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1 **The fate of cumulative applications of ¹⁵N-labelled fertiliser in perennial**
2 **and annual bioenergy crops**

3

4 **List of authors:**

5 Fabien Ferchaud^{1*}, Guillaume Vitte¹, Jean-Marie Machet¹, Nicolas Beaudoin¹, Manuella
6 Catterou² and Bruno Mary¹

7 ¹ *INRA, UR 1158 AgroImpact, site de Laon, F-02000 Barenton-Bugny, France*

8 ² *Université de Picardie Jules Verne, Unit CNRS-FRE 3498 Edysan, 33 rue Saint Leu, F-*
9 *80039 Amiens, France*

10

11 *Corresponding author:

12 INRA, UR 1158 AgroImpact

13 Pôle du Griffon

14 180 rue Pierre-Gilles de Gennes

15 02000 Barenton-Bugny FRANCE

16 Tel.: +33 323240775, fax: +33 323240776

17 Email: fabien.ferchaud@laon.inra.fr

18 **Abstract**

19 The fate of nitrogen (N) fertiliser applied to bioenergy crops is a key issue to allow high
20 biomass production while minimising environmental impacts due to N losses. The aim of this
21 study was to follow the fate in the soil-plant system of N fertiliser applied to perennial
22 (*Miscanthus × giganteus* and switchgrass), “semi-perennial” (fescue and alfalfa) and annual
23 (sorghum and triticale) bioenergy crops. Crops received ¹⁵N-labelled fertiliser (urea
24 ammonium nitrate solution) during 4 or 5 successive years on the same subplots, at a rate
25 varying from 24 to 120 kg N ha⁻¹ yr⁻¹. Biomass production, N and ¹⁵N removal at harvest were
26 measured each year. The ¹⁵N recovery in crop residues, non-harvested crop parts and soil was
27 measured at the end of the ¹⁵N-labelling period. Perennial crops had higher biomass
28 production but generally lower ¹⁵N recovery in harvested biomass than other crops,
29 particularly when harvested late (end of winter). At the end of the 4 or 5-year period, the
30 proportion of ¹⁵N recovered in harvested biomass was 13-34% for perennials, 23-38% for
31 semi-perennials and 34-39% for annual crops. Perennial crops stored large amounts of N in
32 their belowground organs; the mean ¹⁵N recovery in these organs was 12%, corresponding to
33 a N storage flux of 14 kg N ha⁻¹ yr⁻¹. The ¹⁵N recovery in soil (including crop residues) was
34 higher for perennials (average 36%) than semi-perennials (28%) and annual crops (19%),
35 corresponding to a N immobilisation rate of 43, 15 and 12 kg N ha⁻¹ yr⁻¹ respectively. The
36 mean overall ¹⁵N recovery in the soil-plant system was 69% in perennials, 61% in semi-
37 perennials to 56% in annual crops, suggesting that important fertiliser losses occurred through
38 volatilisation and denitrification. Perennial bioenergy crops had the better efficiency by
39 storing fertiliser-N in soil organic matter and living belowground biomass used as N reserves
40 for succeeding years.

41 **Keywords**

42 bioenergy, nitrogen fertiliser, ^{15}N , nitrogen use efficiency, miscanthus, switchgrass

43

44 **Highlights**

45 • The fate of ^{15}N -labelled fertilizer was compared in bioenergy crops over 4-5 years

46 • Perennial crops exported the smallest amounts of ^{15}N in harvested biomass

47 • They stored the highest amounts of ^{15}N in their belowground organs and soil+litter

48 • The overall ^{15}N recovery was greater in perennials than other crops

49 • The early harvested miscanthus gave the highest overall ^{15}N recovery

50 **1. Introduction**

51 Bioenergy production from crops has been supported to contribute to the production of
52 renewable energy in response to the challenges of climate change and depletion of fossil
53 resources (Don *et al.*, 2011). However, the use of conventional food crops to produce biofuels
54 has raised a lot of concerns about its environmental consequences (*e.g.* Galloway *et al.*, 2008;
55 Smith and Searchinger, 2012). The large nitrogen (N) requirements of these first generation
56 bioenergy crops may be harmful to the greenhouse gas balance of biofuels (Crutzen *et al.*,
57 2008). The development of new conversion technologies and biorefineries allows considering
58 a wide range of new bioenergy crops (Ragauskas *et al.*, 2006; Sanderson and Adler, 2008;
59 Somerville *et al.*, 2010). Among them, perennial C4 crops such as miscanthus and switchgrass
60 are considered as promising because of their high biomass production with low nutrient
61 requirements and expected low greenhouse gas emissions (Don *et al.*, 2011; Jørgensen, 2011;
62 Monti *et al.*, 2012; Cadoux *et al.*, 2014). However, even with these crops, N fertilisation may
63 still be necessary to maintain high yields and soil fertility on the long term (Cadoux *et al.*,
64 2012; Monti *et al.*, 2012; Cadoux *et al.*, 2014). Fertiliser-N use efficiency of bioenergy crops
65 is therefore a key issue to allow high biomass production while minimising environmental
66 impacts due to N losses.

67 There are various ways to define and measure fertiliser-N efficiency. Two different
68 approaches are widely used in the literature: (1) the apparent recovery which is based on the
69 difference in N uptake between a crop receiving N fertiliser and a reference plot without N
70 applied (*e.g.* Cassman *et al.*, 2002) and (2) the actual recovery or ¹⁵N recovery which is the
71 fraction of labelled N that is taken up by a crop following application of ¹⁵N-labelled fertiliser
72 (Hauck and Bremner, 1976). Both methods can give similar or dissimilar results whether or
73 not the uptake of inorganic soil N is different between fertilised and unfertilised treatments.
74 "Pool substitution" between fertiliser-N and soil mineral N can lead to a higher apparent than

75 actual recovery (Jenkinson *et al.*, 1985). Nevertheless, only the ^{15}N method allows to
76 determine the fate of the fertiliser-N in the different compartments of the agroecosystem (*e.g.*
77 crop and soil) and therefore the overall losses of fertiliser-N (Gardner and Drinkwater, 2009).
78 Few studies have analysed the fate of ^{15}N -labelled fertiliser applied to bioenergy crops.
79 Christian *et al.* (2006) and Pedroso *et al.* (2014) analysed the effect of a single ^{15}N
80 fertilisation pulse during 3 successive years on miscanthus and switchgrass respectively. They
81 found a rather low ^{15}N recovery in the harvested biomass (14-39%) and that belowground
82 organs and soil represented important N sinks. Pedroso *et al.* (2014) also pointed out the
83 effect of crop management, *i.e.* harvest date, on the ^{15}N recovery and partitioning. However,
84 no study has compared the ^{15}N recovery of different bioenergy crops at the same site.
85 Only a few experiments have followed the recovery of ^{15}N in the soil-plant system on the long
86 term. In arable cropping systems, a small proportion of the residual ^{15}N , *i.e.* the labelled
87 fertiliser-N remaining in soil (mainly in organic form) and crop residues after harvest, is re-
88 mineralised each year from the soil organic matter pool and can be recovered by the following
89 crops or lost through N leaching or gaseous losses (Glendining *et al.*, 2001; Macdonald *et al.*,
90 2002; Sebilo *et al.*, 2013). The amount of ^{15}N remaining in the soil-plant system after the year
91 of application is likely to be greater with perennial bioenergy crops than with annual crops
92 because of the presence of perennial organs. The ^{15}N stored in perennial organs can be used
93 by the crop in the subsequent years and partly recovered at harvest, as shown by Christian *et*
94 *al.* (2006) for miscanthus. Using cumulative applications of ^{15}N in the same plots over several
95 growing seasons could allow to integrate part of the long-term fate of the residual ^{15}N and to
96 reduce the variability in plant N uptake and fertiliser-N losses due to climate conditions and
97 agronomical context (*i.e.* age of the crop, other stresses, etc.). In this study, we used
98 cumulative applications of ^{15}N -labelled fertiliser for four or five years to determine (1) the
99 fate of fertiliser applied to perennial, semi-perennial and annual bioenergy crops in the soil-

100 plant system, and (2) the interaction with crop management, *i.e.* harvest date of perennial
101 crops and N fertiliser rate for all crops. We hypothesised that perennial crops would export
102 smaller amounts of ^{15}N through harvests than the other crops but would store larger amounts
103 of ^{15}N in perennial organs and soil organic matter, leading to an equal or improved overall ^{15}N
104 recovery in the soil-plant system.

105 2. Materials and methods

106 2.1. Site and experimental design

107 The study is based on an ongoing long-term experiment established in 2006 at the INRA
108 experimental station in Estrées-Mons, northern France (49.872 N, 3.013 E) called “Biomass
109 & Environment” (B&E). The soil is a Haplic Luvisol (IUSS Working Group WRB, 2006).
110 Soil characteristics are given in Table S1 (Appendix A). Over the period 2006-2011, the mean
111 annual temperature was 10.6 °C, the mean rainfall and potential evapotranspiration were 673
112 and 737 mm yr⁻¹ respectively. Before 2006, the field had been cultivated for many years with
113 annual crops, winter wheat being the most common crop.

114 The experiment was initiated to study the production and the environmental impacts of a wide
115 range of bioenergy crops. It compares eight “rotations”: four with C4 perennial crops
116 (monocultures), two with C3 “semi-perennial” crops (destroyed every two or three years) and
117 two with C3/C4 annual crops (Table 1). The perennial crops are miscanthus
118 (*Miscanthus* × *giganteus* Greef & Deuter ex Hodkinson & Renvoize) and switchgrass
119 (*Panicum virgatum* cv. Kanlow). They are harvested either early in October (E) or late in
120 February (L). The semi-perennial crops are tall fescue (*Festuca arundinacea*) and alfalfa
121 (*Medicago sativa*). Annual crops are fibre sorghum (*Sorghum bicolor* (L.) Moench cv. H133)
122 and triticale (× *Triticosecale* Wittmack). The experiment also includes two nitrogen
123 treatments (N- and N+) with fertiliser-N rates depending on the crops (Table 2). The rationale
124 for defining the N rates was explained by Cadoux *et al.* (2014).

125 The 2.7 ha field was divided into two parts in order to facilitate cropping operations and limit
126 competition between plants due to differences in canopy height: (1) a split-block design in the
127 west part for perennial crops with “rotations” in the main plots (miscanthus early, miscanthus
128 late, switchgrass early, switchgrass late) and N fertilisation rates in the subplots (N- and N+),
129 and (2) a split-plot design in the east part for the other crops with rotations in the main plots

130 (fescue-alfalfa, alfalfa-fescue, sorghum-triticale and triticale-sorghum) and N fertilisation
131 rates in the subplots (N- and N+). Each of the two parts comprised three replicate blocks and
132 24 subplots of 360 m² (Fig. S1, Appendix A). Soil analyses performed in 2006 revealed a
133 slightly higher clay content in the west than in the east part (180 ± 27 vs. 148 ± 19 g kg⁻¹ in
134 the 0-30 cm layer, Table S1 in Appendix A).

