concentration, as well as excess 15 N atoms and NDFF, were higher 206in leaves collected at the fruit growth than all other sampling data (Table 2). Excess 15 N and NDFF showed similar values between all other sampling data. Leaves total N was 207 similar at budding and end of flowering and higher than the values measured at fruit har- 208 Vest and senescence (Table 2).

Interaction between treatment and sampling time was not significant for total N,
 excess ¹⁵N and NDFF in fruits in 2016, consequently table 3 only reports the effects of main factors. Total N concentration, as well as excess ¹⁵N atoms and NDFF, have ob-211_{served} fruits of trees subjected to 50B + 50F than those supplied with a single treatment 212(Table 3). All values were higher in pulp than in stone (Table 3).

Fruits (pulp and stone), leaves, shoot and stem dry weight was higher in 50B+50F than in 100B; the opposite was observed for thick roots; no significant differences were observed for the 1-year shoot and fine roots (Table 4).

Total N content in leaves, fruit pulp and stone and thick roots was higher in trees 216 subjected to 50B+50F applications than 100B; the opposite was observed for the shoot (Table 5). No significant difference was observed for other organs (Table 5).

N deriving from fertilizer in fruit stone and pulp and thick roots was higher in trees 218subjected to 50B+50F applications in comparison to 100B; no significant differences 219were observed between treatments for other organs (Table 6).

N deriving from soil in fruits (pulp and stone), leaves, shoot and thick roots was higher in 50B+50F than 100B; no differences were observed for other treatments (Table $22\,l_{7}$).

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Total N content in the soil profile between 0 and 0.10 m was higher when N was 231 split in comparison to the entire rate in 2016; while at 0.011-0.02 m of depth the values of total N were higher for 100B than for 50B+50F (Table 8). In 2017 total N was higher 232 in soil 50B+50F than in 100B (Table 8). Total N decreased with depth both in 2016 and 2332017 (Table 8).

The excess ¹⁵N atoms were higher, in 2016 and 2017, in all layers of soil subjected to 100B applications (Table 9) and, for both treatments, the values decreased with depth 235_(Table 9).

Soil NDFF was higher in 50B+50F than 100B along with the entire soil profile 237both in 2016 and 2017; moreover, the values decreased with depth for all treatments and 238in all years (Table 10).

239₄. Discussion

The split of N rates in two different phenological phase enhanced leaves ¹⁵N atoms 241 excess and NDFF in both years probably because this application method improved the synchronism between the availability of mineral N in the soil and N uptake by the root 242 system (Radicetti et al., 2017; Sabahi et al., 2016). According to some authors (Brunetto 243 et al., 2016a; Neto et al., 2008; Roccuzzo et al., 2012) at flowering peach trees start pro-244 ducing new roots that enlarged the surface area and increased the volume of soil explored by the root system, enhancing, consequently, the uptake of water and nutrients. In the 245 period immediately after flowering, there is also intense vegetative activity due to the 246 formation of new shoots that become a sink for nutrients, mainly N (Brunetto et al., 2072016a; Carranca et al., 2018).

In 2017, the year after the application of enriched urea, leaves showed lower ¹⁵N atoms and NDFF in both N application techniques (100B and 50B + 50F) probably be-249 cause plants allocated part of the ¹⁵N assimilated to fruits as also evidenced previously 250 (Muhammad et al., 2015). According to our results, we evidenced that N was mainly allocated to the pulp and, to a lesser extent, to stones with more evident results in 2016 than in 2017. Fruits from plants fertilized with a split mode (50B+50F) evidenced, in both 252 years, higher ¹⁵N values, reinforcing the hypothesis of a greater synchronism between N 253 applications and absorption. Moreover, peach pulp recorded higher excess ¹⁵N atoms and 254 NDFF than stone as also demonstrated previously (Kuo et al., 2016; Pescie et al., 2018). However, excess N allocation to the pulp can lead to worse fruit quality and an increase 255 of fungal diseases in the field or during storage (Brunetto et al., 2015; Bush et al., 2018).

