

Fast assimilate turnover revealed by *in situ* ¹³CO₂ pulse labelling in Subarctic tundra

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1 **Abstract**

2 Climatic changes in Arctic regions are likely to have significant impacts on vegetation
3 composition and physiological responses of different plant types, with implications
4 for the regional carbon (C) cycle. Here we explore differences in allocation and
5 turnover of assimilated C in two Subarctic tundra communities. We used an *in situ* ¹³C
6 pulse at mid-summer in Swedish Lapland to investigate C allocation and turnover in
7 four contrasting tundra plant communities. We found a high rate of turnover of
8 assimilated C in leaf tissues of *Betula nana* and graminoid vegetation at the height of
9 the growing season, with a mean residence time of pulse-derived ¹³C of 1.1 and 0.7
10 days, respectively. One week after the pulse, c. 20 and 15%, respectively, of
11 assimilated label-C remained in leaf biomass, representing most likely allocation to
12 structural biomass. For the perennial leaf tissue of the graminoid communities, a
13 remainder of approximately 5% of the pulse-derived C was still traceable after 1 year,
14 whereas none was detectable in *Betula* foliage. The results indicate a relatively fast C
15 turnover and small belowground allocation during the active growing season of recent
16 assimilates in graminoid communities, with comparatively slower turnover and
17 greater investment in belowground allocation by *Betula nana* vegetation.

18

19 **Keywords:**

20 Carbon cycle; GPP partitioning; Stable Isotopes, Tundra biome;

21

22 **Introduction**

23 Tundra vegetation is a heterogeneous cover of different vegetation types occurring in
24 patches throughout the landscape, reflecting a harsh climate and well-adapted, but
25 species-poor, communities. Changes in the frequency and extents of different
26 vegetation types are likely to result from altered climatic conditions (Street et al.
27 2007). There is already abundant evidence for an increase in shrub expansion in
28 Arctic tundra over the past 50 years (Tape et al. 2006), with direct consequences on
29 the albedo (Chapin et al. 2005), hydrology and cycling of C and nutrients (Post et al.
30 2009; Wookey et al. 2009). Process models indicate that this “greening” of the tundra
31 (Wookey et al. 2009) has led to an increase in net primary productivity (NPP), which
32 is predicted to persist into the near future (Sitch et al. 2007). However, these process
33 models are based predominantly on leaf area relationships, and therefore lack the
34 necessary level of detail for the partitioning of assimilated C (i.e. gross primary
35 productivity, GPP) to accurately predict changes in the net C balance and cycling of
36 nutrients in tundra ecosystems (Euskirchen et al. 2009; Hudson and Henry 2009;
37 McGuire et al. 2009; Roberts et al. 2009; Street et al. 2007). NPP modelling
38 approaches are necessarily based on generalisations of GPP to NPP ratios, with a
39 range of assumptions regarding the apportioning of assimilated C to either respiratory
40 loss or biomass gain, and ultimately turnover rates of C in ecosystems (Gifford 2003;
41 Trumbore 2006; Waring et al. 1998). Moreover, ecophysiological responses used in
42 ecosystem models often lack the necessary detail and have treated vegetation
43 characteristics uniformly (Shaver et al. 2007; Williams et al. 2001), ignoring unique
44 responses of different plant functional types (PFTs) and their individual C allocation
45 patterns. Likely changes in vegetation structure and hence shifts in nutrient input and

46 cycling (van Wijk et al. 2004) following climatic change require new modelling
47 approaches that can capture these dynamic variations. Consequently, there is a need to
48 obtain direct measurements of C allocation and turnover in different tundra vegetation
49 communities, in order to parameterise ecosystem models considering separate PFTs
50 (Williams et al. 2006). So far, little is known about the PFT-specific C partitioning
51 and turnover in tundra vegetation (Douma et al. 2007; Shaver et al. 2006; Street et al.
52 2011; Woodin et al. 2009), but an understanding of the partitioning of GPP into either
53 respiration or biomass is crucial, as this ultimately determines the long-term fate of
54 assimilated C and likely differs between PFTs.

55 Pulse labelling experiments using stable isotope tracers have become a standard tool
56 in ecophysiology to investigate the fate of assimilated C in the plant-soil-atmosphere
57 continuum. They are frequently used to unravel processes of assimilation, allocation
58 and respiration (Bowling et al. 2008; Dawson et al. 2002). We used the stable carbon
59 isotope ^{13}C to quantify, *in situ*, the amount of C incorporated into plant biomass and
60 the rate of respiration of labelled C in relation to the total amount of label uptake
61 during photosynthesis. This approach has previously been successful in forest and
62 shrubland ecosystems (Carbone and Trumbore 2007; Carbone et al. 2007; Höglberg et
63 al. 2008). Whilst C partitioning in tundra vegetation has been addressed in a
64 radiocarbon (^{14}C) pulse labelling experiment (Olsrud and Christensen 2004), there are
65 no published results of GPP partitioning and C turnover for specific plant functional
66 types representative of vascular vegetation. The aims of our study were to apply an *in*
67 *situ* stable isotope ($^{13}\text{CO}_2$) pulse in representative tundra vegetation to (1) trace
68 assimilated C into different biomass pools, (2) estimate the turnover of assimilated C
69 and (3) assess the long-term fate of these assimilates within different PFTs. The
70 presented work concentrates on vascular plant communities but also includes

71 representative lichen dominated plots, whilst an accompanying experiment conducted
72 at the same site and over the same period as this study investigated GPP:NPP
73 partitioning and C turnover in bryophyte communities (Street et al. 2011).

