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Contribution of cover crop residue decomposition to peach trees nitrogen nutrition

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Contribution of cover crop residue decomposition to peach tree nitrogen nutrition

3 Abstract 4 Purpose: Cover crop nitrogen (N) cycling has an important role in agricultural production and 5 contributes to peach (Prunus persica (L.) Batsch.) N nutrition. This study evaluated black oat 6 (Avena strigosa Schreb) and ryegrass (Lolium multiflorum L.) residue decomposition dynamics, 7 N recovery from cover crop residues, and N compartmentalization in peach tree organs. 8 *Methods*: A two-year field trial was developed with labeled (5 atom% ¹⁵N excess) cover crop 9 shoot biomass application in a 5-year old peach orchard. The region's climate is warm temperate 10 (Cfb), and the soil is classified as a Typic Hapludalf. Litter bags with unlabeled shoot residues 11 were also deposited in the orchard to assess biomass, carbon (C), N, lignin, cellulose, and non-12 structural biomass decomposition dynamics. 13 *Results:* After 13 months, the leaves, trunk, and roots showed the greatest proportion of N 14 derived from residues (Ndfr) (35.4, 25.1, and 22.4%, respectively) while the greatest 15 concentrations of 15 N and Ndfr occurred in roots <2 mm (0.0376 and 0.94%, respectively). The 16 N derived from cover crop shoots in the second production cycle was similar among tree organs.

17 Ryegrass residues presented the highest decomposition constant (k) values for dry matter, total
18 organic carbon (TOC), cellulose, and lignin. Hence, black oat residues presented a higher half-

19 life $(t^{\frac{1}{2}})$ for dry matter, TOC, total N, cellulose, and lignin.

20 Conclusions: The N derived from black oat and ryegrass residues in mature trees was
21 expressively low (<1%) and similar between species. Within organs, the highest Ndfr occurred</p>
22 in peach leaves during the flowering stage, when the greatest residue decomposition rate also
23 occurred. Soil N and plant internal N reserves are the major N sources for newly formed organs,
24 but greater contributions to tree N nutrition may occur with long-term cover crop residue
25 deposition and different plant species.

Keywords: Avena strigosa, Lolium multiflorum, N cycling; ¹⁵N recovery, Prunus persica (L.)
Batsch.

29

30 1 Introduction

Nutrient management in orchards has a great impact on plant growth, yields, and fruit quality. Sustainable soil fertility management in orchards requires not only a fine-tuning of fertilizer rates, but also a higher use efficiency of nutrients already present in soil (Tagliavini 2012). There is a multitude of factors, including the cycling of plant residues, that dictate nutrient availability in the soil system. A key step to improving nutrient use efficiency is understanding the nutrient's fate during leaf-litter decomposition on the soil surface and the ability of that nutrient to become available for plant uptake (Tagliavini and Scandellari 2013).

38 Nitrogen (N) is a nutrient with a great impact on growth and reproductive development 39 in fruit trees, such as peach [Prunus persica (L.) Batsch.], as it directly affects the flowering 40 quality, fruit set, fruit quality, and yield potential. However, mature peach trees tend to absorb 41 only small amounts of the N derived from fertilization (Policarpo et al. 2002) as the majority 42 of the N used by the new vegetative organs comes from internal N reserves. Part of the N-43 fertilizer applied to the orchard soils can be lost by volatilization (especially when using urea 44 as N source) (Roccuzzo et al. 2017), leaching (greater loss potential in sandy soils with low 45 organic matter content) (Lorensini et al. 2012), and surface runoff (especially in areas with 46 steep grades) (Martínez et al. 2006). Moreover, greater losses can occur when the N-fertilizers 47 are not applied at the right time and rate to meet trees' nutritional demand.

48 Cover crops, such as black oat (*Avena strigosa* Schreb.) and ryegrass (*Lolium* 49 *multiflorum* L.), are used in the inter-row space of orchards to dissipate the kinetic energy of 50 rainfall and reduce soil erosion. Throughout tree development, cover crop shoots are frequently 51 cut and deposited on the soil surface contributing to increasing soil organic matter (SOM) 52 content and nutrient cycling (Brunetto et al. 2017; Ferreira et al. 2014).

53 During the decomposition of cover crop residues, part of the carbon (C) present in the 54 plant tissue may remain in the soil, especially from residues with lower lability. However, the 55 majority of the C will probably return to the atmosphere as CO₂, depending on the residue composition and mineralization rate (Oliveira BS et al. 2016; Reichert et al. 2015). Part of the 56 57 plant tissue N can increase soil available N (nitrate or ammonium), which can be absorbed by 58 mature trees. The decomposition of plant material and the release of N from cover crop residues 59 depends on the residue biochemical composition, especially the cellulose and lignin content, but also the C/N, lignin/N, and cellulose/lignin ratios (Carranca et al. 2009). Residues with low 60 61 cellulose content, high lignin content, and high C/N ratio typically have a low decomposition 62 rate and may even temporarily immobilize soil N (Bonanomi et al. 2013). On the other hand, 63 more labile residues with higher cellulose content, lower lignin content, and lower C/N and 64 cellulose/lignin ratios can mineralize N to the soil, increasing the plant-available N forms 65 (Cabrera et al. 2005). The residue decomposition and nutrient release is also dependent on 66 edaphoclimatic characteristics, especially soil texture, moisture, aeration, temperature, and 67 nutrient availability, which directly affect soil microbial activity (Brunetto et al. 2011; Cabrera 68 et al. 2005; Ferreira et al. 2014).

69 Studies investigating the N derived from cover crop residues can be accurately 70 performed using the ¹⁵N isotope dilution technique (Brunetto et al. 2014; Neto et al. 2008; 71 Tagliavini et al. 2007). Annual tree organs such as leaves, twigs of the year, and fruits are major sinks for the absorbed N derived from in-season fertilization (Brunetto et al. 2017), SOM 72 73 mineralization, and decomposing cover crop residues. Part of the N stored in perennial organs 74 is remobilized to the tree meristematic tissues in the subsequent growth cycle, reducing the tree 75 N needs from N fertilizers at this vegetative phase. Even though the contribution of cover crops 76 for plant nutrition has been evidenced for different crops and production systems (Amossé et al. 2014; Brunetto et al. 2011, 2014; Oliveira RA et al. 2016; Ovalle et al. 2010), information
on how the N derived from cover crop residues is redistributed in different peach organs is still
scarce.