135 At the start of the experiment, the field was mouldboard ploughed at a depth of *ca.* 25 cm.
136 After seedbed preparation, miscanthus was planted in April 2006 (1.5 rhizome m⁻²) and
137 switchgrass sown in June 2006 (seed rate = 15 kg ha⁻¹). In 2006, perennial crops were not
138 harvested because of the low biomass production during the first year of growth. Their
139 aboveground biomass was cut and left on the soil surface. Semi-perennial crops were sown in
140 2006, 2009 and 2011, usually in April. Before sowing, the previous crop (alfalfa or fescue)
141 was destroyed in late autumn with a cultivator and a disc harrow (15 cm deep) in 2008 and
142 mouldboard ploughed (*ca.* 22 cm deep) in 2010. These crops were harvested in two or three
143 cuttings depending on years, with the last cut in October. Annual crops were tilled
144 superficially (12-15 cm deep) without inversion ploughing. Sorghum was sown in late May
145 and harvested in late September. Triticale was sown in mid-October and harvested in late July
146 or early August. The N fertiliser was surface-applied from 2007 onwards as UAN solution
147 (urea ammonium nitrate) containing 390 g N l⁻¹ (50% urea, 50% NH₄NO₃). Perennial crops
148 received a single annual application in late April. Fescue received N fertiliser at the beginning
149 of each cycle of regrowth and seedling crops were not fertilised before the first cut, so that the
150 total N rate varied between years. Sorghum was fertilised just before sowing and triticale in
151 March at mid-tillering for N- and N+ treatments and in late April at the beginning of stem
152 elongation for N+. Further details about crop management are given by Cadoux *et al.* (2014).
153 ¹⁵N-labelled UAN fertiliser, uniformly labelled on urea, NH₄ and NO₃, was applied to a
154 subplot of 36 m² located north of each plot from 2007 to 2010 (perennial crops) or 2011

155 (other crops). Simultaneously, the unlabelled UAN was added at the same rate in the rest of
156 the plot. The labelled UAN solution was applied with a CO₂-pressurised hand sprayer to
157 mimic the concentration and volume of liquid fertiliser applied in the rest of the plot. The
158 isotopic excess of the labelled fertiliser varied between treatments in order to apply the same
159 amounts of ¹⁵N per surface area in all treatments (Table 2).

160

161 2.2. Sampling and analytical procedures

162 2.2.1. Aboveground biomass at harvest

163 Harvested crop production was measured every year from 2006 to 2011. On each harvest
164 date, the aboveground biomass was collected manually in one micro-plot inside the ¹⁵N-
165 labelled subplot. The size of the micro-plot depended on the crops, according to the amount of
166 biomass produced per unit area and stand homogeneity: 3.84 m² for miscanthus (six plants),
167 2.5 m² for switchgrass, *ca.* 3.6 m² for sorghum and *ca.* 5 m² for fescue, alfalfa and triticale.
168 The cutting height was 7 cm for all crops. The fresh biomass was weighed and a
169 representative subsample was dried at 65 °C for 96 h to determine the dry matter content and
170 ground before analysis. In order to better take into account canopy variability of miscanthus,
171 the measured biomass was corrected by the number of stems determined in a wider
172 undisturbed area of 25 m² according to Strullu *et al.* (2011). The N concentration and ¹⁵N
173 abundance were determined using an elemental analyser (FLASH EA 1112 series, Thermo
174 Electron, Bremen, Germany) coupled to an isotope ratio mass spectrometer (DELTA V
175 Advantage, Thermo Electron, Bremen, Germany).

176

177 2.2.2. Soil

178 The soil was sampled on two dates: at the beginning of the experiment in May 2006 to
179 measure initial ¹⁵N excess and at the end of the ¹⁵N-labelling period, *i.e.* in March 2011 for

180 perennial crops (west part of the field trial) and March 2012 for other crops (east part of the
181 field trial). Soil cores of 8 cm diameter were extracted with depth increments of 20 cm and
182 inserted into plastic tubes using a powered soil corer (Humax soil sampler, Switzerland). In
183 2006, two soil cores were taken in each plot down to 40 cm depth. In 2011 and 2012, six soil
184 cores were taken in each plot down to 60 cm. All cores were located inside the labelled
185 subplot in a 2.6 m² micro-plot and taken in intra-row and inter-row zones.

186 From 2006, the ploughing depth was reduced from ca. 30-35 cm to less than 25 cm in all
187 treatments. The old ploughing depth (referred to below as *Y*) was identified in the soil cores
188 on each sampling date by detecting changes in soil colour and structure. Soil cores removed
189 from the plastic tubes in the laboratory were divided into five layers (0-5, 5-20, 20-*Y*, *Y*-40
190 and 40-60 cm) in 2011 and 2012. Coarse residues (>2 mm), roots and rhizomes were then
191 carefully removed from the soil by handpicking. Soil samples were dried at 38 °C for 96 h,
192 crushed through a 2 mm sieve, subsampled and finely ground with a ball mill (PM 400,
193 Retsch, Germany) before analysis. Soil samples were analysed for total N concentration and
194 ¹⁵N abundance using an elemental analyser (EURO EA, Eurovector, Italy) coupled to an
195 isotope ratio mass spectrometer (Delta Plus Advantage, Thermo Electron, Germany). Bulk
196 densities were also determined at each sampling date either with steel cylinders or a dual
197 gamma probe (LPC-INRA, France). Full details of the methodology are given by Ferchaud *et*
198 *al.* (2015b).

199

200 2.2.3. *Dead and living crop biomass*

201 In order to make a complete ¹⁵N balance in the soil-plant system, crop residues and living
202 crop biomass were sampled at the same time and location as in the final soil sampling. Crop
203 residues included dead plant parts accumulated in soil or at soil surface whereas living crop

204 biomass was composed of living aboveground material in the case of alfalfa, fescue and
205 triticale and living belowground material (roots and rhizomes) for all crops.

206 Crop residues from perennials present at soil surface were collected just before soil sampling
207 in 2011. Stem bases and fragments (>10 mm) as well as fallen leaves (mulch) of miscanthus
208 late were sampled in the whole micro-plot. Small stem fragments (2 to 10 mm) and leaf debris
209 (for miscanthus late) present at soil surface were collected in six areas of 27 × 27 cm within
210 each micro-plot, corresponding to the location of the soil cores. Stem fragments below soil
211 surface (>2 mm) were collected in the 8 cm diameter cores. Aboveground residues from the
212 six areas were pooled together, as well as belowground residues from the six soil cores. The
213 residues from semi-perennial and annual crops, buried by soil tillage, were collected in the
214 soil cores in 2012. All residues were dried at 65 °C for 96 h, weighed and ground before
215 analysis.

216 The aboveground living biomass of fescue and alfalfa was cut just above the soil surface in
217 each micro-plot before soil sampling in 2012. Triticale and catch crop (before sorghum)
218 plants were pulled from the soil in each micro-plot in order to collect aboveground and part of
219 the belowground biomass and washed to eliminate soil contamination. Roots of all crops
220 (including remaining roots of triticale and catch crop) and perennial crop rhizomes collected
221 in the six cores of each micro-plot were pooled for each layer and washed. Given the very
222 large spatial variability of the rhizome biomass of miscanthus, the method proposed by Strullu
223 *et al.* (2011) was used to quantify it more precisely. It consisted in counting the number of
224 stems of all plants in a given subplot and extracting the entire rhizome of the plant having the
225 median number of stems. All plant samples were dried at 65 °C for 96 h, weighed and ground
226 before analysis.

227 The N concentration and ^{15}N abundance of all samples were determined using an elemental
228 analyser (FLASH EA 1112 series, Thermo Electron, Bremen, Germany) coupled to an isotope
229 ratio mass spectrometer (DELTA V Advantage, Thermo Electron, Bremen, Germany).

230

231 2.3. Calculations

232 2.3.1. Biomass, N content and apparent N recovery

233 For each sampling, the measured biomass was expressed in tons of dry matter per hectare and
234 the N content (kg N ha^{-1}) was obtained by multiplying biomass by N concentration.

235 The annual crop production and harvested nitrogen measured in labelled subplots were
236 compared to the measurements achieved in unlabelled subplots of the experiment and
237 presented in an earlier paper (Cadoux *et al.*, 2014). We found a good relationship between the
238 two estimates although the N content was slightly lower in labelled subplots ($y = 0.98 x$, $R^2 =$
239 0.93 for biomass production; $y = 0.94 x$, $R^2 = 0.87$ for harvested N). This difference was
240 considered acceptable. The apparent recovery of fertiliser-N (R_A , in %) was calculated as
241 follows:

$$242 \quad R_A = \frac{T_N - T_0}{F}$$

243 where T_N and T_0 are the amounts of N in the fertilised (N+) and unfertilised (N-) aboveground
244 crop biomass at harvest (kg N ha^{-1}), respectively, and F is the amount of fertiliser-N applied
245 (kg N ha^{-1}). This calculation was applicable only to the crops whose N- treatment was
246 unfertilised, *i.e.* for perennial crops and sorghum.

247

248 2.3.2. Actual ^{15}N recovery

249 The amount of N derived from the ^{15}N -labelled fertiliser in a given crop part or soil layer
250 (N_{dff} , in kg N ha^{-1}) was calculated according to Hauck and Bremner (1976):

251
$$Ndff = T \cdot \frac{(p - q)}{(f - q)}$$

252 where T is the amount of N in the labelled crop part or soil layer (kg N ha⁻¹), p the ¹⁵N excess
253 atom fraction in the labelled crop part or soil layer, q the ¹⁵N excess atom fraction in control
254 crop or soil that did not receive labelled N and f the ¹⁵N excess atom fraction of the labelled
255 fertiliser. The ¹⁵N recovery was calculated as the ratio between $Ndff$ and the amount of N
256 applied. The mean q value in aboveground biomass at harvest was derived from the analyses
257 made in the unlabelled subplots in 2007 and 2009. For the final crop residues and living crop
258 biomass, q was either obtained from corresponding unlabelled samples (N- treatments for
259 perennial crops) or using the mean value calculated for aboveground biomass. The q values in
260 soil samples were obtained with the measurements made in 2006 in the corresponding plots
261 and soil layers.

262 The $Ndff$ in soil samples were calculated in each treatment on an equivalent soil mass (ESM)
263 basis (Ellert and Bettany, 1995). The “reference” soil masses used for calculations were those
264 measured in 2006 (667, 2000, 2002, 884 and 3137 t ha⁻¹ for 0-5, 5-20, 20-Y, Y-40 and 40-60
265 cm respectively). Detailed calculations are given by Ferchaud *et al.* (2015b) for soil organic
266 carbon stocks and carbon isotopic composition. The same equations were applied here
267 replacing carbon concentration by N concentration and $\delta^{13}\text{C}$ by ¹⁵N excess atom fraction. In
268 the following, soil layers on ESM basis are called L1 to L5.