74 **Materials and Methods**

75 *Site description*

76 The study site was located in northern Sweden near Abisko at 68°18'N, 18°51'E at
77 about 700 m asl (Fig. 1). The mean annual temperature in the Abisko Valley is -1 °C
78 (recorded at the Abisko Research Station situated 6 km north of the research site at
79 400 m asl), with mean July air temperatures of +11 °C
80 (<http://www.linnea.com/~ans/ans.htm>). Three replicated plots (0.55 x 0.55 m) were
81 established for each of four contrasting vegetation types, representing typical tundra
82 heath communities dominated by key species or plant types (Table 1): (1) dwarf birch
83 (*Betula nana* L.), (2) crowberry (*Empetrum nigrum* L.), (3) graminoid communities
84 (dominated by sedges), and (4) exposed ridges (dominated by lichen). All 12 plots
85 were located within an area of 15 x 20 m. Soils comprised organic horizons developed
86 on glacial till deposits. Organic layer depth varied between 5 and 30 cm for the three
87 vascular plant communities, whilst there were only small “pockets” of soil material in
88 lichen plots. The patchwork of vegetation communities over relatively short distances
89 relates mostly to the micro-topography between small rocky outcrops and resulting
90 variability in soil drainage. While graminoid communities tended to occur in areas of
91 lower drainage, *E. nigrum* and *B. nana* patches were generally situated in more mesic
92 locations. Areas dominated by mosses also co-occurred where soil moisture was
93 highest; these were not part of this study (but see Street et al. 2011).

94 *¹³CO₂ labelling*

95 Isotope pulse labelling was carried out on 4th July 2007 between 13:20 and 16:00 hrs.
96 For pulse labelling, we placed custom-made Perspex covers (0.55 x 0.55 m, 0.2 m
97 high; York Plastics, York, UK) over the 12 individual vegetation patches without
98 inserting a soil frame, thus preventing potential damage to the branch and root
99 systems, a particular concern in these dwarf shrub communities (Fig. 2a). Larger gaps
100 between the cover edge and the vegetation or ground were filled using clear plastic
101 film. Remaining gaps were small and not significant for an incursion of ambient air
102 given the over-pressured chamber conditions during the labelling gas flow. The pulse
103 gas (artificial air with 371 – 375 µl l⁻¹ CO₂ with 98% ¹³C atom enrichment; Spectra
104 Gases Ltd., Littleport, UK) flushed the Perspex covers at a flow rate of 5 l min⁻¹,
105 resulting in a mean residence time of pulse gas within the covers of 12 minutes.
106 Weather conditions during the pulse period were sunny with air temperatures
107 averaging 20.2 °C, i.e. relatively warm but not an unusual July air temperature for
108 daytime maxima in the region (Fig. 3). Tests under comparable meteorological
109 conditions and with identical gas flow rates showed a mean increase of 2.7 °C inside
110 the labelling hood compared to ambient temperatures, so that temperature conditions
111 inside the sampling hoods are not likely to have exceeded 25 °C.

112 *Pulse chase sampling*

113 Leaf and lichen materials were sampled before the pulse, and then at 0.25, 0.75, 1, 2,
114 3, 5 and 7 days following the pulse. At each sampling occasion, approximate
115 equivalents of 1 g dry mass of foliage of the dominant plant species (or lichen on
116 exposed ridges) were collected, ensuring an even representation of plants within the
117 area covered by the cover. Senesced foliage was removed, and samples were

118 transferred into paper envelopes at the field site and subsequently transferred to the
119 laboratory and oven-dried at 60 °C. The remoteness of the site meant that oven drying
120 of samples occurred between 3 and 12 hours of sampling in the field for individual
121 samples. As we can not exclude the possibility that some of the labelled assimilates
122 were lost as respiration in this period, we treat our estimates of C turnover as low
123 estimates of actual turnover. Soil samples were taken initially before the pulse, and
124 again at 1 and 7 days post-pulse, to a depth of 5 cm using a 5 cm diameter corer.
125 Sampling to this depth resulted in obtaining mainly organic soil material, with only
126 small amounts of mineral content in individual samples. All samples were stored in
127 plastic bags in soil pits at below 5 °C before transportation to the laboratory (within
128 less than 5 hours), where samples were frozen.

129 *¹³CO₂ monitoring*

130 Immediately after the end of the pulse, opaque PVC chambers (20 cm diameter, 10
131 cm high; Fig. 2b) were placed on the vegetation at the centre of the 0.55 x 0.55 m
132 pulse areas. Chambers were operated in flow-through mode, with air being drawn at
133 0.3 l min⁻¹ laterally from the headspace, and ambient air entering the chamber space
134 through a central chamber lid inlet (1 cm diameter) (Subke et al. 2009). Gaps between
135 chamber base and soil were filled using plastic film, whilst any remaining small leaks
136 at the chamber base being considered insignificant for measurements made in flow-
137 through mode. The CO₂ concentration and ¹³C:¹²C isotopic ratio in sample lines from
138 soil chambers were measured directly in the field using a mobile laboratory
139 containing an isotope ratio mass spectrometer (IRMS), referred to here as the York
140 Mobile Lab (see Subke et al. (2009) for more detail of chambers and laboratory). The
141 laboratory comprises a gas handling unit for switching between the 16 input lines,

142 connected to a Continuous-Flow Gas-Chromatograph IRMS (CF-GC-IRMS). A
143 standard laboratory gas chromatograph is coupled to a 12 cm radius magnetic sector
144 mass spectrometer (SIRAS Series2, Micromass, UK), NIER (Non-Ionizing
145 Electromagnetic Radiation) type ion impact source, triple faraday collector system,
146 rotary/turbo-molecular pumping vacuum system, interfaced to Microsoft Windows™
147 data system (model name “PVS12”, built by Pro-Vac Services, Crewe, UK). The CF-
148 GC-IRMS, together with a temperature control system and gas supplies for CO₂
149 reference and helium carrier gas, has been custom built to fit a modified twin-axle
150 trailer unit (Model ‘Tow-A-Van’, Indespension Ltd., Bolton, UK), thus providing the
151 unique opportunity to conduct isotope ratio measurements in real time and under field
152 conditions, with no need for off-line sampling and associated problems of gas
153 handling and storage. Transportation of the York Mobile Lab to the field site was by
154 helicopter. Power for running the instrumentation was provided by two propane
155 fuelled generators (Honda EU10i, converted for propane use; Honda Motor Co Ltd.,
156 Tokyo, Japan) with a total power output of 1.8 kW.