We hypothesize that a) ryegrass and black oat residues will present similar decomposition dynamics, and b) the N released will be used by trees and preferentially redistributed to newly formed organs and trunk. Thus, this study aims to evaluate a) the recovery of N derived from the decomposition of black oat and ryegrass shoot residues in mature trees, b) the N absorption dynamics and the N compartmentalization in different tree organs, and c) to assess black oat and ryegrass shoot residue decomposition dynamics in a mature peach orchard.

87

88 2 Materials and methods

89 **2.1 Cover crop cultivation**

Soil for this experiment was collected from the 0-0.20 m depth of a soil classified as a
Rhodic Paleudalf (Soil Survey Staff 2014) in Southern Brazil (29°42'54" S, 53°42'25" W). The
soil was air-dried, ground to pass through a sieve with 2-mm openings, and reserved for
analysis. Selected soil chemical and physical characteristics are shown in Table 1.
Polypropylene pots (20.0 cm x 25.0 cm) with four kilograms of soil were used to cover crop
cultivation.

96 Seeds of black oat and ryegrass were pre-germinated in plastic germination boxes 97 (Gerbox®) by placing the seeds on paper towels and moistening them with distilled water 98 amounting to 2.5 times the weight of the dry paper. The seed boxes were placed in a BOD 99 (Biochemical Oxygen Demand) chamber at a constant temperature of 25°C and a positive 100 photoperiod of 8 h. A germination test was carried out from 01 May to 13 May 2014, when 25 91 germinated seeds of each plant species were sown in the polypropylene pots. Seven days after 102 emergence, the less developed seedlings were thinned, leaving 20 black oat or 20 ryegrass103 plants in each pot.

Urea containing 46.6% N, labeled with 5 atom%¹⁵N, was applied to 15 pots containing 104 black oat and 15 pots containing ryegrass at a rate of 10 g of N m⁻². Unlabeled urea with 45% 105 106 N was applied to an additional 15 pots grown with black oat and 15 pots grown with ryegrass (control plots). Urea, with and without ¹⁵N, was diluted in deionized water and applied to the 107 soil, thereafter pots were irrigated daily with deionized water to maintain the soil at 70% field 108 109 capacity (Casaroli and Jong van Lier 2008) during the experimental period. The N-fertilizer 110 solution application was split six times throughout the cultivation period: 10, 17, 24, 31, 38, 111 and 45 days after thinning. At 18 and 41 days after thinning, 50 mL of a nutrient solution containing 500, 502, 48, 19, 9, and 14000 mg L⁻¹ of B, Mn, Zn, Cu, Mo, and KH₂PO₄, 112 113 respectively, were applied to each pot. Throughout the crop cycle, weeds were removed by 114 hand from the pots.

Sixty-three days after sowing (*i.e.*, 17 July 2014), the shoots of black oat and ryegrass grown with and without ¹⁵N applications were cut close to the soil surface. The cover crop shoot residues were washed and oven-dried at 65 °C until constant weight. The cover crop ¹⁵N-labeled shoot residues were reserved for Experiment 1, while the unlabeled shoot residues were reserved for Experiment 2. Subsamples of both ¹⁵N-labeled and unlabeled residues were ground with a Wiley mill and separated for chemical characterization and ¹⁵N analysis.

121

122 2.2 Experiment 1 - Mature trees N recovery from black oat and ryegrass shoot residue 123 decomposition

124 2.2.1 Experimental layout

125 The experiment was carried out in a mature peach orchard of 'Chimarrita' [*Prunus* 126 *persica* (L.) Batsch] (Raseira et al. 2014) grafted on 'Capdeboscq' [*Prunus persica* (L.) Batsch]

127 rootstock (Mayer et al. 2014). The orchard was established in 2009 in the experimental area of Embrapa Uva e Vinho, located in the city of Bento Goncalves, Southern Brazil (29°09'44" S, 128 51°31'50" W). Before the installation of the orchard, 30 kg P₂O₅ ha⁻¹ (as triple superphosphate) 129 and 40 kg K₂O ha⁻¹ (as potassium chloride) were applied to increase soil-test P and K to 130 optimum levels. Subsequently, the orchard received annual fertilizer rates of 90 kg N ha⁻¹ (as 131 urea), 40 kg P₂O₅ ha⁻¹ (as triple superphosphate), and 80 kg K₂O ha⁻¹ (as potassium chloride) 132 133 to maintain adequate nutrient levels according to the regional production guidelines (CQFS -134 RS/SC, 2004). After the installation of the experimental area, the adjacent plants received no fertilization. The spacing between tree rows and between plants within rows was 4.0 and 1.5 135 136 m, respectively, totaling 1667 plants per hectare. The Y-shaped pruning system was used for 137 trees. The climate of the region is warm temperate (Cfb), according to the Köppen-Geiger 138 classification system, with an average air temperature of 17.1°C and an average annual rainfall 139 of 1755 mm (Fig. 1). The soil is classified as a Typic Hapludalf (Soil Survey Staff 2014) and 140 its selected physical and chemical characteristics are presented in Table 1. The local landscape 141 is gently sloping.