269

270 2.6. Statistical analyses

271 All statistical analyses were performed using R (R Core Team, 2014). The effects of rotation,
272 nitrogen and their interaction were evaluated by analysis of variance (ANOVA) for the
273 different variables. Two linear mixed-effect models were used: the first one adapted to a split-
274 block design (with blocks, rotation \times blocks and nitrogen \times blocks interactions as random
275 factors) was used for perennial crops and the second, adapted to a split-plot design (with

276 blocks and rotation \times blocks interaction as random factors), was used for the other crops.
277 Rotation, nitrogen and their interaction were treated as fixed factors in both models. The *lme*
278 function from the *nlme* package was used to fit the models (Pineiro *et al.*, 2014). Significant
279 differences ($p < 0.05$) between treatments were detected using the *lsmeans* function (Lenth,
280 2014). The assumptions of ANOVA were checked by visual examination of the residuals
281 against predicted values and using the Shapiro-Wilk and Levene tests. Log-transformed data
282 or Box-Cox transformation were used if necessary to satisfy these assumptions.

283 3. Results

284 3.1. Crop production and N removal at harvest

285 The mean harvested biomass was calculated from 2007 (first year with all crops present and
286 beginning of N applications) to the end of the period during which ¹⁵N-labelled fertiliser was
287 applied, *i.e.* 2010 for perennial crops and 2011 for the other crops (Table 3). The mean
288 harvested biomass represented 19.0 t DM ha⁻¹ yr⁻¹ in perennial crops and 10.3 t DM ha⁻¹ yr⁻¹
289 in other crops. Rotation, fertiliser-N rate and their interaction had a significant effect on
290 biomass production for both crop types (Table S2, Appendix A). Among perennial crops, Mis
291 E N+ was the most productive, yielding 26.6 ± 2.6 t ha⁻¹ yr⁻¹. Miscanthus produced generally
292 more than switchgrass, particularly the early harvest (E) treatments. An interaction between
293 harvest date and N fertilisation was observed: biomass production was significantly higher in
294 N+ than in N- for E treatments whereas N fertilisation had no significant effect for L
295 treatments. Among semi-perennial and annuals crops, Tri-Sor N+ was the most productive
296 treatment with 12.6 ± 1.3 t ha⁻¹ yr⁻¹. The higher level of fertilisation (N+) significantly
297 enhanced biomass production in annual crops compared to N-, but not in semi-perennials
298 crops. Fescue alone had a small and significant response to N rate (+1.5 t ha⁻¹ yr⁻¹ in N+).
299 Annual data are given in Table S3 (Appendix A).

300 The amount of N exported at harvest varied widely, from 38 ± 10 kg ha⁻¹ yr⁻¹ in Mis L N- in
301 2007-2010 to 228 ± 15 kg ha⁻¹ yr⁻¹ in Alf-Fes N+ in 2007-2011 (Table 3). It was significantly
302 affected by rotation, fertiliser-N rate and their interaction (Table S2, Appendix A). Fertilised
303 perennial crops exported systematically higher amounts of N than unfertilised ones (+24 kg
304 ha⁻¹ yr⁻¹ on average). The amount of N exported at harvest was greater in early than in late
305 harvest treatments, particularly with miscanthus. It was high for semi-perennial crops (204 kg
306 ha⁻¹ yr⁻¹ on average) but did not change significantly with N fertilisation. On the contrary,

307 fertilised annual crops (N+) exported more N than low fertilised ones (+50 kg ha⁻¹ yr⁻¹ on
308 average). Annual data are given in Table S4 (Appendix A).

309

310 3.2. Dead and living crop biomass

311 The amount of crop residues found at soil surface or within soil at the end of the ¹⁵N-labelling
312 period was much higher in perennial than in other crops: 13.2 vs. 2.4 t DM ha⁻¹ on average
313 respectively (Table 4). It did not change significantly with N fertilisation for perennials and
314 other crops (Table S2, Appendix A). The amount of aboveground residues was particularly
315 important in Mis L because of the presence of senescent leaves accumulated in mulch at soil
316 surface. The total living crop biomass was also much higher in perennial crops: rhizomes and
317 roots (0-60 cm) of perennial crops represented 12.9 to 24.7 t DM ha⁻¹ (18.5 t ha⁻¹ on average)
318 in March 2011 whereas the total living crop biomass of the other crops was only 0.4 to 6.7 t
319 DM ha⁻¹ in 2012.

320 The N content in crop residues was significantly higher in N+ than in N- for perennial crops
321 but not for the other crops (Table 5). It was higher in Mis L than in the other perennial
322 treatments (105 vs. 48 kg N ha⁻¹ respectively on average). Residues of semi-perennial and
323 annual crops contained 5 to 30 kg N ha⁻¹. Large amounts of nitrogen were stored in rhizomes
324 and roots of perennial crops: 264 kg ha⁻¹ in Mis L and 148 kg ha⁻¹ in Mis E. A greater amount
325 of N in the L treatment was also found for switchgrass but was not significant (p<0.05). N
326 fertilisation increased N stocks in belowground organs of miscanthus and switchgrass by 65
327 kg ha⁻¹ on average. Nitrogen was mainly stored in rhizomes for miscanthus (73%) and in roots
328 for switchgrass (65%). The N content in the living biomass of the other crops ranged from 21
329 kg ha⁻¹ (Sor-Tri: triticale sown in October 2011) to 158 kg ha⁻¹ (Fes-Alf: fescue sown in April
330 2011) and did not differ significantly between N- and N+ (Table S2, Appendix A).

331

332 3.3. *Ndff* and ¹⁵N recovery in the exported biomass

333 The amount of N derived from fertiliser exported at harvest was calculated each year from
334 2007 to 2011 (Table 6). From 2008 onwards, the *Ndff* could derive either from the fertiliser
335 applied during the same year or from preceding applications because the ¹⁵N-labelled fertiliser
336 was applied every year in the same subplots. In unfertilised crops (alfalfa or sorghum N-) following fertilised ones, the *Ndff* was low (between 0.4 and 2.8 kg ha⁻¹), except in 2011 for
337 fescue and alfalfa N+ (5.6 kg ha⁻¹ on average), showing that the carry over effect of fertiliser
338 was much smaller than its direct effect. The *Ndff* tended to increase with time for miscanthus
339 but not for switchgrass. On average over the whole period, *Ndff* in the exported biomass
340 represented 26, 18 and 28 kg N ha⁻¹ yr⁻¹ for perennial, semi-perennial and annual crops
341 respectively. This corresponds to 28-30% of the exported N for perennial crops, 3-17% for
342 semi-perennials and 11-37% for annual crops. The exported N derived from other sources
343 (soil and atmosphere) was greater: 62, 186 and 65 kg ha⁻¹ yr⁻¹ for perennial, semi-perennial
344 and annual crops respectively.

346 The actual ¹⁵N recovery in the harvested biomass was on average 21.8% for perennial and
347 33.5% for the other crops (Table 6). It was significantly affected by the rotation (Table S5,
348 Appendix A). Perennial crops harvested late had a significantly lower recovery than the early
349 harvested: 13.2 ± 1.4% for Mis L vs. 34.1 ± 8.5% for Mis E. The lower ¹⁵N recovery in the
350 Alf-Fes than in the Fes-Alf rotation (24.9 vs. 39.7%) could be due to the lower yields of
351 fescue (6.8 vs. 11.7 t DM ha⁻¹ yr⁻¹) in this rotation. The ¹⁵N recovery was significantly higher
352 in N+ than in N- treatments (mean difference of 4.4%). For each crop independent of the
353 rotation, the mean ¹⁵N recovery was 29.9 and 33.6% for fescue N- and N+ respectively,
354 31.4% for sorghum N+ and 32.3 and 46.0% for triticale N- and N+ respectively.

355 The ¹⁵N recovery was compared to the apparent recovery (R_A) calculated for perennial crops
356 and sorghum (crops with an unfertilised control) (Fig. 1). The two methods gave very similar

357 results: the regression equation was $y = 0.95 R_A$ ($R^2 = 0.69$). This good agreement confirmed
358 the veracity of the low efficiency of fertiliser-N detected with the ^{15}N data.

359

360 3.4. ^{15}N recovery in dead and living crop biomass

361 A significant share of the ^{15}N fertiliser was found in residues of perennial crops (4.2% on
362 average) whereas it was almost negligible for the other crops (0.3%, Table 7). As expected,
363 the ^{15}N recovery in crop residues was higher in Mis L (6.6%) than in the other perennial
364 crops. The ^{15}N recovery in the living biomass of perennial crops (belowground organs) was
365 also important. It was higher in Mis L (17.5%) than in other perennials (9.8% on average).
366 The *Ndff* was mainly located in rhizomes for miscanthus and in roots for switchgrass. In spite
367 of their deep rooting system (Ferchaud *et al.*, 2015a), miscanthus and switchgrass allocated a
368 very small fraction of fertiliser in roots below 40 cm (0.2% on average). Finally the ^{15}N
369 recovery in the living biomass of semi-perennial and annual crops (above and belowground)
370 was low: 0.3 to 1.2%. The whole ^{15}N recovery in dead and living crop biomass ranged from
371 11.2 to 24.1% in perennial crops and 0.5 to 1.3% in the other crops. The *Ndff* represented 27-
372 32% of the total N content in dead and living biomass of perennial crops, and 1-15% for the
373 other crops.

374

375 3.5. ^{15}N recovery in soil

376 The average ^{15}N recovery of labelled fertiliser in all soil layers (L1-5, *ca.* 0-58 cm depth) was
377 31.7% for perennial crops and 23.1% for the other crops (Table 8), corresponding to 38 and
378 13 kg N ha⁻¹ yr⁻¹ respectively. Under perennial crops, the ^{15}N recovery did not differ between
379 treatments whatever the soil layer and was mainly located (85%) in the two upper layers (*ca.*
380 0-18 cm). Under semi-perennial and annual crops, the ^{15}N recovery in soil was significantly
381 affected by the rotation, the fertiliser-N rate and their interaction (Table S5, Appendix A). It

382 was higher under semi-perennial than annual crops (27.7 vs. 18.6% respectively on average in
383 L1-5) and higher in N- than in N+ (+5.6% on average) although the difference was only
384 significant for Sor-Tri. Similarly to perennial crops, 83% of the fertiliser recovered under
385 annual crops was found in the upper soil layers (*ca.* 0-19 cm). It was only 62% under semi-
386 perennial crops due to the soil ploughing event in 2011 which incorporated a part of the
387 labelled N below 19 cm.

388

389 3.6. Overall ¹⁵N recovery

390 The overall ¹⁵N recovery in the soil-plant system at the end of the ¹⁵N-labelling period, *i.e.* the
391 sum of the labelled N exported in the harvested biomass during the four or five year period
392 and stored in living crop biomass, crop residues and soil at the end of the period, is presented
393 in Fig. 2. It ranged from 51.6 ± 4.4% (Sor-Tri N+) to 82.1 ± 6.5% (Mis E N+). It was
394 significantly higher for Mis E N+ than for the other perennial crops (82.1 vs. 65.2%
395 respectively on average). Overall ¹⁵N recovery in semi-perennial and annual crops did not
396 differ between treatments and was 58.3% on average. The unrecovered ¹⁵N is attributed to
397 losses towards the groundwater and the atmosphere, *i.e.* leaching, volatilization and
398 denitrification. It represented a large part of the fertiliser: 17.9% for Mis E N+, 34.8% for
399 other perennial crops and 41.7% for semi-perennial and annual crops.