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Peach trees subjected to split N applications (50B + 50F) accumulated more N than those subjected to a single application mainly in leaves and pulp, which are annual organs that present intense growth and increased dry mass throughout the phenological 266_{stages} (Pescie et al., 2018; Roccuzzo et al., 2017) being the main N sinks during the veg-267 etative season.

The N plant content in the second year was derived mainly from sources other than the fertilizer, probably from organic matter mineralization and from waste decomposition, since the NDFF rate in most organs did not exceed 2.5%, except for the thick 270 roots, which recorded values close to 3.5%.

Decreased total N, excess ¹⁵N atoms and NDFF values in leaves at senescence, or even at fruit growth, are probably due to protein degradation and remobilization of N 272 forms from leaves to reserve organs such as stems, branches older than one year, or roots 273 (Brunetto et al., 2016a; Roccuzzo et al., 2017). Part of the N accumulated in storage or-274 gans may be redistributed to annual organs growing in the subsequent cycle, a fact that can decrease fruit tree dependence on N applied in the year (Carranca et al., 2018; Roccuzzo et al., 2017).

It is well known that in spring, peach trees use N stored in perennial organs and it 277was estimated that the majority of N remobilization occurs before root uptake starts 278 (Rufat and DeJong 2001. The remobilization process is little affected by the amount of soil N (Tagliavini and Millard, 2005) but the duration depends on the amount of stored 279N, being longer in trees with large storage pools (Grassi et al., 2003). Once remobilization 280 finishes, root uptake provides the remainder of the N used for growth; consequently, from 281 this stage until the end of the season it is important to maintain adequate N in the soil.

As a consequence of soil OM mineralization in the soil there is an increase of N 282 availability that not derives from enriched urea and determines an increase of the amount 283 of N deriving from the soil in the plant (García-Orenes et al., 2016). This partly justifies 284 the low leaf NDFF rates in the N application year (2016) and in the following year (2017) which did not exceed 17% and 3% respectively, throughout leaf collection times in both N supply techniques. Moreover, it is also evidenced by the higher NDFS content that 286 NDFF showed in plants at the end of the experiment.

The higher excess ¹⁵N atoms and NDFF values observed in topsoil layers, mainly in soil subjected to split N application (50B+50F), can be attributed to the complexation/adsorption of part of the N applied to organic compounds of organic matter on the 289_{soil} (Zhang et al., 2015), a fact that was observed in orchards subjected to different N

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298 rates and application times (Brunetto et al., 2016a; Roccuzzo et al., 2017). Thus, the split mode should induce greater soil mineral N availability in the most superficial layers, those more explored by roots, explaining the greater incidence of applied N in split mode $300_{50B+50F}$) on leaves and fruits N concentration.

 $301 \qquad \text{The lower values measured in 2017 could be due to loss of the N supplied in 2016} \\ 302 \\ \text{due to volatilization (Carranca et al., 2018; Dominghetti et al., 2016; Pescie et al., 2018),} \\ \text{denitrification (Nevison et al., 2016), leaching (Sparks, 2018; Lynch and Wojciechowski, 303_{2015)} \\ \text{and runoff (Baram et al., 2016; Dominghetti et al., 2016; Pescie et al., 2018)}.$

Data from this experiment evidence that N should be applied yearly and possibly 305 in split mode to reduce as much as possible loss of N in the environment and at the same time, meet plant needs. Moreover, the maintenance of high soil OM levels with different techniques such as minimum soil tillage, organic fertilization, ground cover plant culti-307 vation and maintenance of plant residues on the soil (Baldi et al., 2016; Brunetto et al., 3082014; Radicetti et al., 2017) could help to reduce N loss in the environment and improve plant nutritional status.

310_{5. Conclusion}

311 Peach trees allocated more N derived from urea in leaves and fruits than in other 312 organs, to a higher extent in the year of fertilizer application more than in the following year, probably due to N loss in the environment or absorbed by plants. The best results in 313 terms of N concentration were observed as a consequence of the split mode showing that 314 with this technique it is easier to meet plants' need.

For the purpose of recommendation, it is important to make it clear that although the plants show better results with the fractionation of N rates; however, at the time of 316 application it should be checked whether the costs for fertilizer supply (which includes 317 double use of machinery and labor) offset the productivity gain and guarantee profits for 318 the producer.