142 Before the experiment set up, the orchard included cover crops, mainly grasses (e.g., black oat and ryegrass). In the evaluation area, however, all vegetation, including roots, was 143 removed, so that no plants were competing with the trees. On 17 July 2014, ¹⁵N-labeled shoot 144 145 residues of black oat and ryegrass (sampled 63 days after sowing) were deposited at the soil surface to an area of 0.96 m² (0.8 m x 1.2 m) surrounding the tree trunk. Each plot was 146 147 composed of three trees that each received 0.13 kg dry matter of either black oat or ryegrass 148 shoot residue, corresponding to 1354 kg of dry matter per hectare. A nylon net (2 mm mesh) 149 covered the residue and was fixed to the soil with metal clamps to prevent residue displacement 150 by the wind, rain, and animals. For the duration of the experiment, non-residual herbicide was applied to the area of residue deposition to manage weeds and avoid ¹⁵N absorption by weeds. 151

152 2.2.2¹⁵N recovery by mature trees

To follow the pattern of ¹⁵N uptake during the mature tree development, the first peach 153 154 leaf sampling was carried out on 17 August 2014 (30 days after cover crop shoot residue 155 deposition), while subsequent leaf sampling was performed monthly from September 2014 to 156 April 2015. At each leaf sampling time, twenty mature leaves per plant were collected from the 157 middle position on twigs of the year, surrounding the tree canopy. The leaves were oven-dried with forced air at 65 °C until constant weight, ground, and reserved for total N and ¹⁵N analysis. 158 159 On 17 December 2014,115 days after cover crop shoot residue deposition, a PVC pipe 160 (250 mm diameter) was inserted into the soil to collect a soil core sample from the 0-0.20 m 161 depth of each plot. Three soil cores were collected in the central position at one side of the trees row, where ¹⁵N-labeled black oat and ryegrass shoot residues were deposited. Each soil core 162 163 was separated into 0.05 m layers (*i.e.*, 0-0.05, 0.05-0.10, 0.10-0.15, and 0.15-0.20 m depths) 164 and combined for a composite sample of each depth. Thereafter, the soil samples were air-dried, ground to pass a 2-mm sieve, and reserved for total N and ¹⁵N analysis. 165

166 On 23 November 2015 (15 months after cover crop shoot residue deposition), the 167 trunks of trees that received cover crop residue were cut close to the soil surface and separated 168 into the stem, twigs of the year, 2-year-old branches, and leaves. The fruits were collected from 169 each tree and weighed to determine fruit yield. Thereafter, ten fruits per tree were randomly 170 selected to determine total N content and ¹⁵N enrichment in the pulp (*i.e.*, mesocarp plus 171 exocarp). The tree roots were removed from the soil, washed with distilled water, and separated 172 into three diameter classes (< 2 mm, 2 - 5 mm, > 5 mm). Root material was reserved for analysis of total N and ¹⁵N enrichment. The same stratified soil sampling procedure and analysis, 173 174 described above, was repeated before peach roots were removed from the soil.



177 a mature peach orchard

178 2.3.1 Experimental layout

179 Experiment 2 was laid out in the same orchard used for Experiment 1. Unlabeled black 180 oat and ryegrass shoot residues were placed in nylon litter bags (2 mm mesh) covering an area 181 of 0.16 m² (0.4 m x 0.4 m) and deposited on 22 July 2014 on the soil surface of planting rows. 182 Each plot was composed of four trees that received either black oat or ryegrass shoot residue. 183 Each tree received 64 g of dry matter from black oat or ryegrass shoot residue, corresponding 184 to 4,000 kg dry matter per hectare. The bags were fixed to the soil with a metal clamp to increase 185 the area of contact with the soil surface and prevent displacement by the wind. The litter bags 186 were collected at 30-day intervals (*i.e.*, 0, 30, 60, 90, 120, 150, and 180 days after deposition), 187 and at each sampling time, four litter bags of each cover crop were removed from the field. In 188 the laboratory, the residues were washed to remove the adhering soil particles and then oven-189 dried at 65 °C, ground, sieved through 2-mm openings, and reserved for further analysis.

190

191 **2.4 Plant and soil analysis**

192 Before cover crop residue deposition in the experiments, a subsample was subjected 193 to sulfuric acid digestion to determine total N, phosphorus (P), potassium (K), calcium (Ca), 194 and magnesium (Mg) contents (Tedesco et al. 1995). Total N was determined by steam 195 distillation (TE-0364, Tecnal, Brazil). Total P was determined in a spectrophotometer (Bell 196 Photonics, 1105, Brazil) at 882 nm (Murphy and Riley 1962). Total K was determined in a 197 flame photometer (Digimed, BM-62, Brazil). Calcium and Mg were determined in an atomic 198 absorption spectrophotometer (PerkinElmer, AAnalyst 200, USA). Total organic carbon (TOC) 199 was determined by wet combustion (Yeomans and Bremner 1988). The determination of lignin, 200 cellulose, and non-structural biomass was performed according to the methodology described 201 by Aber and Martin (1999). Total N and ¹⁵N were analyzed in an elemental analyzer (Thermo Scientific, Flash EA 1112, Milan, Italy) and by isotope-ratio mass spectrometry (Thermo
Scientific, Delta V Advantage, Bremen, Germany), respectively. Cover crop shoot residue
chemical characteristics are presented in Table 2.

The tree organs' tissue and soil samples from Experiment 1 were also analyzed for total N and %¹⁵N following the above-mentioned procedures. The remaining cover crop residues from litter bags in Experiment 2 were also analyzed for C content, lignin, cellulose, and non-structural biomass following the above-mentioned methodologies. The decomposition of residue dry matter and the release of C, N, cellulose, lignin, and non-structural biomass were estimated by subtracting the initial content from the amount determined after each sampling time.

212

213 **2.5 Calculations and statistical analysis**

Atom% ¹⁵N excess in soil and plant tissue samples was calculated based on the natural ¹⁵N abundance (Mariotti 1983). The N derived from residue (Ndfr) and the N derived from soil (Ndfs) was calculated by the following equations:

217

218
$$15 N$$
 excess in sample (%) = % $15 N$ in sample - 0.3663% (Eq. 1)

219

220
$$Ndfr(\%) = (\% \stackrel{15}{\square} N \text{ excess in sample } / \% \stackrel{15}{\square} N \text{ excess in residue}) x 100$$
 (Eq. 2)

221

222
$$Ndfs(\%) = 100 - Ndfr$$
 (Eq. 3)

223

The residual percentage of each variable (total C and N, cellulose, lignin, and nonstructural biomass) were adjusted by the exponential mathematical model described by Wieder and Lang (1982): 227

228
$$X = X_0^{(-kt)}$$
 (Eq. 4)

229

Where: X is the amount of dry matter or nutrient remaining in the residue after a period
t (days); X₀ is the initial amount of dry matter or nutrient in the residue; k is the decomposition
constant.