157 The air drawn continuously from all chambers was directed sequentially to the CF-
158 GC-IRMS. One measuring cycle (length of 1 h) consisted of measurements from all
159 12 respiration chambers as well as one reference gas injection, one measurement of
160 ambient air, and two measurements from respiration chambers placed on un-pulsed
161 control vegetation to measure natural abundance (NA) isotope ratios of respired CO₂.
162 While the respiration chambers on the labelling plots remained in place for the entire
163 duration of the monitoring period, the two NA chambers rotated between different
164 locations to capture all four vegetation types. The continuous CO₂ monitoring
165 chambers remained in place for four full days following the pulse, and an additional
166 8-hour period 7 days after the pulse.

167 The isotope ratio of the CO₂ flux was calculated using a 2-source mixing model:

$$\delta_F = \frac{\delta_{Sample} C_{Sample} - \delta_{Air} C_{Air}}{C_{Sample} - C_{Air}} \quad (\text{Equation 1})$$

168
169 where C and δ are, respectively, the CO₂ concentration and ¹³C/¹²C isotopic mixing
170 ratio of a gas, and the suffixes relate to ecosystem respiration flux CO₂ (F), CO₂ in
171 ambient air (Air), and CO₂ in the sample line ($Sample$). Weather conditions during the
172 pulse-chase period, with high wind speeds over extended periods (Fig. 3), meant that
173 there was significant leakage of air from the chamber head space. Since the ambient
174 air concentrations of ¹²C and ¹³C were subtracted from the sample gas, this leakage
175 had no effect on the observed isotope ratio detected in the CO₂ enrichment above
176 ambient concentrations. However, the absolute amount of C recorded in the sample
177 lines cannot be regarded as an accurate estimate of respiration from the area enclosed
178 by the chamber, as we cannot correct for the amount of air that leaked from each
179 chamber over time under high winds. We therefore concentrate here on isotopic ratio
180 changes of respiration-derived CO₂. For quality control purposes, $\delta^{13}\text{C}$ estimates were
181 only considered valid if the respiration-derived CO₂ in the sample gas amounted to
182 more than 50 $\mu\text{mol mol}^{-1}$ (an apparent CO₂ flux of 0.2 $\mu\text{mol m}^{-2} \text{s}^{-1}$). This requirement
183 led to the exclusion of isotopic data from lichen plots, which had very low CO₂ flux
184 rates, and therefore only small CO₂ concentration increases compared to ambient air
185 in the sample gas.

186 *Solid sample $\delta^{13}\text{C}$ analysis*

187 Coarse and fine roots and stones were removed from soil cores by sieving. The
188 organic soil material was mixed with any mineral soil fractions present, and sub-
189 samples of homogenised soil samples were ground for isotopic analyses. Fine roots

190 (<1 mm) were separated from coarser root material and also ground. For plots with
191 mixed communities, leaf material was sorted and analyses performed for the dominant
192 species only. For leaf samples, only green parts were used for sample processing (i.e.
193 excluding senesced leaves). For lichen plots, where only small biomass quantities
194 could be obtained, soil particles were carefully removed prior to sample processing.
195 Dried and ground leaf, root, and soil samples were analysed for their isotopic
196 composition on an elemental analyser (EA: FlashEA1112, ThermoFinnigan,
197 Germany) linked to a custom built IR-MS (with components identical to those
198 described above for the IRMS in the 'York Mobile Lab', and also constructed by Pro-
199 Vac Services Ltd., Crewe, UK). The instrumental output was calibrated against
200 certified isotopic material (NIST sucrose ANU 8542). Isotope results relate to the
201 organic matter contained in samples, i.e. they exclude mineral components present in
202 soil samples. All isotopic ratios are expressed in the delta notation (in ‰) and
203 reported relative to the Vienna Pee Dee Belemnite standard.

204 *Turnover calculations of assimilated C*

205 In order to assess differences in C turnover, we consider two general pools of C
206 turnover within the vegetation: (1) a labile C pool, and (2) leaf biomass C. All
207 assimilated C initially forms part of the labile, non-structural C pool (Pool 1), which
208 includes all water-soluble forms of carbohydrates (i.e. also sugars stored in cell
209 vacuoles). As this labile pool turns over, a small fraction of it becomes incorporated
210 into leaf biomass (Pool 2), whilst the remainder is either exported to other plant parts
211 via the phloem, or is respired by the leaf. In order to derive the fraction of assimilated
212 C built into structural biomass (carbon use efficiency, CUE), as well as calculating the