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325 Author Contributions

- Betania Vahl de Paula carried out the assembly and experimental collection, analysis and 326 interpretation of data and co-wrote the paper. Beatriz Baticini Vitto, Paula Beatriz Sete,
- 327 Talita Trapp and Jovani Zalamena collected the data, laboratory analysis and statistical
- 328 analysis. Gustavo Brunetto and Elena Baldi analyzed the data and co-wrote the paper.
- 329 George Wellington Bastos de Melo, Moreno Toselli, Danilo Eduardo Rozane revised the paper.

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336 Conflicts of Interest

337_{The authors have declared that no competing interests exist.}

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Table 1. Effect of urea application and sampling time on total N, excess ¹⁵N atoms and N deriving from fertilizers (NDFF) on peach leaves in 2016.

Treatment	Total N (% DW)	Atom ¹⁵ N excess (% DW)	NDFF (% DW)
50B+50F	2.92	0.326	10.9
100B	3.08	0.242	8.08
Significance	*	***	***
Sampling Time			
Budding	3.55 a	0.120 с	4.00 c
End of flowering	3.51 a	0.403 a	13.4 a
Fruit growth	2.20 d	0.313 b	10.4 b
Fruit harvest	3.32 b	0.401 a	13.4 a
Senescence	2.42 c	0.184 с	6.14 c
Significance	***	***	***
<i>Treatment</i> × <i>Time</i>	n.s.	n.s.	n.s.

Values followed by the same letter are not statistically different according to Student Neuman Keul test (P \leq 0.05). n.s., *, ***: effect not significant, significant at P \leq 0.05 and P \leq 0.001, respectively.

Table 2. Effect of urea application and sampling time on total N, excess ¹⁵N atoms and N deriving from fertilizers (NDFF) on peach leaves in 2017.

Treatment	Total N (% DW)	Atom ¹⁵ N excess (% DW)	NDFF (% DW)
50B+50F	3.15	0.048	1.61
100B	3.30	0.039	1.30
Significance	***	***	***
Sampling Time			
Budding	3.42 b	0.0388 b	1.29 b
End of flowering	3.45 b	0.0345 b	1.15 b
Fruit growth	3.74 a	0.0745 a	2.49 a
Fruit harvest	2.76 c	0.0341b	1.14 b
Senescence	2.76 с	0.0357 b	1.19 b
Significance	***	***	***
Treatment×Time	n.s.	n.s.	n.s.

Values followed by the same letter are not statistically different according to Student Neuman Keul test ($P \le 0.05$). n.s., *, ***: effect not significant, significant at $P \le 0.001$, respectively.

Table 3. Effect of urea application and organ on total N, excess ¹⁵N atoms and N deriving from fertilizers (NDFF) on peach fruit in 2016.

Treatment	Total N (% DW)	Atom ¹⁵ N excess (% DW)	NDFF (% DW)
50B+50F	0.738	0.234	7.88
100B	0.580	0.182	5.68
Significance	**	*	***
Organ			
Pulp	0.893	0.242	8.17
Stone	0.424	0.174	5.40
Significance	***	*	***
Treatment×Organ	n.s.	n.s.	**

n.s., *, **, ***: effect not significant, significant at P \le 0.05, P \le 0.01 and P \le 0.001, respectively.

Table 4. Effect of urea application on organ biomass (g pt⁻¹) at the end of the experiment.

Treatment	Stone	Pulp	Leaves	Branche s	Shoot	Stem	Fine roots	Thick roots
50B+50F	929	6447	3142	698	1335	8966	305	4480
100B	712	4699	2554	518	1230	5836	349	4961
Significance	*	*	*	*	n.s.	*	n.s.	*

n.s., *: effect not significant and significant at P $\!\leq\! 0.05,$ respectively.

Table 5. Effect of urea application on organ total N content (g pt⁻¹) at the end of the experiment.