The half-life $(t^{\frac{1}{2}})$ was calculated with the value of k (Paul and Clark 1996) (Equation 5). The $t^{\frac{1}{2}}$ expresses the time required for half of the residue to decompose and half of the nutrients contained in the residue to be released.

236

237
$$t^{1/2} = 0.693/k$$
 (Eq. 5)

238

239 All data were submitted to normality and homogeneity of variance by the Lilliefors and Shapiro-Wilk tests prior to the analysis of variance (ANOVA). The dry matter, atom% ¹⁵N, 240 total N (mg tree⁻¹), and total soil N (Experiment 1) variables were transformed [log10(x)] to fit 241 242 a normal distribution before running the ANOVA. Experiment 1 had a completely randomized 243 block design with five replicates. ANOVA was performed to determine the influence of black oat and ryegrass shoot residues on leaf total N, atom%¹⁵N excess, and Ndfr for each leaf's 244 sampling time. ANOVA was also conducted to determine the influence of black oat and 245 ryegrass shoot residue, tree organs, and their interaction on dry matter, total N, atom% ¹⁵N 246 247 excess, Ndfr, and Ndfs. Experiment 2 had a completely randomized block design with five 248 replicates. ANOVA was performed to determine the influence of black oat and ryegrass shoot 249 residues on the remaining dry matter, TOC, total N, cellulose, lignin, and non-structural 250 biomass decomposition constant rate (k) and half-life (t¹/₂). Means were compared by the Tukey-test (p < 0.05). When no difference was observed among cover crop species, the mean 251

value of the treatments was compared. ANOVA was also conducted to determine the influence of black oat and ryegrass shoot residues, residue deposition time, and their interaction on C/N ratio, dry matter, TOC, and total N. When the interaction was significant, two times the standard error of means (SEM) was used as the minimum difference between means statistically different for $p \le 0.05$.

257

3 Results

259 3.1 Experiment 1 - Mature trees N recovery from black oat and ryegrass shoot residue 260 decomposition

261 Following the pattern of leaf response of mature trees cultivated in soils with black oat and ryegrass ¹⁵N-labeled shoot residue deposition, the highest N concentration was found in 262 trees cultivated with ryegrass shoot residue, especially in November and December 2014 (Fig. 263 264 2a). However, the highest leaf N concentration occurred in August 2014, shortly after the 265 deposition of both cover crop residues on the soil surface. Thereafter, leaf N concentration decreased over time in all trees. The highest atom% ¹⁵N and the Ndfr were observed in 266 267 September in leaves of trees that received the deposition of black oat residue (Fig. 2b and 2c). In subsequent evaluations, there were no differences for leaf concentration of total N, atom % 268 ¹⁵N, and % Ndfr between trees cultivated under both cover crop treatments. 269

After 75 weeks (*i.e.*, December 2015), the dry matter, atom % ¹⁵N, and Ndfr of mature peach organs did not differ between the two treatments, but significant differences were observed within tree organs (Table 3). The highest total N concentration occurred in mature tree leaves. The highest dry matter yield was observed in the peach trunk, followed by roots > 5 mm and leaves (7677.6, 2243.1, and 1212.2 g tree⁻¹, respectively). Mature trees leaves and trunk presented the highest Ndfr (mg tree⁻¹) and Ndfs (mg tree⁻¹) while the highest atom % ¹⁵N excess and % Ndfr were observed in roots < 2 mm. 277 Soil characteristics were not affected by the type of cover crop residue, but differences 278 were observed among the soil layers (Table 4). For the mean effect of crop residues, the soil 279 samples collected at 23 and 75 weeks (i.e., 115 and 375 days, respectively) after cover crop residue deposition showed the highest atom% ¹⁵N and Ndfr at the 0-0.05 m depth (Table 4), 280 281 with little variation among other soil depths. It should be noted that 23 weeks after the deposition of black oat and ryegrass shoot residues, atom%¹⁵N and Ndfr values were higher 282 283 than the values observed 75 weeks after the residue deposition. Overall, the percentage of soil 284 Ndfr below the 0.05-m depth was low.

285

3.2 Experiment 2 - Black oat and ryegrass shoot residue decomposition and N release in a mature peach orchard

288 The temporal dynamics of shoot residue dry matter, TOC, N, lignin, cellulose, and 289 non-structural biomass contents were explained by the exponential decay model (Eq. 4). Cover 290 crop residue dry matter and TOC, N, cellulose, lignin, and non-structural biomass contents 291 decreased rapidly during the experimental period, with slight differences between cover crop 292 species (Fig. 3). At the end of the trial (i.e., 150 days after the residue deposition), only 293 approximately 15% of the cover crop residue dry matter and TOC remained in the soil surface 294 (Fig. 3a and 3b, respectively). There was an 82 and 79% decrease in the N content of black oat 295 and ryegrass remaining residues, respectively (Fig. 3c). Lignin and cellulose presented 296 comparable decomposition dynamics, with remaining percentages close to zero after 150 days 297 of residue decomposition. The remaining non-structural biomass followed a similar trend as 298 dry matter, TOC, and total N, with slightly higher values (20 and 30% for black oat and ryegrass 299 shoot residues, respectively) at the end of the experiment (Fig. 3f). The highest k values for dry matter, TOC, cellulose, and lignin were observed for ryegrass shoot residue (Table 5). Black 300 301 oat residue presented the highest $t^{\frac{1}{2}}$ values for dry matter, TOC, total N, cellulose, and lignin