213 rate of turnover of labile C, we fitted the following exponential decay function to
214 isotopic enrichment results of both the biomass and respiration data:

$$215 \quad \delta = \delta_0 + ae^{-bt}, \quad (\text{Equation 2})$$

216 where δ is the $\delta^{13}\text{C}$ value of either biomass or respired CO_2 , t is the time since pulse
217 labelling (in days), and δ_0 , a , and b are fitted parameters. In doing so, we assume that
218 the overall pool sizes do not change over the observation period, i.e. that the reduction
219 in ^{13}C abundance in Pool 1 is caused by respiratory loss, phloem export or
220 incorporation into Pool 2, and not by dilution due to an increase in (unlabelled) C
221 subsequent to the ^{13}C pulse. From this, we estimate the mean residence time (MRT)
222 of labile C in foliage and total respiratory substrate as the inverse of the decay
223 constant (b):

$$224 \quad \text{MRT} = b^{-1}. \quad (\text{Equation 3})$$

225 For foliage data, MRT refers to labile C in leaves, whilst the MRT calculated for
226 ecosystem respiration is indicative of the turnover of labile C pools in both plants and
227 soil. The asymptote of the exponential decay function (δ_0) indicates the amount of
228 labelled C not affected by the exponential decay, i.e. the amount of C that would
229 remain in the leaf once the labile pool of labelled C is completely turned over. We
230 consider this to represent C fixed in more permanent molecular structures, mainly
231 structural components of the plant biomass. We therefore calculate the fraction of C
232 fixed as GPP during our pulse period and subsequently allocated to leaf biomass
233 within each plot according to:

$$234 \quad f_{SC} = (\delta_0 - \delta_{NA}) / (\delta_i - \delta_{NA}), \quad (\text{Equation 4})$$

235 where f_{SC} is the fraction of GPP allocated to structural carbon, δ_i is the initial
236 abundance of ^{13}C measured in the leaf biomass, δ_0 is the asymptote value obtained
237 from Equation 2, and δ_{NA} is the NA $\delta^{13}\text{C}$ value of the vegetation obtained from

238 samples taken before the pulse. We note that this calculation relates to leaf biomass
239 only, and does not represent the CUE of the whole plant. For biomass data harvested 1
240 year after the pulse, we also applied Equation 4, substituting δ_0 with measured
241 isotopic abundances in the biomass for each plot.

242 *Statistical methods*

243 Differences between average MRTs and f_{SC} values were assessed using two-way
244 ANOVA for vegetation type and C pool (MRT) and vegetation type and sampling
245 time (f_{SC}) as independent variables. Differences between vascular plant communities
246 in $\delta^{13}C$ values of root tissue and SOM were analysed using a repeated-measures
247 ANOVA (within-subjects factor = time post pulse; between-subjects factor = plant
248 community). Where ANOVAs indicated significance, differences between means
249 were assessed *post-hoc*, using the Holm-Sidak *t*-test. Although the assumption of
250 normality (i.e. Kolmogorov-Smirnov test) was not always within the $P > 0.05$
251 threshold (i.e. $n = 3$) we report those data as significance levels were high. All
252 statistical tests and regression parameter fittings were carried out using Sigma plot
253 11.0 software (Systat Software Inc., San Jose, California).

254 **Results**

255 *Short-term $\delta^{13}C$ changes in foliage samples*

256 Initial foliage enrichment in ^{13}C was found to be highest in *Betula*, indicating greater
257 assimilation rates per leaf biomass in these dwarf shrubs than in the other vegetation
258 types. Despite considerable spatial variation in leaf tissue enrichment, $\delta^{13}C$ values in
259 *Betula* leaves were significantly greater than all other vegetation types until 48 hours
260 after the pulse (Fig. 4). A sharp decline in enrichment over the following days meant

261 that this difference disappeared after this period. $\delta^{13}\text{C}$ values in both *Betula* and
262 graminoid foliage showed a consistent trend of diminishing ^{13}C abundance, following
263 an exponential decay curve over the sampling period (Fig. 4). In contrast to these
264 communities, no consistent trend was observed for either *Empetrum* or lichen plots,
265 the latter lacking any considerable increase in $\delta^{13}\text{C}$ (Fig. 4). C turnover tended to be
266 higher in *Betula* leaves, compared to graminoid foliage, but differences were not
267 statistically significant (Table 2). In contrast, the fraction of GPP allocated to plant
268 biomass in leaves was higher in graminoid compared to *Betula* plots. One year after
269 the pulse, less than 25% of pulse-derived C found after one week could be detected in
270 graminoid leaf biomass, whilst none was detected in *Betula* foliage (Table 2).

271 *Short-term $\delta^{13}\text{C}$ changes in ecosystem respiration*

272 Given the remoteness of the site and resulting logistical problems of transportation
273 and power provision, the York Mobile Lab performed consistently, enabling
274 measurement of the isotope ratio and CO_2 flux rates at frequent time resolution, with
275 data loss occurring on two separate occasions due to failure of the generators. This
276 caused data gaps of about 12 hours over the total monitoring period of 134 hours.

277 Figure 4 shows the changes in isotopic abundance of respiration over the 8-day period
278 for all *Betula*, graminoid, and *Empetrum* plots. Since total respiration on the lichen
279 plots was very low, no significant increase in CO_2 concentration in the sample gas
280 was recorded, and all data were removed by the data quality requirement (see above).
281 Isotope ratios immediately after the pulse showed extremely high values with a sharp
282 decline lasting about 16 hours post pulse. This pattern of a sharp decline from extreme
283 values, followed by a moderate increase, has been reported in other pulse chase
284 experiments (Högberg et al. 2008) and has been shown to result from physical tracer

285 diffusion in and out of soil pores (Subke et al. 2009). We included only data after this
286 initial period for the exponential decay regression in order to capture the return of C
287 allocated belowground by plants only. Given the shallow depth of soils, its
288 comparatively low porosity, and the likelihood of high rates of flushing of soil pores
289 under the windy conditions following the pulse (Fig. 3), we consider the abiotic tracer
290 return after this period to be insignificant.