400 The *Ndff* exported at harvest represented 20% (Mis L N+) to 74% (Sor-Tri N+) of the overall
401 recovery (50% on average for all treatments). The *Ndff* stored in living crop biomass in 2011
402 or 2012 was 17% of the overall recovery for perennial crops and only 1% for the other crops.

403 The *Ndff* stored in crop residues was 6% and <1% of the overall recovery for perennials and
404 other crops respectively. Finally, the *Ndff* stored in soil in 2011 or 2012 ranged between 25%
405 (Sor-Tri N+) and 59% (Alf-Fes N-) of the overall recovery (42% on average for all
406 treatments).

407 **4. Discussion**

408 4.1. Crop production and nitrogen removal at harvest

409 *4.1.1. Crop production*

410 Perennial C4 crops were the most productive crops in this experiment. The crop yields were
411 in the range of those reviewed in the literature by Gabrielle *et al.* (2014), except for fibre
412 sorghum which had a lower production in our experiment compared to literature data
413 originating from southern Europe.

414 The interactive effect of harvest date and N fertilisation on the yield of perennial crops was
415 probably the result of the harvest date on belowground N reserves. Early harvest impedes a
416 complete N translocation from aboveground to belowground organs in autumn, reducing N
417 reserves for the succeeding year (Strullu *et al.*, 2011; Pedroso *et al.*, 2014). However, yields
418 of fertilised, early harvested treatments were higher than those of fertilised, late harvested
419 treatments of miscanthus because the aboveground biomass decreased in autumn and winter
420 due to C translocation towards rhizomes and leaf fall (Strullu *et al.*, 2011).

421

422 *4.2.2. N removal at harvest*

423 As already shown previously (Cadoux *et al.*, 2014), N exported by late harvested perennial
424 crops was particularly low because N concentration in the aboveground biomass was very low
425 at the end of winter. This is due to N translocation in autumn (Garten *et al.*, 2010; Strullu *et*
426 *al.*, 2011; Pedroso *et al.*, 2014). N exported by perennial crops harvested early was closer to
427 that observed for annual crops due to the higher N concentrations and yields than in late
428 harvest, as a result of incomplete N and C translocation. Semi-perennial crops showed the
429 highest N removal with high N concentrations, particularly for alfalfa. This result is in
430 accordance with previous studies showing high N concentrations in the harvested biomass of
431 these crops (Da Silva Perez *et al.*, 2010; Kanapeckas *et al.*, 2011). However, a large part of

432 the N removed by alfalfa probably originated from the atmosphere through symbiotic N
433 fixation (Anglade *et al.*, 2015).

434

435 4.2. N content of dead and living crop biomass

436 4.2.1. *Crop residues*

437 Although the biomass of crop residues was much higher in perennial than other crops, the
438 difference was smaller for their N content because residues of perennial crops had a higher
439 C:N ratio than other crops (85 vs. 21). The greatest amount of crop residues was found in
440 miscanthus late: 19.9 t DM ha⁻¹ and 105 kg N ha⁻¹ (average of N- and N+). About half of this
441 amount was contained in senescent leaves accumulated in a thick mulch at soil surface (8.1 t
442 DM ha⁻¹ and 50 kg N ha⁻¹) and the rest was located in stem residues. The values obtained in
443 our study for the leaf mulch were almost identical to those measured by Amougou *et al.*
444 (2012) one year earlier in the same experiment and close to the measurements of Christian *et*
445 *al.* (2006) on a 4-year-old miscanthus in late harvest (6.9 t DM ha⁻¹ and 57 kg N ha⁻¹). The
446 biomass and amount of N in switchgrass residues (10.9 t DM ha⁻¹ and 49 kg N ha⁻¹) were very
447 close to the measurements reported by Garten *et al.* (2010) for a 4-year-old switchgrass (10.7 t
448 DM ha⁻¹ and 52 kg N ha⁻¹).

449

450 4.2.2. *Living crop biomass*

451 Perennial crops were also characterized by a large amount of N stored in living belowground
452 organs. The biomass and N content of the miscanthus rhizomes observed in our experiment
453 were comparable to those reported by Himken *et al.* (1997) (16 t DM ha⁻¹ and 179-227 kg N
454 ha⁻¹) and higher than the values reported by Christian *et al.* (2006) (9.9 t DM ha⁻¹ and 140 kg
455 N ha⁻¹) also in a 4-year-old plantation with late harvests. The root biomass and N content
456 found in our experiment were intermediate between those reported by Christian *et al.* (2006)

457 and Neukirchen *et al.* (1999). For switchgrass, the amount of N stored in the rhizome was
458 higher than that reported by Garten *et al.* (2010) but the root N content over 0-60 cm was
459 similar. To our knowledge, the combined effects of N fertilisation and harvest date on the
460 belowground N content of switchgrass have not been studied in other experiments. Pedroso *et*
461 *al.* (2014) compared a two-harvest system to a single post-anthesis harvest system and found
462 that the two-harvest system increased the N removal at harvest by 51 kg N ha⁻¹ yr⁻¹ and
463 reduced belowground N stock over 0-100 cm by 36%, in accordance with our results.
464 The belowground N content of semi-perennial crops measured in 2012 over 0-60 cm (47 and
465 38 kg N ha⁻¹ for fescue and alfalfa respectively) was smaller than that of perennial crops.
466 However, these crops were re-sown in spring 2011 and the dry spring of that year caused
467 difficulties in alfalfa establishment. Indeed, the root biomass measured in 2012 was twice
468 lower than that reported by Thiébeau *et al.* (2011) at the end of the first year of growth.

469

470 4.3. ¹⁵N recovery in the soil-plant system

471 4.3.1. ¹⁵N recovery in crops

472 Perennial crops harvested late were characterized both by a low ¹⁵N recovery in the harvested
473 biomass and a high ¹⁵N recovery in living belowground organs. This is consistent with the
474 observations made for total N and apparent recovery and attributed to the important N
475 remobilisation from aboveground to belowground organs occurring in autumn. Christian *et al.*
476 (2006) also observed that the greatest part of the labelled fertiliser taken up by miscanthus
477 was located in the belowground biomass at the end of winter. Using their results, we could
478 calculate that the ¹⁵N recovery in the cumulative harvested biomass over 3 years was 28.4%.
479 This is much higher than the 13.2% observed in our experiment for miscanthus late. A similar
480 difference between the two studies was observed for the total N removed at harvest (73 vs. 38
481 kg N ha⁻¹ yr⁻¹) which suggests a greater N remobilisation in autumn in our experimental

482 conditions. Pedroso *et al.* (2014) found a similar effect of the harvest mode on a 2-year-old
483 switchgrass: ^{15}N recovery at harvest increased from 18.4 to 39.1% with a two-harvest system
484 compared to a single post-anthesis harvest system, and simultaneously the ^{15}N recovery in
485 belowground organs decreased from 27.0 to 10.4%. The ^{15}N recovery in harvested biomass in
486 the single harvest treatment was consistent with the 16.6% observed in our study (switchgrass
487 late) whereas the ^{15}N recovery in belowground organs was higher than ours (27.0 vs. 10.6%).
488 The difference is mainly due to the high ^{15}N recovery in deep roots (*ca.* 10% below 60 cm).
489 Finally, the ^{15}N recovery in the whole plant including crop residues measured in our study for
490 miscanthus late (37.3%) was smaller than the 55.8% which can be calculated using the data
491 reported by Christian *et al.* (2006) in a 4-year-old crop. For switchgrass late, there was also a
492 large gap between our result (31.3%) and the value reported by Pedroso *et al.* (2014) (47.8%)
493 that could be partly due to the difference in root sampling depth.

494 Concerning semi-perennial crops, the ^{15}N recovery in the harvested biomass ranged between
495 22.6 and 41.1%, with a significant difference between the two rotations. In the literature, the
496 ^{15}N recovery by forage crops has been mainly studied in perennial ryegrass (*Lolium perenne*).
497 Reported values of ^{15}N recovery at harvest range generally between 50 and 60% (Whitehead
498 and Dawson, 1984; Webster and Dowdell, 1985; Bristow *et al.*, 1987; Stevens and Laughlin,
499 1989). However, smaller values have been observed by Dawson and Ryden (1985) (11 to
500 48%) who showed an effect of the date of application, the ^{15}N recovery at harvest being
501 higher for spring than for summer or autumn applications. These authors also showed that ^{15}N
502 recovery in summer was much smaller in case of water stress. In our experiment, the lower
503 fescue yields observed in 2009 and 2010 and the equivalent repartition of the N applied
504 between the spring, summer and autumn cuts may have reduced the ^{15}N recovery at harvest.

505 The ^{15}N recovery at harvest of annual crops ranged between 33.6 and 39.4% but sorghum had
506 lower recovery (31.4%) than triticale (32.3-46.0%). Our results for sorghum fall in the lower

507 range of results reported for maize (*Zea mays*) which vary between 32 and 71% (Balabane
508 and Balesdent, 1992; Timmons and Baker, 1992; Reddy and Reddy, 1993; Normand *et al.*,
509 1997; Sen Tran and Giroux, 1998; Stevens *et al.*, 2005; Rimski-Korsakov *et al.*, 2012). This
510 variability is not fully understood although ¹⁵N recovery is lower for N applications at sowing
511 than at later stages and for surface than injected applications (Jokela and Randall, 1997; Seo
512 *et al.*, 2006). In our experiment, the timing of N application (at sowing) and the low growth of
513 sorghum in May and June may explain the low ¹⁵N recovery observed for this crop. The ¹⁵N
514 recovery measured for triticale in our study was also rather low (at least for N-) compared to
515 previous results reported in the literature for winter wheat (*Triticum aestivum*), ranging from
516 36 to 68% (Recous *et al.*, 1988b, 1992; Macdonald *et al.*, 1989, 1997; Powlson *et al.*, 1992;
517 Thomsen and Christensen, 2007; Giacomini *et al.*, 2010). The ¹⁵N recovery was lower for
518 applications at tillering than at stem elongation. For example, Recous *et al.* (1988b) reported
519 that ¹⁵N recovery increased from 36% for 50 kg N ha⁻¹ applied at tillering to 55% for 100 kg
520 N ha⁻¹ applied at stem elongation. This may explain the difference observed in our study for
521 triticale between N- and N+ treatments because N- received 60 kg N ha⁻¹ at tillering whereas
522 the 120 kg N ha⁻¹ for N+ were split between tillering and stem elongation.

523

524 4.3.2. ¹⁵N recovery in soil

525 Between 12.9 and 34.6% of the ¹⁵N fertiliser applied was recovered in the soil. After N
526 applications, the fertiliser inorganic N in soil is rapidly depleted due to plant uptake and
527 immobilisation of N by the soil heterotrophic microflora (Bristow *et al.*, 1987; Recous *et al.*,
528 1988a; Recous and Mchet, 1999). Microbial N is then incorporated into soil organic matter
529 and slowly mineralised in subsequent years (Glendining *et al.*, 2001; Jenkinson *et al.*, 2004;
530 Sebilo *et al.*, 2013). The ¹⁵N recovered in soil could also derive from labelled crop residues
531 returned to the soil after harvest (or crop destruction for fescue) and incorporated into the soil

532 organic matter (Macdonald *et al.*, 2002). Almost certainly, the great majority of the ¹⁵N
533 recovered in soil in our experiment was in organic rather than inorganic forms since residual
534 inorganic ¹⁵N is negligible at harvest time for optimal or sub-optimal N rates (Recous *et al.*,
535 1988b; Macdonald *et al.*, 1989; Normand *et al.*, 1997).