(Table 5). There was no statistical difference for k and $t^{\frac{1}{2}}$ values of the non-structural biomass 302 in both cover crop residues. The C/N ratio of litter bag residues over time was greater in black 303 304 oat than in ryegrass, except at 90 and 180 days after deposition, when the two cover crops 305 presented similar values (Table 6). The C/N ratio constantly decreased until 90 days of 306 deposition, then remained stable for a month and slightly increased at the end of the trial. Black 307 oat shoot residue showed greater dry matter than ryegrass, with values decreasing over time in 308 both cover crops. Total organic C decreased across time and was different between cover crop 309 species. Total organic C of black oat shoot residue was higher than ryegrass from 30 to 120 310 days after deposition; at the beginning of the experiment, ryegrass showed higher values than 311 black oat, while at the end of the experiment (150 and 180 days after deposition), the values 312 were similar between the two cover crops species. Black oat total N content was higher than 313 ryegrass at 0, 30, 90, and 120 days after deposition; in other sampling times, the situation was 314 inverted. The remaining residue N content decreased over time, the only exception was black 315 oat that showed a slight increase 90 days after deposition and then again decreased until the end 316 of the trial (Table 6).

317

318 4 Discussion

319 Different from annual crops, perennial plants, such as fruit trees, use two main sources 320 of N for their vegetative growth and reproduction: the root N uptake and the internal N cycling 321 (Carranca et al. 2018). This behavior can be observed in our study by the highest total N 322 concentration in mature tree leaves at the first sampling time, which is probably related to the 323 plant's internal N reserves. Likewise, the highest N derived from cover crop residues measured 324 in leaves at 30 and 60 days after black oats and ryegrass shoot deposition, respectively, indicates 325 that the cover crop residues were rapidly decomposing and releasing N to the soil, which was 326 taken up by trees during an active absorption phase. This timing coincided with the flowering 327 stage when plants increase young root growth, which is responsible for the absorption of water 328 and nutrients, including soil N (Bravo et al. 2012). The trees' superficial roots, which were in 329 proximity with the cover crops' decomposing residues, may have contributed to a greater extent 330 to the N absorption at this stage. Furthermore, this behavior indicates that synchronizing cover 331 crop termination (*e.g.*, mowing) so that a greater decomposition rate occurs in a period of high 332 tree N demand can contribute to enhancing N recovery.

333 A higher soil N availability favors plant dry matter production by increasing the 334 formation and growth of new vegetative organs such as leaves and twigs, which have a high 335 nutritional N demand due to intense cell division and elongation (Nario et al. 2003; Roccuzzo 336 et al. 2017). However, the highest leaf N content in mature trees at different phenological stages 337 was mostly derived from other N forms than cover crop shoot residue decomposition. This is 338 because the mineral N fertilization that the orchard received over the years may have had a 339 residual effect on our study. The low N recovery from the cover crop shoot residue 340 decomposition by the peach organs might also be related to N losses from the plant-soil system, 341 especially by denitrification, runoff, and leaching (Carranca et al. 2018; Roccuzzo et al. 2017) 342 as only a small proportion of Ndfr (below 0.8%) remained in the soil. On the other hand, the 343 high N content in newly formed organs may also be a response to internal N remobilization.

344 The similar behavior observed for the black oat and ryegrass shoot residue 345 decomposition dynamics, dry matter, organic C, N, cellulose, lignin, and non-structural 346 biomass contents is related to the residue initial lignin concentration and C/N ratio (Table 2). 347 The decrease of dry matter, organic C, N, cellulose, lignin, and non-structural biomass that 348 occurred over time is mediated by the activity of soil fauna and the degradation by the microbial 349 population (Carranca et al. 2009; Nguyen and Marschner 2017; Oliveira RA et al. 2016). During 350 the decomposition period, cover crop residues had a C/N ratio below 20 (Table 6) at all 351 sampling times, which facilitates the colonization and mineralization by microbial population 352 (Ferreira et al. 2014; Oliveira RA et al. 2016). Consequently, the cover crop shoot residue dry 353 matter reduction over the 150 days of the trial contributed to the decrease of soil cover (this if 354 there is no deposition of residues in short intervals of time) and the potential to dissipate the 355 kinetic energy of the raindrops, leaving the soil more susceptible to wind and water erosion 356 (Ferreira et al. 2014; Oliveira RA et al. 2016), and therefore N loss by runoff. This process was 357 probably even more intense in the surrounding area of the tree, without cover crop residues. 358 Frequent rainfall and warm temperatures during spring and summer are known to increase soil 359 microbial activity and therefore intensify residue decomposition rate (Chen et al. 2020). Hence, 360 continuous cover crop residue deposition in orchards is paramount to increase both soil cover 361 and SOM content, while reducing nutrient losses over time, which ultimately will positively 362 affect N use efficiency.

363 Using cover crops in orchards is economically feasible since, in addition to the 364 protection against soil erosion, the residue deposition can contribute to the cycling of different 365 nutrients, increasing soil health and potentially reducing the costs associated with fertilizer 366 application. On average, during the experimental period, black oat and ryegrass residues released an equivalent of 120 kg N ha⁻¹ and 136 kg N ha⁻¹, corresponding to 261 and 296 kg 367 368 urea ha⁻¹, respectively. However, this is a substantial amount of N released in 150 days for the 369 present climatic conditions, and therefore a major part of the N derived from cover crop residue 370 was probably lost from the soil-plant system, which is supported by the low N recovery by 371 mature trees. The greater N recovery by mature trees during the first 60 days was related to the 372 50% N released from both cover crops during this period (see Table 6 and Fig. 3), which 373 corresponds to the flowering phase where new roots are active and are responsible for water 374 and nutrient uptake (Bravo et al. 2012). Furthermore, during this phase the trees present an 375 intensive leaf growth, which increases leaf area and consequently surface area for transpiration, 376 resulting in a greater accumulation of minerals within the plant body. These findings agree with those observed in other fruit studies, such as the grapevines, in soils amended with other typesof residue (Brunetto et al. 2011, 2014).