291 Similar to the results found in leaf biomass, despite *Betula* vegetation tending to show
292 higher values than graminoid communities, there were no significant differences
293 between the mean residence times of labile C as evidenced in enrichment of
294 ecosystem respiration for these two vegetation types (Table 2).

295 *Root and SOM results*

296 Fine root tissue of the three vascular plant plots showed only a slight isotopic
297 enrichment, with no statistical differences between either plant communities or
298 sampling day (Fig. 5a). SOM results indicated significantly higher $\delta^{13}\text{C}$ values for
299 lichen plots, which are independent of the pulse treatment (Fig. 5b). Grouping all
300 results of vascular communities (i.e. excluding lichen plots), the $\delta^{13}\text{C}$ signal of SOM
301 across all communities was significantly enriched 7 days and still after 365 days post
302 pulse labelling.

303 **Discussion**

304 With a mean residence time of less than one day, our results document a fast rate of C
305 cycling at the peak of the short growing season at this site representing two key tundra
306 PFTs. The CUE of less than 20% at the leaf level is lower than observed in previous
307 experiments including herbaceous plants (Van Iersel 2003) and perennial grass

308 species (Lehmeier et al. 2010a; Lehmeier et al. 2010b). These leaf sample based
309 estimates mean that our CUE estimates are conservative, as phloem export and C
310 storage in other plant organs is not included in this estimate. However, the rapid
311 decline in the ^{13}C label detected in plant respiratory substrate (i.e. as $^{13}\text{CO}_2$) supports
312 the observation that the majority of assimilated C cycles through plants at a high rate.
313 Our findings contrast with results reported from $^{13}\text{CO}_2$ pulse-labelling in temperate
314 peatland communities of 1.9 and 4.7 days, respectively (Fenner et al. 2007; Ward et
315 al. 2009). A critical factor affecting the turnover of C between these contrasting
316 systems is likely to be the seasonality of GPP as well as NPP. The already
317 considerable difference between the two cited temperate peatland studies may relate
318 to the different seasons during which the ^{13}C pulse experiment was conducted (June in
319 Fenner et al. 2007 and October in Ward et al. 2009). Our results are representative for
320 conditions at the height of the arctic summer, and it is likely that the high rate of
321 diurnal assimilation combined with a very N limited system (Michelsen et al. 1996)
322 resulted in the observed high turnover of C, with the majority of initially fixed C
323 being respired. During periods of relatively lower diurnal GPP, a higher proportion of
324 fixed C may be turned into plant biomass (NPP), resulting in a longer overall MRT
325 and higher CUE of C in the ecosystem.

326 Our results further indicate that C available for respiration (which is likely to include
327 respiration from all plant organs as well as rhizospheric respiration involving
328 heterotrophic organisms) turns over at a faster rate than labile C found in leaves
329 (Table 2). Whilst both community types show the same pattern of shorter MRTs of
330 the respiratory substrate pool, the difference in turnover tended to be greater in the
331 dwarf birch plots compared to the graminoid communities. Such differences in
332 turnover both between PFTs and between different substrate pools within a particular

333 PFT require further investigation, as a better understanding of these rates is required
334 in order to enable meaningful process modelling of anticipated vegetation distribution
335 and shifts in environmental conditions (Dorrepaal 2007; Nobrega and Grogan 2008).
336 Our treatment of the labile C pool, which includes recent assimilates as well as
337 carbohydrates in other storage forms (e.g. sugars stored in cell vacuoles) is simplistic
338 compared to other schemes (Lehmeier et al. 2008; Lehmeier et al. 2010b), and may
339 confound effects of carbon use efficiency (NPP/GPP) with rates of respiratory
340 processing of different substrates. However, whilst we acknowledge this limitation,
341 the results nonetheless allow a comparative investigation of C allocation in different
342 PFTs. This is likely to relate to differences in C transport to belowground plant
343 organs. Despite lacking statistical significance, a trend towards faster allocation of
344 recent assimilates to roots in *B. nana* emerges, which is not observed in the graminoid
345 community (Fig. 5a). This is corroborated by the trend of slightly faster turnover of
346 labile C in graminoid plots compared to *B. nana* dominated vegetation, as evidenced
347 in the marginally shorter mean residence time of labile substrates, even if here also the
348 differences are not significant. Whilst the low level of replication ($n = 3$) means that
349 these trends show statistical significance in only a few instances, a general difference
350 in C allocation pattern emerges, with fast C turnover and a relatively small
351 belowground allocation of recent assimilates in graminoid communities, and
352 comparatively slower turnover and greater investment in belowground allocation by
353 the dwarf shrub communities. Previous work by Shaver and Chapin (1991) has shown
354 that distinct differences in allocation patterns and C turnover between vegetation
355 forms in the tundra at leaf level can be compensated by storage processes in stems,
356 resulting in similar C turnover dynamics between communities. The results for label
357 retention in foliar biomass after 1 year are consistent with the deciduous habit of *B.*

358 *nana*, where no label-derived ^{13}C could be detected, whilst the graminoid
359 communities still showed significant enrichment. These findings corroborate results
360 reported on the basis of biomass and nutrient abundances for characteristic divergence
361 in plant C pools above- and below ground between PFTs, and seasonally different
362 allocation patterns related to growth form and leaf habit (Chapin et al. 1980).