536 For miscanthus, we measured a higher recovery in soil (28.7% over 0-60 cm) than that
537 calculated from Christian *et al.* (2006), *i.e.* 20.6% over 0-50 cm. Pedroso *et al.* (2014)
538 reported values for switchgrass (25 to 38% over 0-300 cm) closer to our results (34.6% over
539 0-60 cm). Surprisingly, they found that a large part of the soil ¹⁵N was located in deep soil
540 layers, whereas our results and other studies showed that the great majority of the ¹⁵N
541 recovered in soil was found in the topsoil (Glendining *et al.*, 1997; Christian *et al.*, 2006).

542 The ¹⁵N recovery measured in soil for semi-perennial crops (23.3-33.9%) falls in the range of
543 the values reported in the previously cited studies for ryegrass (16 to 32%).

544 For annual crops, our results (12.9-24.4%) are consistent with those reported for wheat
545 (between 9 and 36%) and maize (between 15 and 37%) in the studies cited earlier. We
546 hypothesise that the gradient observed in our study between annual, semi-perennial and
547 perennial crops is linked to the amount and composition of crop residues. The accumulation
548 under perennial crops of undecomposed residues with a high C:N ratio probably created a
549 high microbial demand for N. On the contrary, the small amount of residues returning to the
550 soil with annual crops results in a small microbial N immobilisation, explaining the lower ¹⁵N
551 recovery in soil.

552

553 4.3.3. Overall ¹⁵N recovery

554 In our experiment, the overall ¹⁵N recovery in the crop-soil system was rather low (average
555 60%), except for miscanthus early (82%). The recovery by miscanthus late (66%) was smaller
556 than that which can be calculated using data of Christian *et al.* (2006) (77%). The values

557 reported by Pedroso *et al.* (2014) for switchgrass over three growing seasons (62-72%) were
558 quite comparable to ours (64-66%). The overall recovery that we found in semi-perennial
559 crops (average 61%) was lower than those reported in the previously cited studies for ryegrass
560 (66-95% with an average of *ca.* 80%). Finally, the lowest recovery was found in annual crops
561 (average 56%), falling in the lower range of values reported over one growing season for
562 maize (47-100%) and wheat (62-96%) averaging *ca.* 75% for both crops. In a meta-analysis
563 of published ¹⁵N field experiments on temperate climate grain crops, Gardner and Drinkwater
564 (2009) found a mean total ¹⁵N recovery of 62%, with a large variability.

565 The hypotheses provided earlier to explain the low ¹⁵N recovery in fescue, sorghum and
566 triticale can also apply to the overall recovery. Recous and Machet (1999) and Limaux *et al.*
567 (1999) showed that any increase in plant ¹⁵N uptake by winter wheat results in an increase in
568 plant and soil ¹⁵N recovery. This was confirmed by Gardner and Drinkwater (2009) who
569 showed in their meta-analysis that practices increasing ¹⁵N recovery in the crop, such as
570 improved timing or knifed-in applications, also increased the overall ¹⁵N recovery.

571 The low overall ¹⁵N recovery observed in our experiment suggests that important losses took
572 place. We believe that in a multi-annual study like ours, ¹⁵N losses are likely to be higher than
573 during a single growing season because of the remineralisation of the ¹⁵N previously
574 immobilised in soil. The losses of ¹⁵N could be due to nitrate leaching and gaseous losses, *i.e.*
575 volatilization and denitrification.

576 Losses of ¹⁵N through nitrate leaching were probably low. We evaluated nitrate leaching in
577 the site during the same period as for the ¹⁵N study (Ferchaud and Mary, 2016). The mean
578 amount of total N leached (unlabelled + labelled) below 210 cm was 2, 1 and 3 kg N ha⁻¹ yr⁻¹
579 for perennial, semi-perennial and annual crops respectively, which represented 2, 2.5 and 5%
580 of the fertiliser-N inputs. Indeed, nitrate leaching was not favoured in our context with
581 moderate winter rainfall, large soil water content and deep rooting depth (Ferchaud *et al.*,

582 2015a). However, a small part of the added ^{15}N could have moved downwards in the soil and
583 could be located between 60 and 210 cm at the time of soil sampling. This fraction was not
584 measured and therefore not included in the ^{15}N recovery.

585 Regarding gaseous losses, ammonia volatilisation could have been favoured by the type of
586 fertiliser used in our study, *i.e.* UAN containing 50% urea N. Urea is known to increase the
587 risk of ammonia volatilisation compared to other forms of N fertiliser because of the
588 temporary increase in soil pH during urea hydrolysis, particularly in a slightly alkaline soil
589 like ours. In their review, Harrison and Webb (2001) found that volatilisation represented 0-
590 4% and 6-47% of the N applied as ammonium nitrate and urea fertilisers respectively. They
591 suggested that volatilisation losses from UAN applications were intermediate between the two
592 other forms. Fox *et al.* (1996) reported ammonia volatilisation losses of 22% of the N
593 fertiliser applied as UAN on average for three years in a grain maize. Vaio *et al.* (2008)
594 measured losses ranging from 6 to 33% of the N applied as UAN to a tall fescue. Additional
595 ^{15}N losses could have also occurred through denitrification but their importance in field
596 conditions is largely unknown.

597 **5. Conclusion**

598 This study provides an original evaluation of the fate of the N fertiliser applied to different
599 perennial and annual bioenergy crops over 4-5 years. The fertiliser N recovery in the
600 harvested biomass, determined using either the ¹⁵N or the difference method, was generally
601 lower for perennial than other crops. The difference between crops was lower when
602 belowground organs of perennial crops were taken into account. Fertiliser-N immobilised in
603 soil was greater under perennial than annual crops. The overall fertiliser-N recovery (exported
604 + stored in living and dead biomass + stored in soil) tended to be greater with perennial than
605 other crops, consistently with our initial hypothesis, but crop management also affected the
606 overall recovery. Treatments ranked as follows: miscanthus harvested early > other perennial
607 crops ≥ semi-perennial and annual crops. Globally, N recovery was rather low for all crops
608 compared to achievable efficiency reported for conventional crops. It could probably be
609 increased by improvements in cropping practices (rate, timing and form of fertiliser
610 application). The effect of these practices and the partitioning of the N losses between
611 leaching, volatilisation and denitrification deserve further investigations.

612

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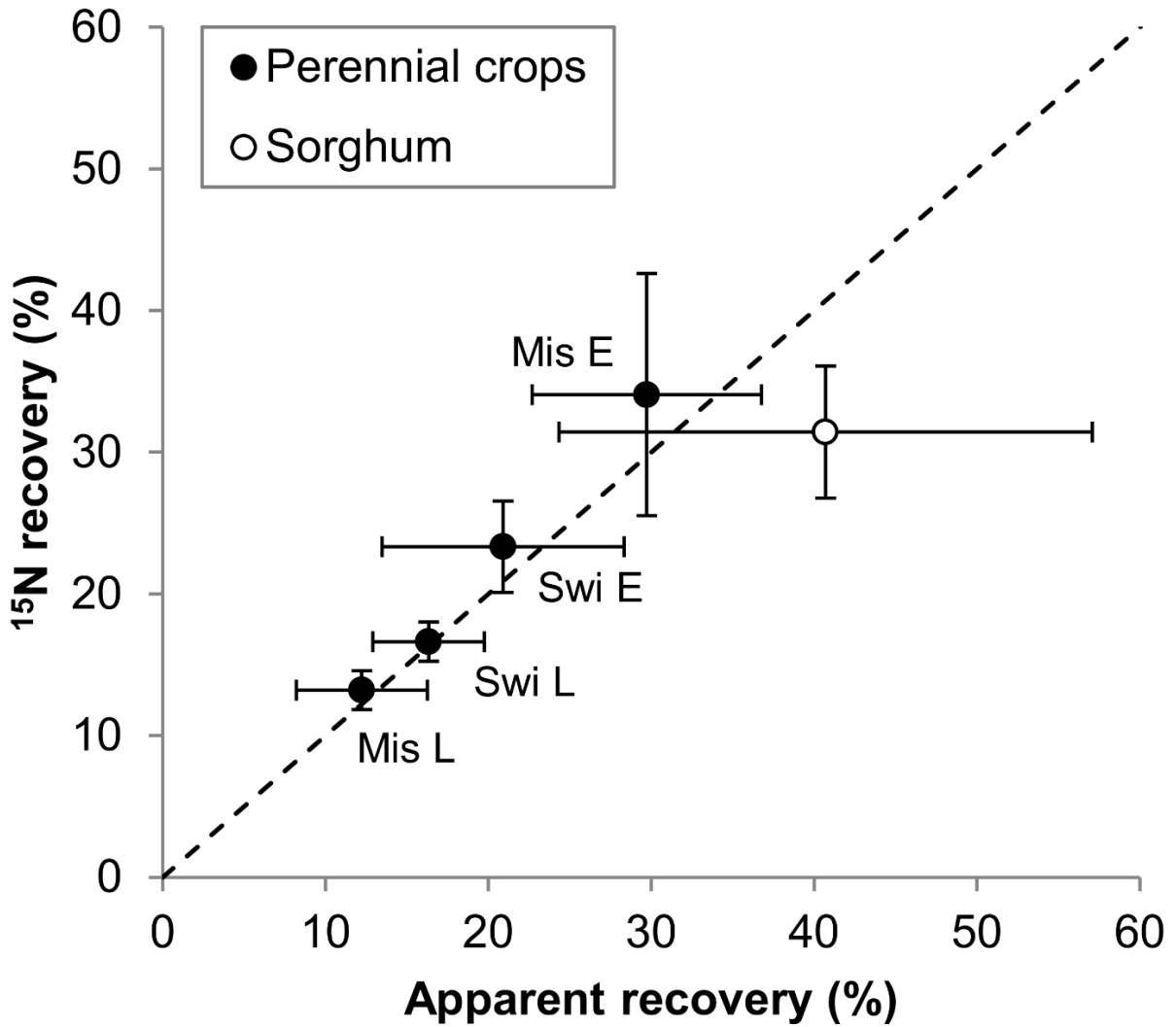
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807 **Figures**