The overall contribution of cover crop residue to peach nutrition in our study was low (below 1%). However, a greater contribution might occur with different cover crop species and residue management that synchronizes the residue N release with trees nutrient demand. It is also important to highlight that even with a low contribution to N nutrition in a short term, cover crops play an important role in cycling other nutrients such as P, K, and S, increasing SOM content and reducing soil and water loss in the system, which is paramount for sustainable food production.

386

387 **5** Conclusions

388 The highest percentage of N derived from the decomposition of black oat and ryegrass 389 shoot residues occurred in leaves at the flowering stage, *i.e.*, about 30 to 60 days after the shoot 390 residues deposition on the soil surface. The majority of peach leaf N and the N in other young 391 tree organs was derived from other N sources than the cover crop residues, such as soil available 392 N, organic matter, plant internal N reserves, or even the residual effect of mineral fertilizers 393 applied to the trees adjacent to the study area. This is partially explained by the shoot residue 394 decomposition rate on the soil surface, which was reduced to 25-50% during the flowering 395 period.

Most of the N contained in leaves in the first evaluation cycle and annual and perennial organs in the second evaluation cycle was also derived from other N sources than cover crop decomposing shoot residue. It is expected that cover crop grasses with a similar C/N ratio to black oat and ryegrass will have comparable residue decomposition dynamics and contribution to tree N nutrition.

401

Further studies need to be performed to evaluate other cover crop species and residue

- 402 management to synchronize cover crop N mineralization with the stages of greater plant N
- 403 demand to avoid N losses and increase the N recovery and plant nutritional status.

- 405 **Conflict of Interest** The authors declare that they have no conflict of interest.
- 406
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525	Table 1 Selected phy	sical and chemical	characteristics of e	experimental	soils at the 0-0.20 m

526 depth

Soil properties	Unit	Experiment1 and 2
		0-0.20 m layer
Clay (pipette method)	g kg ⁻¹	333
Silt (pipette method)	g kg ⁻¹	405
Sand (pipette method)	g kg ⁻¹	262
Organic C (Walkley and Black 1934)	g kg ⁻¹	17.3
pH in H ₂ O (1:1)	-	5.2
Total N (Kjeldahl method)	g kg ⁻¹	2.0
NO ₃ ⁻ -N (extracted by KCl 1 mol L ⁻¹)	mg kg ⁻¹	34
NH4 ⁺ -N (extracted by KCl 1 mol L ⁻¹)	mg kg ⁻¹	56
Alkaline hydrolizable N (Roberts et al. 2009)	mg kg ⁻¹	183
Aluminum (exchangeable) (extracted by KCl 1 mol L-1)	mg kg ⁻¹	15
Magnesium (exchangeable) (extracted by KCl 1 mol L ⁻¹)	mg kg ⁻¹	150
Calcium (exchangeable) (extracted by KCl 1 mol L ⁻¹)	mg kg ⁻¹	760
Phosphorus (available) (extracted by Mehlich 1)	mg kg ⁻¹	15
Potassium (available) (extracted by Mehlich 1)	mg kg ⁻¹	100

528	Table 2 Chemical characterization of black oat and ryegrass shoot residues at the beginning of
529	the experiment, and the amount of dry matter and nutrients added to the soil for residues
530	decomposition (0-20 cm depth).

Variable	Black oat	Ryegrass
	g	g kg ⁻¹
TOC ^a	463.2 ± 3.5^{k}	431.9±2.0
Total N ^b	42.9±07	47.1±0.6
P°	4.3±0.2	4.0±0.2
K ^d	24.6±0.4	27.9±1.7
Ca ^e	4.9±0.2	5.4±0.1
Mg^{f}	5.8±0.1	5.6±0.1
Cel ^g	388.9±0.6	444.7±1.2
Lig ^h	141.4±0.3	140.5±1.8
Bio ⁱ	569.7±0.3	414.8±0.8
C/N	10.8±0.2	9.2±0.1
Lig/N	1.0 ± 0.01	3.0±0.04
C/P	109.0±5.9	108.3±5.7
Cel/Lig	$9.4{\pm}0.8$	3.2±0.6
At % N ¹⁵	40.1±0.1	36.0±0.04
	Amount of residue and nutrient	ts added to the soil surface (kg ha ⁻¹)
DM ^j	5176.8	5607.4
TOC	220.1	228.9
Ν	13.6	14.9
Р	2.4	2.2
Κ	20.1	22.0
Ca	2.0	2.5
Mg	2.1	2.5

^aTotal organic carbon; ^bTotal nitrogen; ^cTotal phosphorus; ^dTotal potassium; ^eTotal calcium; ^fTotal magnesium;
 ^gCellulose; ^hLignin; ⁱNon-structural biomass; ^jDry matter; ^kmean standard error (n = 3)

Table 3 Dry matter, total N, atom 15 N excess, 15 N derived from residues (Ndfr), and N 533

534 derived from other sources (Ndfs) in mature peach trees organs after 375 days of black oat