363 The absence of significant enrichment in *Empetrum* leaf samples was a
364 surprising result. The CO_2 flux results clearly showed a respiratory return of ^{13}C
365 similar to the other vascular plots, indicating that significant amounts of the isotopic
366 tracer were assimilated by the vegetation. This is corroborated by CO_2 gas exchange
367 measurements obtained by automated NEE chambers at the same site, where
368 *Empetrum*, *Betula*, and graminoid communities show similar assimilation values (both
369 in terms of mean daily NEE or peak daytime peak NEE rates; data not shown; R.
370 Poyatos Lopez, *pers. com.*). The results suggest that only small amounts of
371 assimilated C were stored in the foliage of *E. nigrum*. We suspect that the inconsistent
372 temporal pattern of enrichment in *E. nigrum* foliage results from a high degree of
373 heterogeneity in photosynthetic activity, particularly during relatively dry conditions,
374 as in our case. Whilst care was taken to sample representative parts of the *Empetrum*
375 canopy, the absence of significant tracer assimilations in the sampled foliage could
376 therefore be caused by an inadvertent preference for less active areas within the
377 canopy.

378 Notwithstanding the small-scale experimental work, this study shows different rates
379 of C turnover and belowground allocation of recent assimilates between key tundra
380 plant communities, and warrants further long-term validation in relation to C pool
381 turnover modelling in such mosaic landscapes containing considerable C stocks.

382

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393

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- 516

517 **Table 1**

518 Four contrasting plant communities studied during $^{13}\text{CO}_2$ pulse-labelling experiment.

519 All plots (n = 3 per community type) were located within an area of 15 x 20 m.

520 Species names in brackets indicate subdominant species. LAI estimates are mean \pm 1

521 SD, and in the “Exposed Ridge” community refers to vascular plants located within

522 the plot; biomass sampling in these plots following the isotope pulse was carried out

523 for lichen biomass only.

524

Community	Dominant species	LAI
Dwarf birch	<i>Betula nana</i> (<i>Empetrum nigrum</i>)	2.09 \pm 0.31
Empetrum heath	<i>Empetrum nigrum</i>	1.66 \pm 0.13
Graminoid	<i>Carex nigra</i> (<i>Tofieldia pusilla</i> , <i>Scirpus cespitosus</i>)	1.60 \pm 0.32
Exposed ridge	Lichen (predominantly <i>Cetraria</i> and <i>Cladonia</i> spp.)	0.27 \pm 0.15

525

526

527 **Table 2**

528 Carbon turnover in foliage of *Betula nana* and graminoid spp. Mean residence times
 529 (MRTs) and fractions of GPP incorporated into structural carbon (f_{SC}) derived from
 530 isotopic results. Data are mean \pm 1 SE (n = 3). Two-way ANOVA showed significant
 531 differences in MRT between C pools (i.e. between leaf biomass and respiration
 532 substrate pools, $p = 0.011$) across vegetation types, but no differences between
 533 vegetation types or interactions between pools and vegetation ($p = 0.117$ and
 534 $p = 0.226$, respectively). The amount of GPP incorporated into biomass differed
 535 between time points (1 week or 1 year; $p = 0.001$), with no significant difference
 536 between vegetation types or interaction between both ($p = 0.163$). However, a Post-
 537 hoc Hiolm-Sidak test revealed a significant difference between vegetation types 1
 538 year after the pulse ($p = 0.007$)

539

		<i>Betula nana</i>	Graminoid
MRT (leaf biomass)	days	1.093 \pm 0.223	0.667 \pm 0.164
MRT (respiration)	days	0.456 \pm 0.007	0.394 \pm 0.025
f_{SC} (1 week)	-	0.160 \pm 0.020	0.207 \pm 0.054
f_{SC} (1 year)	-	-0.002 \pm 0.002	0.046 \pm 0.016

540

541

542 **Figure legends**

543 **Figure 1**

544 Map of the study area in Swedish Lapland (inset map). The experiment was sited at
545 about 700 m asl on the slope from the Nissoncorru mountain in the south-east to lake
546 Torneträsk in the north. The hatched line indicates the approximate position of the
547 altitudinal tree line of mountain birch (*Betula pubescens* Ehrh.).

548

549 **Figure 2**

550 Schematic of a Perspex hood used for $^{13}\text{CO}_2$ pulse labelling (a) and a PVC respiration
551 chamber (b). $^{13}\text{CO}_2$ pulse gas was flushed through Perspex hoods at approximately
552 5 L min^{-1} . Sample air from PVC respiration chambers was drawn from two points in
553 the chamber (only one shown here) to the York Mobile Lab, and ambient air entered
554 the chamber headspace via a central vent. Plastic film (“bubble wrap”) was used to fill
555 larger gaps of both the pulse hoods and the respiration chambers.

556

557 **Figure 3**

558 Meteorological conditions during July 2007 at the research site. The dark shaded area
559 indicates the period of the 3-h pulse period on the 4th July 2007, whilst lighter shaded
560 areas represent the CO_2 flux monitoring periods.

561

562 **Figure 4**

563 Carbon isotope ratios following the 3-hour pulse period in leaf biomass (left column)
564 and ecosystem respiration (right column). Results are means and error bars indicate
565 Standard Errors for three replicate plots of *Betula nana* (a, b), graminoid vegetation

566 (c, d), *Empetrum nigrum* (e, f), and lichen plots (g). Lines are fitted exponential decay
567 curves for means. No meaningful isotope ratios were obtained for lichen plots due to
568 the low CO₂ flux (see text). Enrichment was significantly greater in *Betula* foliage
569 until 48 hours after the pulse ($p = 0.016$ ANOVA with Tukey post-hoc test at 48 h
570 post-pulse). Note difference in y-axis scale between graphs in left-hand and right-
571 hand columns.