808

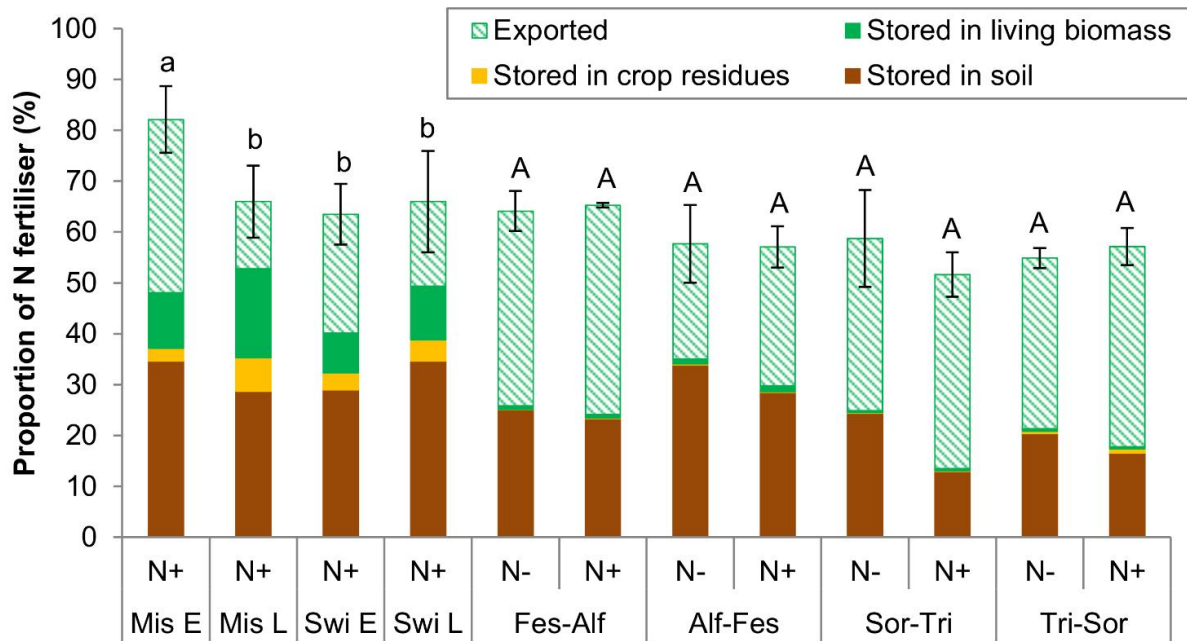
809 Fig. 1. Relationship between the ^{15}N recovery and the apparent recovery (%) in the exported
810 biomass of perennial crops (2007-2010) and sorghum N+ (2007-2011). The dashed line
811 represents the 1:1 line. Bars represent the standard deviations.



812

813

814 Fig. 2. Overall ^{15}N recovery (%) measured for perennial (2007-2010) and semi-
 815 perennial/annual crops (2007-2011). See Table 1 for abbreviations and Table 2 for fertiliser-N
 816 rates. Bars represent the standard deviations. Different letters indicate significant differences
 817 ($p < 0.05$) between treatments (lower case: perennial crops; upper case: semi-perennial and
 818 annual crops).



819

820 **Tables**

821

822 Table 1. Rotations of the B&E long term experiment (Mis = miscanthus, Swi = switchgrass,
 823 Fes = fescue, Alf = alfalfa, Sor = fiber sorghum, Tri = triticale, CC = catch crop; E = early
 824 harvest, L = late harvest, n.h. = not harvested).

Rotation	2006	2007	2008	2009	2010	2011
Mis E	Mis n.h.	Mis E	Mis E	Mis E	Mis E	Mis E
Mis L	Mis n.h.	Mis L	Mis L	Mis L	Mis L	Mis L
Swi E	Swi n.h.	Swi E	Swi E	Swi E	Swi E	Swi E
Swi L	Swi n.h.	Swi L	Swi L	Swi L	Swi L	Swi L
Fes-Alf	CC/Fes	Fes	Fes	Alf	Alf	Fes
Alf-Fes	Alf	Alf	Alf	Fes	Fes	Alf
Sor-Tri*	CC	Sor	Tri/CC	Sor	Tri/CC	Sor
Tri-Sor*	Sor	Tri/CC	Sor	Tri/CC	Sor	Tri/CC

825 *Rotations with catch crops (oats in 2006, rye in 2007, mustard in 2008, oat-vetch mixture in
 826 2009 and mustard-clover mixture from 2010 to 2011) which were sown every year in late
 827 August or early September between triticale and sorghum.

828

829 Table 2. Nitrogen fertilisation rates applied to the B&E long term experiment using ¹⁵N-
 830 labelled UAN. See Table 1 for abbreviations.

Rotation	N	N fertiliser rate (kg ha ⁻¹)						Total N applied (kg ha ⁻¹)	¹⁵ N excess atom fraction (%)
		2006	2007	2008	2009	2010	2011		
Mis E	N-	0	0	0	0	0		0	
	N+	0	120	120	120	120		480	0.395
Mis L	N-	0	0	0	0	0		0	
	N+	0	120	120	120	120		480	0.395
Swi E	N-	0	0	0	0	0		0	
	N+	0	120	120	120	120		480	0.395
Swi L	N-	0	0	0	0	0		0	
	N+	0	120	120	120	120		480	0.395
Fes-Alf	N-	0	120	80	0	0	0	200	0.395
	N+	0	240	160	0	0	0	400	0.197
Alf-Fes	N-	0	0	0	40	120	0	160	0.395
	N+	0	0	0	80	240	0	320	0.197
Sor-Tri	N-	0	0	60	0	60	0	120	0.790
	N+	0	120	120	120	120	120	600	0.395
Tri-Sor	N-	0	60	0	60	0	60	180	0.790
	N+	0	120	120	120	120	120	600	0.395

831

832

833 Table 3. Mean harvested biomass (t DM ha⁻¹ yr⁻¹) and nitrogen exported (kg ha⁻¹ yr⁻¹) from
 834 2007 to 2010 (perennial crops) or 2011 (other crops). See Table 1 for abbreviations and Table
 835 2 for fertiliser-N rates. Values in brackets are standard deviations. Different letters indicate
 836 significant differences (p<0.05) between treatments (lower case: perennial crops; upper case:
 837 other crops).

Rotation	Mean harvested biomass (t DM ha ⁻¹ yr ⁻¹)				Mean N exported (kg N ha ⁻¹ yr ⁻¹)			
	N-		N+		N-		N+	
Mis E	24.2 (3.0)	b	26.6 (2.6)	a	100 (10)	b	135 (18)	a
Mis L	19.0 (2.2)	cd	18.7 (1.6)	cdef	38 (10)	e	52 (5)	cd
Swi E	15.6 (0.6)	dfh	18.1 (1.4)	ceg	70 (4)	cd	95 (5)	b
Swi L	14.8 (0.8)	gh	15.1 (1.3)	efgh	51 (6)	de	71 (2)	c
Fes-Alf	9.8 (0.2)	BC	10.4 (0.3)	B	180 (6)	C	190 (12)	BC
Alf-Fes	8.9 (0.4)	C	9.6 (0.2)	BC	217 (12)	AB	228 (15)	A
Sor-Tri	9.8 (0.3)	BC	11.9 (0.5)	A	71 (6)	E	122 (20)	D
Tri-Sor	9.3 (0.6)	BC	12.6 (1.3)	A	66 (7)	E	114 (13)	D

838

840 Table 4. Dead (crop residues) and living crop biomass (t DM ha⁻¹) measured in March 2011 for perennial crops and March 2012 for semi-
 841 perennial/annual crops. See Table 1 for abbreviations and Table 2 for fertiliser-N rates. Values in brackets are standard deviations. Different
 842 letters indicate significant differences (p<0.05) between treatments (lower case: perennial crops; upper case: other crops).

	Mis E		Mis L		Swi E		Swi L	
	N-	N+	N-	N+	N-	N+	N-	N+
Aboveground crop residues	5.3 (1.3) b	6.5 (1.3) b	16.1 (1.2) a	13.3 (1.2) a	5.5 (2.4) b	6.9 (2.3) b	6.6 (0.9) b	6.8 (0.6) b
Belowground crop residues	6.6 (4.8)	4.1 (2.3)	5.7 (2.2)	4.5 (1.5)	3.4 (0.1)	4.0 (1.3)	5.8 (3.9)	4.6 (2.4)
<i>a) Total crop residues</i>	<i>11.9 (3.5) b</i>	<i>10.6 (2.6) b</i>	<i>21.9 (1.6) a</i>	<i>17.8 (2.7) a</i>	<i>8.9 (2.4) b</i>	<i>10.8 (1.2) b</i>	<i>12.5 (4.7) b</i>	<i>11.4 (2.5) b</i>
Rhizome	17.0 (6.4) a	14.2 (0.9) a	19.7 (1.3) a	16.8 (2.9) a	4.6 (2.3) b	3.0 (2.7) b	5.9 (1.7) b	3.4 (3.6) b
Roots (0-20 cm)	4.7 (0.4) b	3.3 (0.6) b	4.1 (1.6) b	3.6 (1.7) b	7.5 (1.5) a	7.6 (2.9) a	8.9 (3.6) a	6.7 (3.1) a
Roots (20-40 cm)	0.9 (0.2) b	0.4 (0.1) b	0.7 (0.2) b	0.5 (0.1) b	2.8 (0.5) a	2.7 (0.6) a	2.4 (0.9) a	1.9 (0.3) a
Roots (40-60 cm)	0.3 (0.1) b	0.2 (0.1) b	0.2 (0.2) b	0.3 (0.1) b	1.1 (0.2) a	0.8 (0.1) a	1.0 (0.3) a	0.9 (0.3) a
<i>b) Total living crop biomass</i>	<i>22.9 (6.2) ab</i>	<i>18.1 (0.5) ab</i>	<i>24.7 (3.0) a</i>	<i>21.2 (3.3) a</i>	<i>16.0 (3.2) b</i>	<i>14.2 (6.1) b</i>	<i>18.2 (6.4) b</i>	<i>12.9 (6.3) b</i>
<i>Total (a+b)</i>	<i>34.8 (9.7) b</i>	<i>28.8 (2.1) b</i>	<i>46.6 (3.0) a</i>	<i>39.0 (4.0) a</i>	<i>24.9 (5.7) b</i>	<i>25.0 (5.5) b</i>	<i>30.7 (11.1) b</i>	<i>24.3 (8.9) b</i>

	Fes-Alf		Alf-Fes		Sor-Tri		Tri-Sor	
	N-	N+	N-	N+	N-	N+	N-	N+
<i>a) Total crop residues</i>	<i>3.5 (1.2) A</i>	<i>4.0 (1.7) A</i>	<i>0.8 (0.2) B</i>	<i>0.6 (0.2) B</i>	<i>1.6 (0.3) B</i>	<i>1.6 (0.6) B</i>	<i>3.3 (0.8) A</i>	<i>3.7 (0.7) A</i>
Aboveground living biomass	3.6 (0.5) A	3.9 (0.7) A	1.1 (0.6) B	1.1 (0.6) B				
Belowground living biomass	2.4 (0.7)	2.8 (0.3)	1.5 (1.2)	1.3 (0.6)				
<i>b) Total living crop biomass</i>	<i>6.0 (1.2) A</i>	<i>6.7 (0.9) A</i>	<i>2.6 (1.7) B</i>	<i>2.4 (1.2) B</i>	<i>0.4 (0.0) C</i>	<i>0.4 (0.1) C</i>	<i>1.2 (0.5) BC</i>	<i>1.2 (0.2) BC</i>
<i>Total (a+b)</i>	<i>9.5 (2.3) A</i>	<i>10.6 (0.9) A</i>	<i>3.4 (1.7) BC</i>	<i>3.1 (1.4) BC</i>	<i>2.0 (0.3) C</i>	<i>2.0 (0.5) C</i>	<i>4.5 (1.1) B</i>	<i>5.0 (0.9) B</i>

843 Table 5. N content in dead (crop residues) and living crop biomass (kg ha⁻¹) measured in March 2011 for perennial crops and March 2012 for
844 semi-perennial/annual crops. See Table 1 for abbreviations and Table 2 for fertiliser-N rates. Values in brackets are standard deviations. Different
845 letters indicate significant differences (p<0.05) between treatments (lower case: perennial crops; upper case: other crops). The signs - and +
846 indicate a significant effect of N fertilisation (without interaction with rotations).