C	Peach tree organs								CT I
Cover crop	Pulp	Leaves	Twigs of the year	Branches of the year	Trunk	Roots > 5 mm	Roots 2 - 5 mm	Roots < 2 mm	CV (%)
				Dry	matter (g tree	-1)			
Black Oat	54.43	1243.18	523.14	1163.23	6387.33	1891.52	137.58	145.95	
Ryegrass	55.33	1181.28	556.66	985.35	8967.83	2594.65	111.17	98.33	
Average	54.88f ⁽¹⁾	1212.23bc	539.90d	1074.29c	7677.58a	2243.08b	124.37e	122.14e	6.12 ⁽²⁾
				Т	otal N (g kg ⁻¹)				
Black Oat	11.16	26.56	7.44	4.07	2.82	5.90	8.04	8.65	
Ryegrass	12.18	27.87	7.69	4.47	3.12	8.83	11.22	10.54	
Average	11.67b	27.22a	7.60d	4.27e	2.98f	7.37d	9.63c	9.60c	7.36
				Tot	tal N (mg tree ⁻	1)			
Black Oat	575.39	33012.08	3839.95	4733.8	18244.51	11110.02	1109.17	1257.05	
Ryegrass	671.81	32957.52	4288.27	4420.87	28052.09	22870.01	1228.49	1027.03	
Average	623.60e	32984.80a	4064.11c	4577.33c	23148.30ab	16990.01b	1168.83d	1142.04d	5.5
				aton	n ¹⁵ N excess (9	%)			
Black Oat	0.019	0.0138	0.0164	0.0211	0.0156	0.0216	0.0279	0.0387	
Ryegrass	0.017	0.0170	0.0154	0.0190	0.0151	0.0172	0.0266	0.0364	
Average	0.0180c	0.0154c	0.0159c	0.0200bc	0.0153c	0.0194bc	0.0272b	0.0376a	
		1:	⁵ N derived f	rom shoot re	sidues (Ndfr)	(% total N in	the organ)		
Black Oat	0.47	0.35	0.41	0.53	0.39	0.54	0.7	0.96	
Ryegrass	0.42	0.42	0.38	0.47	0.37	0.43	0.66	0.91	
Average	0.45 bc	0.38 c	0.39 c	0.50 bc	0.38 c	0.48 bc	0.67 ab	0.94 a	
			¹⁵ N de	rived from s	hoot residues ((Ndfr) (mg tre	ee ⁻¹)		
Black Oat	2.69	110.88	15.99	25.17	73.86	60.21	7.98	12.01	
Ryegrass	2.84	140.2	16.46	21.09	104.39	98.99	8.4	8.92	
Average	2.76e	125.54a	16.22bc	23.13b	89.12a	79.60a	8.19cd	10.46bc	21.1
		1	⁵ N derived t	from other so	ources (Ndfs) (% total N in t	the organ)		
Black Oat	99.53	99.65	99.59	99.47	99.61	99.46	99.3	99.04	
Ryegrass	99.58	99.58	99.62	99.53	99.63	99.57	99.34	99.09	
Average	99.56ab	99.62a	99.60a	99.50ab	99.62a	99.52ab	99.32b	99.06c	
			¹⁵ N de	erived from o	other sources (Ndfs) (mg tre	e ⁻¹)		
Black Oat	572.71	32901.20	3823.96	4708.63	18170.65	11049.81	1101.19	1245.04	
Ryegrass	668.97	32817.32	4271.81	4399.78	27947.71	22771.01	1220.09	1018.12	
Average	620.84e	32859.26a	4047.89c	4554.20c	23059.18ab	16910.41b	1160.64d	1131.58d	5.49

and ryegrass shoot residues deposition on the soil surface (Experiment 1) 535

536 537 ⁽¹⁾Means (n = 5) followed by the same lowercase letter in the column, for each plant organ, do not differ by the Tukey-test (p > 0.05).⁽²⁾Coefficient of variation (CV) of cover crop shoot residue error 1

- 538 **Table 4** Soil total nitrogen (TN) concentration, atom ¹⁵N excess, and ¹⁵N derived from residues
- 539 (Ndfr) in the 0-0.20 m depth of a mature peach tree orchard with black oat and ryegrass shoot

Cover crop	Soil depth (m)								
Cover crop	0-0.05	0.05-0.10	0.10-0.15	0.15-0.20	CV (%)				
		23 Weeks (i.e., 115 days)							
		Total N ((g kg ⁻¹)						
Black Oat	4.1	3.3	2.0	1.8					
Ryegrass	4.7	2.6	2.0	1.3					
Average	$4.4 a^{(l)}$	2.9 b	2.0 c	1.6 c	6.22(2)				
		¹⁵ N (atom%	⁵ N excess)						
Black Oat	0.0224	0.0097	0.0056	0.0051					
Ryegrass	0.0406	0.0098	0.0057	0.0050					
Average	0.0215 a	0.0097 b	0.0056 b	0.0050b	0.40				
	Ndfr (%)								
Black Oat	0.56	0.24	0.14	0.13					
Ryegrass	1.01	0.24	0.14	0.13					
Average	0.79 a	0.24 b	0.13 b	0.13 b	5.92				
		75 Weeks (i.e., 375 days)							
		Total N ((g kg ⁻¹)						
Black Oat	4.6	3.4	2.4	1.9					
Ryegrass	4.0	3.0	2.3	1.9					
Average	4.3 a	3.2 b	2.3 c	1.9 с	5.66				
	¹⁵ N (atom% ¹⁵ N excess)								
Black Oat	0.0074	0.0041	0.0052	0.0257					
Ryegrass	0.0263	0.0058	0.0051	0.0044					
Average	0.0169 a	0.0050 b	0.0051 b	0.0051 b	0.64				
		Ndfr	(%)						
Black Oat	0.18	0.10	0.13	0.14					
Ryegrass	0.66	0.15	0.13	0.11					
Average	0.42 a	0.12 b	0.13 b	0.13 b	9.83				

540 residues deposition (Experiment 1)

541 ⁽¹⁾Means (n = 5) followed by the same lowercase letter within columns not differ by Tukey test (p < 0.05). 542 ⁽²⁾Coefficient of variation (CV) of cover crop residue error 1

Table 5 Dynamics of black oat and ryegrass shoot residues decomposition in a mature peach orchard. Remaining dry matter, total organic carbon, nitrogen, cellulose, lignin, and nonstructural biomass were adjusted to the model $X = Xo^{(-kt)}$; where X is the amount of dry matter or nutrient remaining in the residue after a period t (days), X₀ is the initial amount of dry matter or nutrient in the residue, k is the decomposition constant rate and t¹/₂ is the half-life for each compartment (Experiment 2)