Treatment	Ston e	Pulp	Leaves	Shoot	Shoot (1 year old)	Stem	Fine roots	Thick roots
50B+50F	9.52	62.8	94.4	5.60	7.84	26.2	3.21	51.9
100B	5.11	39.0	64.4	8.53	6.14	15.9	3.21	40.5
Significance	*	*	*	*	n.s.	n.s.	n.s.	*

n.s., *: effect not significant and significant at P \leq 0.05, respectively.

Table 6. Effect of urea application on organ N deriving from fertilizer (g pt^{-1}) at the end of the experiment.

Treatment	Ston e	Pulp	Leaves	Shoot	Shoot (1 year old)	Stem	Fine roots	Thick roots
50B+50F	0.140	1.12	0.964	0.056	0.129	0.397	0.114	1.24
100B	0.053	0.528	0.877	0.100	0.115	0.240	0.074	0.638
Significance	*	*	n.s.	n.s.	n.s.	n.s.	n.s.	**

n.s., *, **: effect not significant, significant at P \le 0.05 and P \le 0.01, respectively.

Table 7. Effect of urea application on organ N deriving from soil (g pt⁻¹) at the end of the experiment.

Treatment	Stone	Pulp	Leaves	Shoot	Shoot (1 year old)	Stem	Fine roots	Thick roots
50B+50F	9.38	61.6	93.5	5.55	7.72	25.8	3.10	50.6
100B	5.05	38.4	63.5	7.84	6.01	15.6	3.14	39.9
Significance	*	*	*	n.s.	n.s.	n.s.	n.s.	*

n.s., *: effect not significant and significant at P \leq 0.05, respectively.

Table 8. Effect of urea application and depth (m) on soil total N content (g $kg^{\text{-1}}$) in 2016 and 2017.

Treatment	0-0.025	0.026-0.05	0.051-0.01	0.011-0.02
		2016		
50B+50F	20.5	19.5	14.6	12.7
100B	19.0	15.6	13.4	13.4
Significance		2SEM	=0.412	
		2017		
50B+50F	20.7	18.3	16.8	15.5
100B	18.7	17.0	15.7	15.3
Significance		2SEM	=0.146	

Values differing by 2 standard error of means (SEM) are statistically different. Interaction treatment*depth significant at P < 0.05 in 2016 and P < 0.01 in 2017.

Table 9. Effect of urea application and depth (m) on soil 15 N atom excess (%) in 2016 and 2017.

Treatment	0-0.025	0.026-0.05	0.051-0.01	0.011-0.02
		2016		
50B+50F	0.032	0.018	0.006	0.003
100B	0.043	0.032	0.024	0.010
Significance		2SEM=	=0.0011	
		2017		
50B+50F	0.019	0.009	0.007	0.005
100B	0.029	0.021	0.011	0.009
Significance		2SEM=	=0.0010	

Values differing by 2 standard error of means (SEM) are statistically different. Interaction treatment*depth significant at P < 0.05 for both 2016 and 2017.

Table 10. Effect of urea application and depth (m) on soil N deriving from fertilizer (%) in 2016 and 2017.

Treatment	0-0.025	0.026-0.05	0.051-0.01	0.011-0.02
		2016		
50B+50F	1.44	1.05	0.806	0.333
100B	1.08	0.602	0.211	0.103
Significance		2SEM	=0.038	
		2017		
50B+50F	0.965	0.715	0.353	0.291
100B	0.634	0.295	0.231	0.181
Significance		2SEM	=0.035	

Values differing by 2 standard error of means (SEM) are statistically different. Interaction treatment*depth significant at P < 0.05 for both 2016 and 2017.

Suplementary

Table S1. Mean monthly rainfall, air temperature and relative humidity during the experimental months in 2016 and 2017.

Month	Phenological	Avera	ge air	Average	rainfall	Relati	ive air
Wionth	stage	tempera	ture (°C)	(m	m)	humid	ity (%)
		2016	2017	2016	2017	2016	2017
February	Start of senescence	22.6	22.2	146	147	79.2	79.7
June	Budding	10.0	13.8	7	160	78.2	81.5
July	Flowering	12.8	14.5	192	29	77.5	69.5
September	Fruit Growing	14.2	19.1	84	107	74.1	73.9
November	Start of fruiting	18.6	18.2	104	1620	70.9	70.4