	Mis E		Mis L		Swi E		Swi L	
	N-	N+	N-	N+	N-	N+	N-	N+
Aboveground crop residues	14 (4) e	27 (6) cde	71 (9) b	85 (3) a	20 (9) de	34 (13) c	23 (3) cde	35 (2) bc
Belowground crop residues	20 (13)	18 (7)	24 (9)	30 (11)	13 (1)	19 (4)	23 (16)	28 (18)
<i>a) Total crop residues</i>	34 (10) b-	44 (1) b+	95 (2) a-	115 (13) a+	33 (10) b-	54 (10) b+	46 (18) b-	63 (18) b+
Rhizome	63 (24) b-	147 (39) b+	167 (8) a-	247 (66) a+	30 (18) c-	34 (27) c+	52 (17) c-	51 (51) c+
Roots (0-20 cm)	35 (2) b	38 (2) b	40 (13) ab	56 (24) ab	34 (5) b	58 (11) B	52 (19) a	79 (32) a
Roots (20-40 cm)	6 (1) bc	4 (0) c	7 (2) bc	7 (2) bc	12 (1) b	22 (5) a	13 (4) b	21 (4) a
Roots (40-60 cm)	2 (1) c	1 (1) c	2 (1) c	3 (1) bc	4 (1) bc	5 (1) ab	4 (1) bc	7 (2) a
<i>b) Total living crop biomass</i>	106 (27) b-	190 (37) b+	216 (11) a-	313 (71) a+	79 (19) b-	119 (33) b+	120 (39) b-	159 (77) b+
<i>Total (a+b)</i>	140 (36) b-	234 (37) b+	311 (11) a-	428 (73) a+	113 (27) b-	173 (27) b+	166 (57) b-	221 (95) b+
	Fes-Alf		Alf-Fes		Sor-Tri		Tri-Sor	
	N-	N+	N-	N+	N-	N+	N-	N+
<i>a) Total crop residues</i>	21 (7) A	30 (14) A	7 (2) B	5 (1) B	5 (0) B	6 (2) B	11 (2) A	23 (10) A
Aboveground living biomass	82 (4) A	90 (31) A	18 (9) B	21 (12) B				
Belowground living biomass	44 (13)	50 (8)	43 (37)	34 (20)				
<i>b) Total living crop biomass</i>	126 (16) A	140 (39) A	61 (46) B	55 (32) B	15 (0) B	15 (4) B	26 (7) B	27 (7) B
<i>Total (a+b)</i>	147 (24) A	170 (25) A	68 (48) B	60 (33) B	20 (0) C	22 (3) C	37 (8) BC	50 (16) BC

847

848 Table 6. *Ndff* in the exported biomass (kg N ha⁻¹) during each year, N derived from fertiliser
849 and other sources and ¹⁵N recovery (%) in the exported biomass over the whole period. See
850 Table 1 for abbreviations and Table 2 for fertiliser-N rates. Values in brackets are standard
851 deviations. Different letters indicate significant differences (p<0.05) between treatments
852 (lower case: perennial crops; upper case: other crops). The signs - and + indicate a significant
853 mean effect of N fertilisation (without interaction with rotations).

Rotation	N	<i>Ndff</i> in the exported biomass (kg N ha ⁻¹)					Total N exported (kg N ha ⁻¹ yr ⁻¹)		¹⁵ N recovery in exported biomass (%)
		2007	2008	2009	2010	2011	From fertiliser	From other sources	
Mis E	N+	30 (8)	28 (8)	45 (12)	60 (19)		41 (10)	94 (11)	34.1 (8.5) a
Mis L	N+	8 (1)	11 (3)	17 (0)	28 (3)		16 (2)	36 (4)	13.2 (1.4) c
Swi E	N+	37 (6)	19 (8)	24 (2)	32 (12)		28 (4)	67 (1)	23.3 (3.2) b
Swi L	N+	23 (1)	23 (4)	14 (3)	20 (0)		20 (2)	51 (3)	16.6 (1.4) c
Fes-Alf	N-	48 (1)	24 (6)	0 (0)	2 (0)	3 (1)	15 (1)	165 (5)	38.3 (2.9) A-
	N+	96 (2)	61 (7)	1 (0)	2 (1)	5 (1)	33 (2)	157 (11)	41.1 (1.9) A+
Alf-Fes	N-	0 (0)	0 (0)	8 (1)	25 (8)	3 (3)	7 (2)	210 (13)	22.6 (6.1) B-
	N+	0 (0)	0 (0)	20 (3)	61 (9)	6 (6)	17 (2)	211 (15)	27.3 (2.5) B+
Sor-Tri	N-	0 (0)	17 (6)	1 (1)	21 (4)	1 (1)	8 (2)	63 (6)	33.8 (7.0) A-
	N+	37 (12)	46 (7)	49 (0)	57 (5)	39 (9)	46 (7)	77 (14)	38.2 (5.5) A+
Tri-Sor	N-	23 (2)	1 (0)	18 (6)	1 (0)	17 (1)	12 (1)	54 (6)	33.6 (2.6) A-
	N+	59 (5)	34 (4)	54 (1)	30 (9)	60 (7)	47 (4)	67 (10)	39.4 (3.6) A+

854

855 Table 7. ¹⁵N recovery (%) measured in dead (crop residues) and living crop biomass in March 2011 for perennial crops and March 2012 for semi-
856 perennial/annual crops. See Table 1 for abbreviations and Table 2 for fertiliser-N rates. Values in brackets are standard deviations. Different
857 letters indicate significant differences (p<0.05) between treatments (lower case: perennial crops; upper case: other crops).

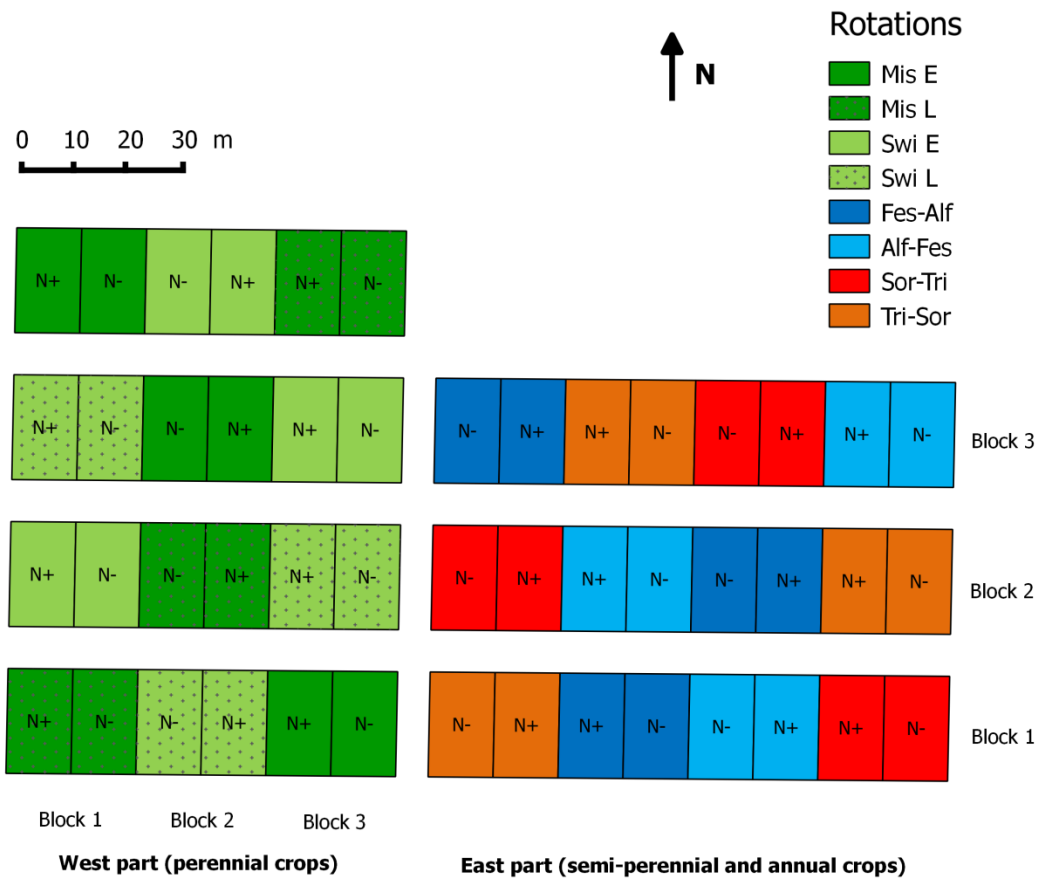
	Mis E		Mis L		Swi E		Swi L	
	N-	N+	N-	N+	N-	N+	N-	N+
Aboveground crop residues		1.5 (0.4) b		4.7 (0.2) a		2.2 (1.0) b		2.3 (0.1) b
Belowground crop residues		1.0 (0.4)		1.9 (0.7)		1.2 (0.2)		1.8 (1.4)
<i>a) Total crop residues</i>		2.6 (0.1) c		6.6 (0.6) a		3.4 (1.1) bc		4.1 (1.5) b
Rhizome		8.4 (2.8) ab		13.8 (2.8) a		2.3 (1.9) c		3.7 (3.7) bc
Roots (0-20 cm)		2.3 (0.1) b		3.2 (1.0) ab		4.0 (0.8) ab		5.2 (2.1) a
Roots (20-40 cm)		0.2 (0.1) b		0.3 (0.0) b		1.3 (0.3) a		1.2 (0.2) a
Roots (40-60 cm)		0.0 (0.0) b		0.1 (0.1) b		0.3 (0.1) a		0.5 (0.2) a
<i>b) Total living crop biomass</i>		10.9 (2.8) b		17.5 (2.5) a		7.9 (2.2) b		10.6 (5.4) b
<i>Total (a+b)</i>		13.4 (2.7) b		24.1 (2.4) a		11.2 (1.8) b		14.7 (6.9) b
	Fes-Alf		Alf-Fes		Sor-Tri		Tri-Sor	
	N-	N+	N-	N+	N-	N+	N-	N+
<i>a) Total crop residues</i>	0.1 (0.0) B	0.1 (0.1) B	0.2 (0.1) B	0.1 (0.0) B	0.1 (0.0) B	0.1 (0.0) B	0.5 (0.0) A	0.8 (0.5) A
Aboveground living biomass	0.6 (0.1)	0.5 (0.2)	0.4 (0.1)	0.4 (0.2)				
Belowground living biomass	0.2 (0.0)	0.2 (0.0)	0.6 (0.4)	0.8 (0.5)				
<i>b) Total living crop biomass</i>	0.7 (0.1) AB	0.7 (0.2) AB	1.0 (0.5) A	1.2 (0.8) A	0.3 (0.1) B	0.4 (0.2) B	0.5 (0.2) B	0.4 (0.1) B
<i>Total (a+b)</i>	0.8 (0.1)	0.8 (0.2)	1.2 (0.6)	1.3 (0.8)	0.5 (0.1)	0.5 (0.1)	1.0 (0.3)	1.2 (0.6)

858

859 Table 8. ^{15}N recovery (%) measured in soil layers in 2011 for perennial crops and in 2012 for semi-perennial/annual crops. See Table 1 for
 860 abbreviations and Table 2 for fertiliser-N rates. Layers L1, L2, L3, L4 and L5 correspond to *ca.* 0-5, 5-18, 18-32, 32-38 and 38-58 cm
 861 respectively. Values in brackets are standard deviations. Different letters indicate significant differences ($p < 0.05$) between treatments (lower
 862 case: perennial crops; upper case: semi-perennial and annual crops).