Treatment	k	$t^{1/2}$	R ²⁽³⁾	k	t ^{1/2}	R ²
	g g ⁻¹	days	-	g g ⁻¹	days	-
	Rem	aining dry ma	atter	Remaining cellulose		
Black Oat	0.0164 b ⁽¹⁾	42 a	0.98^{*}	0.0278 b	25 a	0.95^{*}
Ryegrass	0.0208 a	33 b	0.98^{*}	0.0356 a	19 b	0.98^{*}
CV (%) ⁽²⁾	2.61	2.72		9.05	5.34	
	Remainin	g total organi	c carbon	Re	emaining ligni	in
Black Oat	0.0170 b	41 a	0.97^*	0.0325 b	23 a	0.88^*
Ryegrass	0.0211 a	33 b	0.97^*	0.0614 a	12 b	0.98^{*}
CV (%)	2.26	2.33		13.62	14.16	
	Ren	naining nitrog	gen	Remaining	g non-structura	al biomass
Black Oat	0.0118 a	54 a	0.94^{*}	0.0117 a	59 a	0.91*
Ryegrass	0.0129 a	59 a	0.97^{*}	0.0459 a	50 a	0.93*
CV (%)	6.06	11.38		41.36	29.48	

549 ⁽¹⁾Means (n = 5) followed by the same lowercase letter do not differ by the Tukey test (p < 0.05). ⁽²⁾ Coefficient

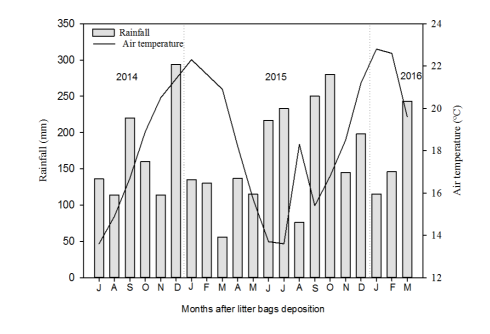
of variation (CV) of the decomposition constant rate (k) and the half-life for each compartment $(t\frac{1}{2})$. (3)

551 Coefficient of determination (\mathbb{R}^2) of the residue's decomposition dynamics model. * significant at p < 0.05

Table 6 Carbon/nitrogen (C/N) ratio, dry matter (DM), total organic carbon (TOC), and nitrogen (N) in the remaining shoot residues of black oat and ryegrass deposited on the soil surface in mature peach trees planting rows (Experiment 2)

Treatments	Day after litter bags deposition on the soil surface								
-	0	30	60	90	120	150	180		
				C/N ratio					
Black oat shoot	16.3	8.7	7.0	5.5	5.8	6.7	6.1		
Ryegrass shoot	15.7	6.6	4.3	5.1	5.0	7.6	5.9		
Significance ⁽¹⁾			-	2 SEM = 0.479					
CV (%)	8.20	15.3	23.9	24.4	23.0	40.8	10.		
				DM (kg ha ⁻¹)					
Black oat shoot	5177	2989	1573	1277	1120	745	161		
Ryegrass shoot	5607	2702	1277	1078	838	825	142		
Significance				2 SEM = 61.0					
CV (%)	3.79	7.13	16.1	9.72	12.1	20.3	26.		
				TOC (kg ha ⁻¹)					
Black oat shoot	2201	1126	562	512	443	290	44.		
Ryegrass shoot	2289	976	420	419	327	280	69.		
Significance				2 SEM = 35.8					
CV (%)	16.80	7.62	16.81	9.79	11.56	23.53	3.1		
				N (kg ha ⁻¹)					
Black oat shoot	136	131	74.8	94.9	77.2	42.7	9.9		
Ryegrass shoot	149	150	88.8	83.6	61.1	47.2	12.		
Significance				2 SEM = 4.12					
CV (%)	9.85	13.2	7.34	18.4	20.6	9.00	23.		

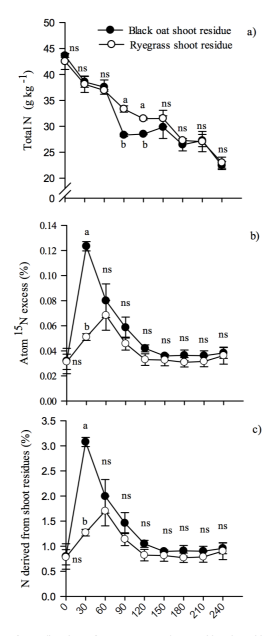
555 ⁽¹⁾ Values differing by 2 standard error of means (SEM) are statistically different



556

557 Fig. 1 Average air temperature and cumulative monthly rainfall after the deposition of litter

558 bags in the experimental area



559

Leaf sampling time after cover crop shoot residue deposition (days)

Fig. 2 Total N a), atom% ¹⁵N excess b), and ¹⁵N derived from shoot residues c) in mature peach trees leaves grown with black oat and ryegrass shoot residues deposition on the soil surface (Experiment 1). Vertical bars indicate the standard error of the mean (n = 3). Lowercase letters compare black oat and ryegrass residues within each sampling time by the Tukey-test (p < 0.05), ns = not significant

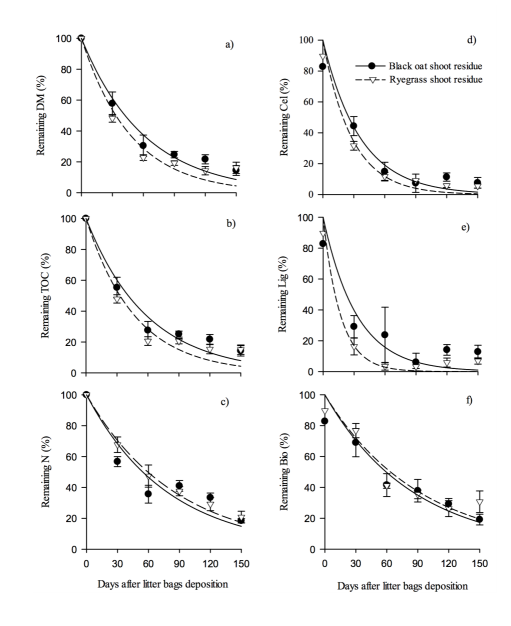


Fig. 3 Remaining percentage of dry matter (DM) a), total organic carbon (TOC) b), nitrogen
(N) c), cellulose (Cel) d), lignin (Lig) e), and non-structural biomass (Bio) f) in black oat and
ryegrass shoot residues deposited on the planting rows of a mature peach tree orchard
(Experiment 2)