Layer	Soil mass (t ha ⁻¹)	Mis E	Mis L	Swi E	Swi L	Fes-Alf		Alf-Fes		Sor-Tri		Tri-Sor	
		N+	N+	N+	N+	N-	N+	N-	N+	N-	N+	N-	N+
L1	667	20.4 (6.2)	16.5 (2.2)	15.4 (3.5)	17.3 (2.5)	2.0 (0.3)	1.9 (0.2)	5.0 (3.4)	3.0 (0.5)	6.7 (1.8)	4.5 (0.6)	7.8 (2.2)	5.3 (1.2)
L2	2000	9.6 (1.6)	8.5 (2.6)	9.0 (2.2)	11.7 (2.3)	14.6 (1.9)	12.4 (2.2)	15.3 (4.9)	14.0 (5.3)	12.8 (2.7)	6.0 (2.0)	9.7 (0.6)	9.0 (0.8)
L3	2002	2.7 (0.2)	2.6 (1.1)	2.7 (0.5)	3.7 (0.6)	7.6 (1.6)	8.1 (2.7)	11.3 (7.9)	9.0 (3.3)	3.3 (0.3)	1.7 (0.4)	2.0 (0.6)	1.6 (0.3)
L4	884	0.3 (0.1)	0.3 (0.1)	0.5 (0.1)	0.7 (0.3)	0.2 (0.2)	0.3 (0.0)	0.7 (0.2)	0.8 (0.3)	0.5 (0.1)	0.2 (0.1)	0.3 (0.1)	0.2 (0.1)
L5	3137	1.6 (0.8)	0.8 (0.2)	1.3 (0.7)	1.2 (0.4)	0.7 (0.3)	0.6 (0.2)	1.6 (0.6)	1.8 (0.9)	1.2 (0.1)	0.6 (0.2)	0.6 (0.1)	0.5 (0.2)
L1-5	8690	34.6 (7.0)	28.7 (6.0)	28.9 (5.8)	34.6 (3.0)	25.1 (1.8)	23.3 (1.8)	33.9 (2.0)	28.5 (3.8)	24.4 (4.6)	12.9 (2.5)	20.4 (2.1)	16.6 (1.3)
		<i>a</i>	<i>a</i>	<i>a</i>	<i>a</i>	<i>BC</i>	<i>BC</i>	<i>A</i>	<i>AB</i>	<i>BC</i>	<i>E</i>	<i>CD</i>	<i>DE</i>

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866

867 Fig. S1. Map representing the experimental design of the B&E long-term experiment (see

868 Table 1 for abbreviations and Table 2 for fertiliser-N rates). All plots are 12×30 m (360 m^2)

869 and the whole field is 2.7 ha

870

871 Table S1. Physical and chemical soil characteristics measured in 2006 for the two parts of the
 872 field trial. Values in brackets are standard deviations between the 24 plots in each part

Part of the field trial	Soil layer (cm)	Clay <2 µm (g kg ⁻¹)	Fine silt 2-20 µm (g kg ⁻¹)	Coarse silt 20-50 µm (g kg ⁻¹)	Fine sand 50-200 µm (g kg ⁻¹)	Coarse sand 200-2000 µm (g kg ⁻¹)	CaCO ₃ (g kg ⁻¹)	pH water
West	0-30	180 (27)	319 (14)	447 (22)	40 (8)	12 (2)	2 (1)	7.8 (0.2)
	30-60	233 (20)	311 (19)	409 (14)	39 (12)	6 (2)	2 (2)	7.8 (0.2)
East	0-30	148 (19)	331 (14)	471 (14)	34 (10)	14 (4)	3 (2)	7.9 (0.2)
	30-60	187 (35)	340 (61)	430 (60)	36 (15)	7 (3)	1 (1)	8.0 (0.2)

873

874

875 Table S2. Statistical analysis of the effects of rotation, nitrogen fertilisation and their
 876 interaction on mean harvested biomass and mean N exported for perennial crops and
 877 annual/semi-perennial crops. Asterisks indicate probability levels: * p<0.05; ** p<0.01; ***
 878 p<0.001; NS = not significant

Factor or interaction		Biomass		Nitrogen content	
		Perennials	Other crops	Perennials	Other crops
Harvested biomass					
Rotation	1	***	***	***	***
Nitrogen	2	***	***	***	***
	1 x 2	**	**	*	**
Total crop residues					
Rotation	1	***	**	***	**
Nitrogen	2	NS	NS	*	NS
	1 x 2	NS	NS	NS	NS
Total living biomass					
Rotation	1	*	***	***	**
Nitrogen	2	NS	NS	*	NS
	1 x 2	NS	NS	NS	NS

879

880

881 Table S3. Biomass harvested (t DM ha⁻¹) from 2006 to 2011. See Table 1 for abbreviations
 882 and Table 2 for fertiliser-N rates. Values in brackets are standard deviations.

Rotation	N	2006	2007	2008	2009	2010	2011
Mis E	N-	0 (0)	23.0 (4.9)	23.6 (2.4)	24.0 (3.3)	26.1 (1.5)	
	N+	0 (0)	21.7 (4.5)	25.2 (2.6)	28.8 (3.3)	30.6 (2.0)	
Mis L	N-	0 (0)	14.3 (4.5)	18.5 (1.7)	20.9 (1.9)	22.2 (2.2)	
	N+	0 (0)	13.9 (2.3)	18.7 (2.4)	19.6 (1.4)	22.4 (1.7)	
Swi E	N-	0 (0)	19.6 (2.4)	18.9 (0.2)	14.9 (1.2)	9.2 (0.7)	
	N+	0 (0)	21.5 (2.9)	16.7 (5.9)	19.2 (1.2)	15.2 (4.2)	
Swi L	N-	0 (0)	15.9 (1.0)	16.7 (0.8)	13.8 (0.6)	12.6 (1.2)	
	N+	0 (0)	15.2 (1.4)	15.9 (2.0)	15.2 (2.3)	14.0 (0.7)	
Fes-Alf	N-	0 (0)	16.1 (0.6)	7.7 (0.7)	3.5 (0.4)	12.4 (1.0)	9.2 (1.9)
	N+	0 (0)	17.3 (0.4)	12.1 (0.6)	2.7 (0.0)	11.8 (0.8)	8.1 (0.9)
Alf-Fes	N-	8.0 (0.6)	14.6 (0.9)	15.8 (0.2)	5.8 (1.1)	6.3 (2.0)	2.0 (1.8)
	N+	7.5 (1.2)	14.6 (0.4)	16.0 (0.3)	6.4 (0.9)	8.6 (0.6)	2.4 (2.1)
Sor-Tri	N-	0 (0)	14.0 (2.3)	9.7 (0.6)	12.3 (1.8)	9.3 (0.3)	3.6 (0.6)
	N+	0 (0)	12.8 (1.8)	14.8 (1.5)	14.8 (2.1)	12.7 (0.8)	4.5 (0.6)
Tri-Sor	N-	15.2 (0.6)	11.5 (1.2)	11.1 (2.2)	8.3 (1.1)	7.8 (1.2)	7.7 (0.4)
	N+	15.2 (0.5)	13.5 (0.2)	14.2 (1.6)	12.3 (0.6)	13.4 (2.6)	9.9 (2.0)

883

884

885 Table S4. Nitrogen exported (kg ha^{-1}) from 2006 to 2011. See Table 1 for abbreviations and
 886 Table 2 for fertiliser-N rates. Values in brackets are standard deviations.

Rotation	N	2006	2007	2008	2009	2010	2011
Mis E	N-	0 (0)	108 (15)	90 (10)	75 (19)	126 (11)	
	N+	0 (0)	99 (25)	108 (18)	137 (25)	197 (12)	
Mis L	N-	0 (0)	22 (8)	33 (12)	33 (9)	61 (17)	
	N+	0 (0)	24 (5)	44 (9)	55 (2)	86 (14)	
Swi E	N-	0 (0)	105 (14)	89 (8)	50 (9)	37 (1)	
	N+	0 (0)	149 (20)	83 (18)	71 (5)	78 (21)	
Swi L	N-	0 (0)	80 (12)	60 (8)	29 (2)	37 (5)	
	N+	0 (0)	95 (10)	86 (8)	48 (7)	54 (3)	
Fes-Alf	N-	0 (0)	218 (15)	66 (9)	86 (8)	320 (33)	210 (47)
	N+	0 (0)	275 (35)	118 (10)	68 (5)	315 (25)	175 (25)
Alf-Fes	N-	212 (13)	408 (15)	417 (32)	86 (6)	99 (29)	76 (67)
	N+	198 (31)	409 (7)	441 (6)	80 (15)	136 (16)	75 (65)
Sor-Tri	N-	0 (0)	134 (23)	53 (7)	67 (6)	54 (3)	49 (10)
	N+	0 (0)	149 (52)	108 (26)	140 (18)	104 (11)	110 (26)
Tri-Sor	N-	170 (7)	92 (10)	89 (21)	56 (11)	37 (8)	55 (7)
	N+	170 (6)	134 (7)	124 (15)	107 (7)	97 (34)	109 (13)

887

888

889 Table S5. Statistical analysis of the effects of rotation, nitrogen fertilisation and their
 890 interaction on ¹⁵N recovery for perennial crops and annual/semi-perennial crops. Asterisks
 891 indicate probability levels: * p<0.05; ** p<0.01; *** p<0.001; NS = not significant

		Perennials	Other crops
Harvested biomass			
Rotation	1	**	*
Nitrogen	2		*
	1 x 2		NS
Total crop residues			
Rotation	1	**	***
Nitrogen	2		NS
	1 x 2		NS
Total living biomass			
Rotation	1	*	*
Nitrogen	2		NS
	1 x 2		NS
Soil			
Rotation	1	NS	***
Nitrogen	2		***
	1 x 2		*
Overall recovery			
Rotation	1	**	NS
Nitrogen	2		NS
	1 x 2		NS

892

893