572

573 **Figure 5**

574 $\delta^{13}\text{C}$ values of (a) fine root tissue from soil cores sampled in the three vascular plant
575 communities, and (b) soil organic matter in all plots. Data show isotopic abundance
576 immediately before the pulse (day 0, representing natural isotopic abundance), and at
577 1 (roots only), 7, and 365 days after the pulse. There were no significant differences
578 or interactions between sampling days and plant species for root tissue. For SOM
579 results (b), a 2-way ANOVA indicate a significant effect of plant community
580 ($p < 0.001$), with lichen plots having significantly enriched values compared to all
581 other plots ($p < 0.02$, Holm-Sidak post-hoc test). A two-way ANOVA performed for
582 SOM data grouping all vascular plant communities (i.e. excluding the lichen plots)
583 showed a significant influence of the sampling day ($p = 0.018$), with values on day 0
584 being significantly lower than either of the other days across all vascular
585 communities.

Figure 1

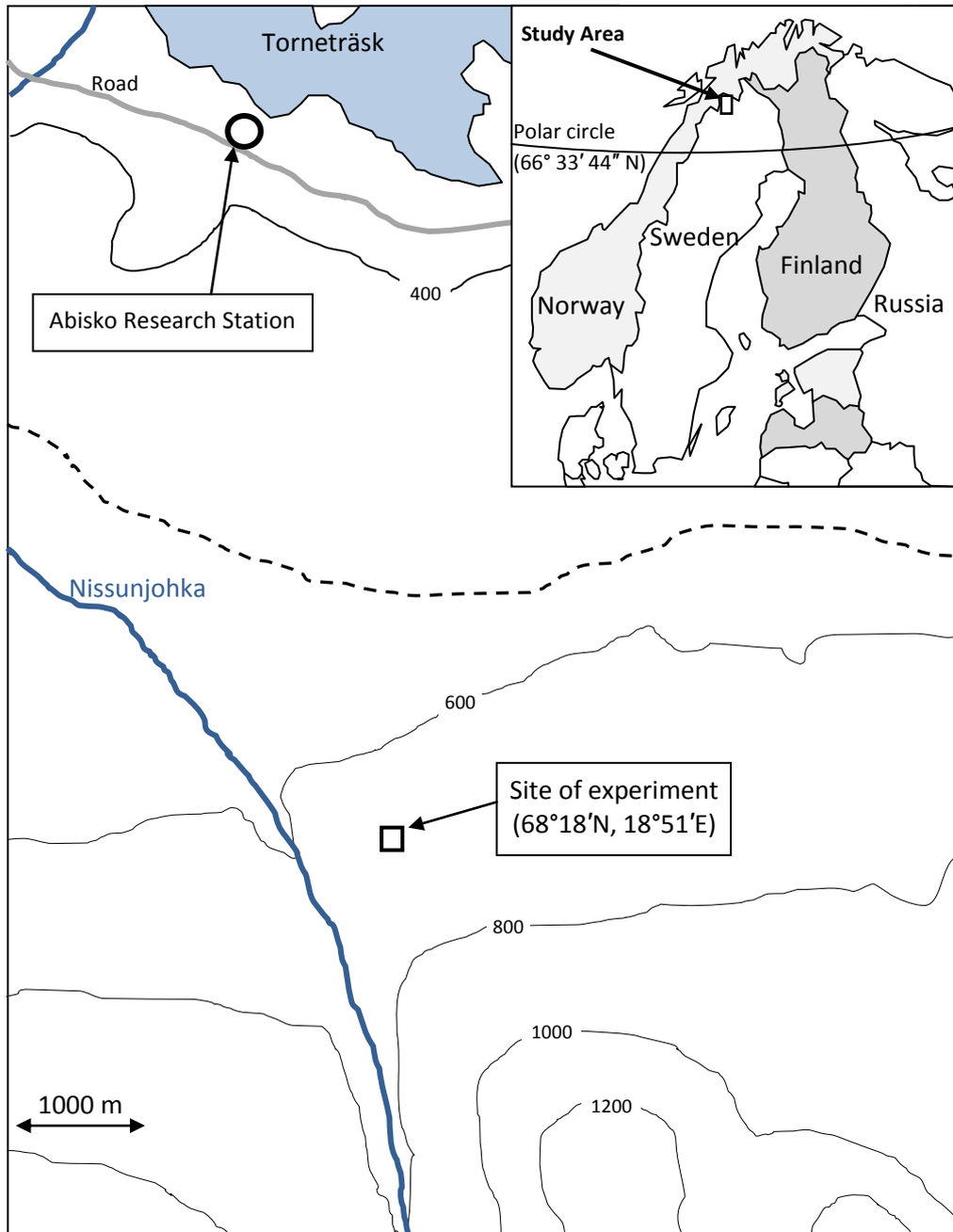


Figure 2

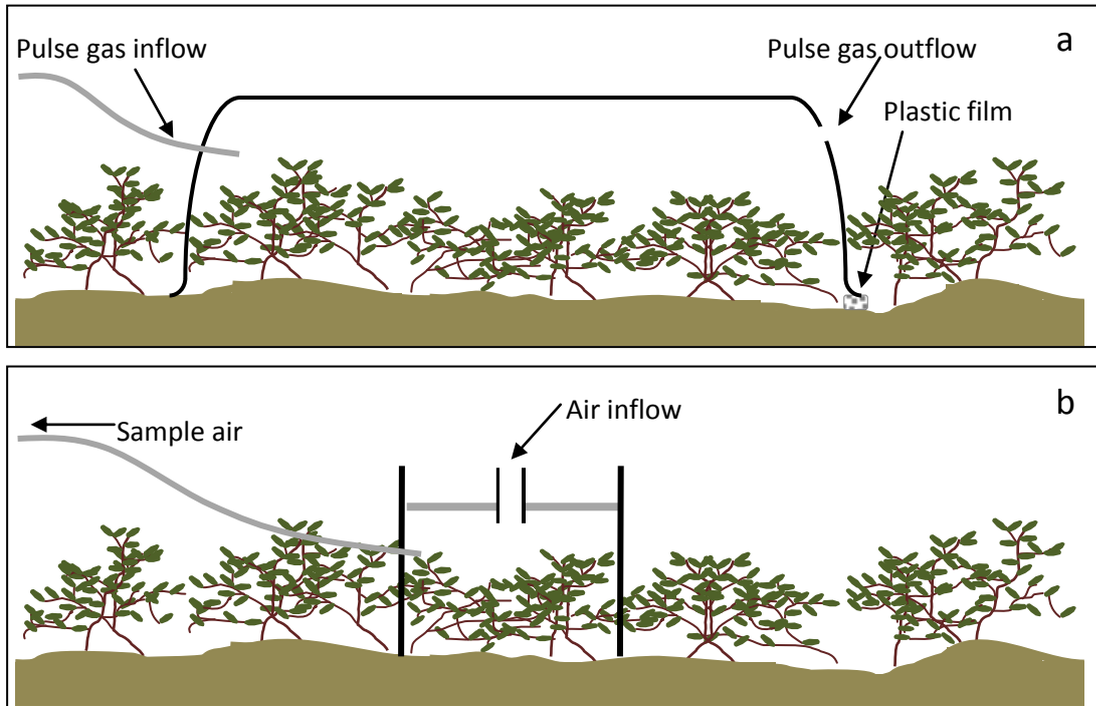


Figure 3

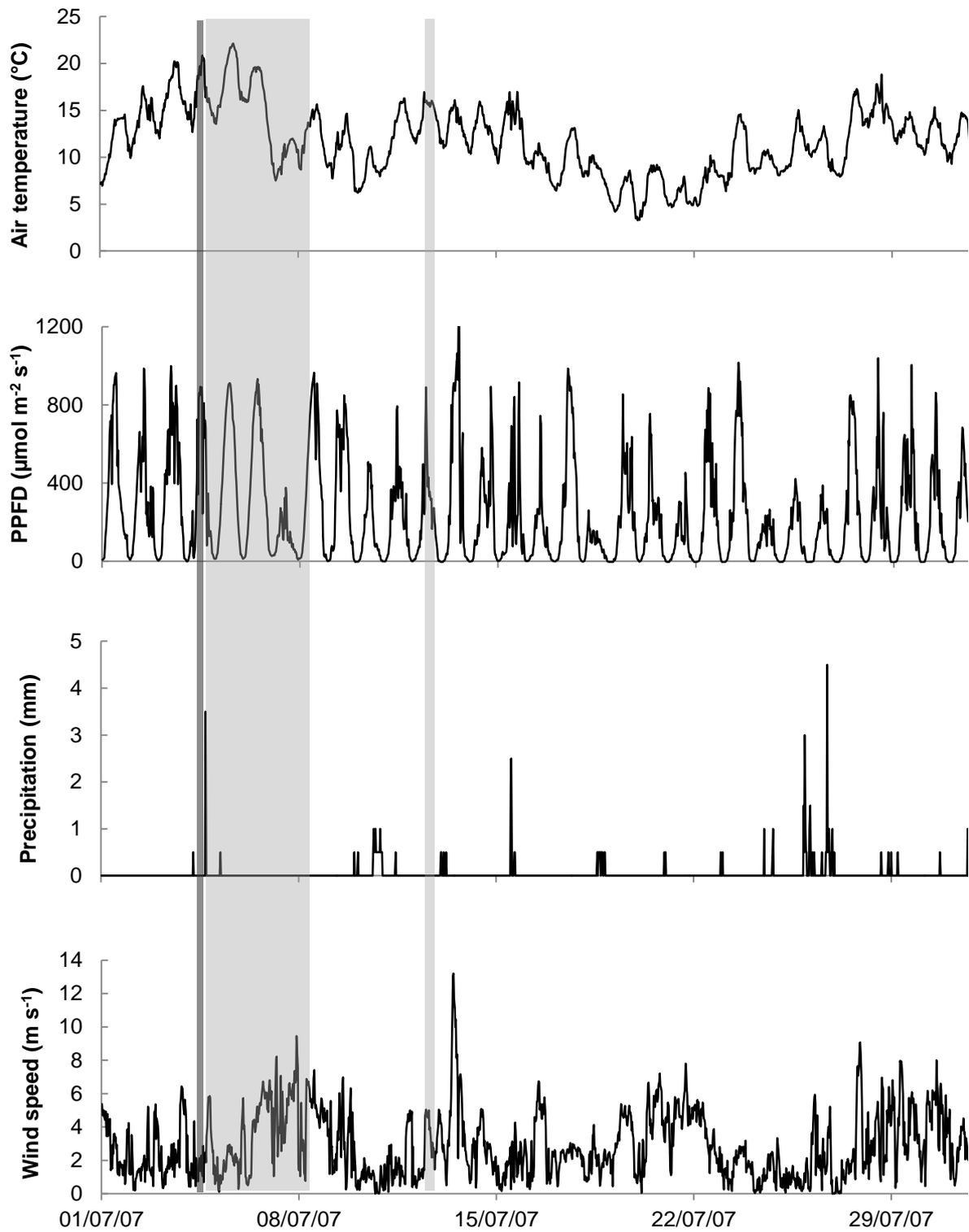


Figure 4

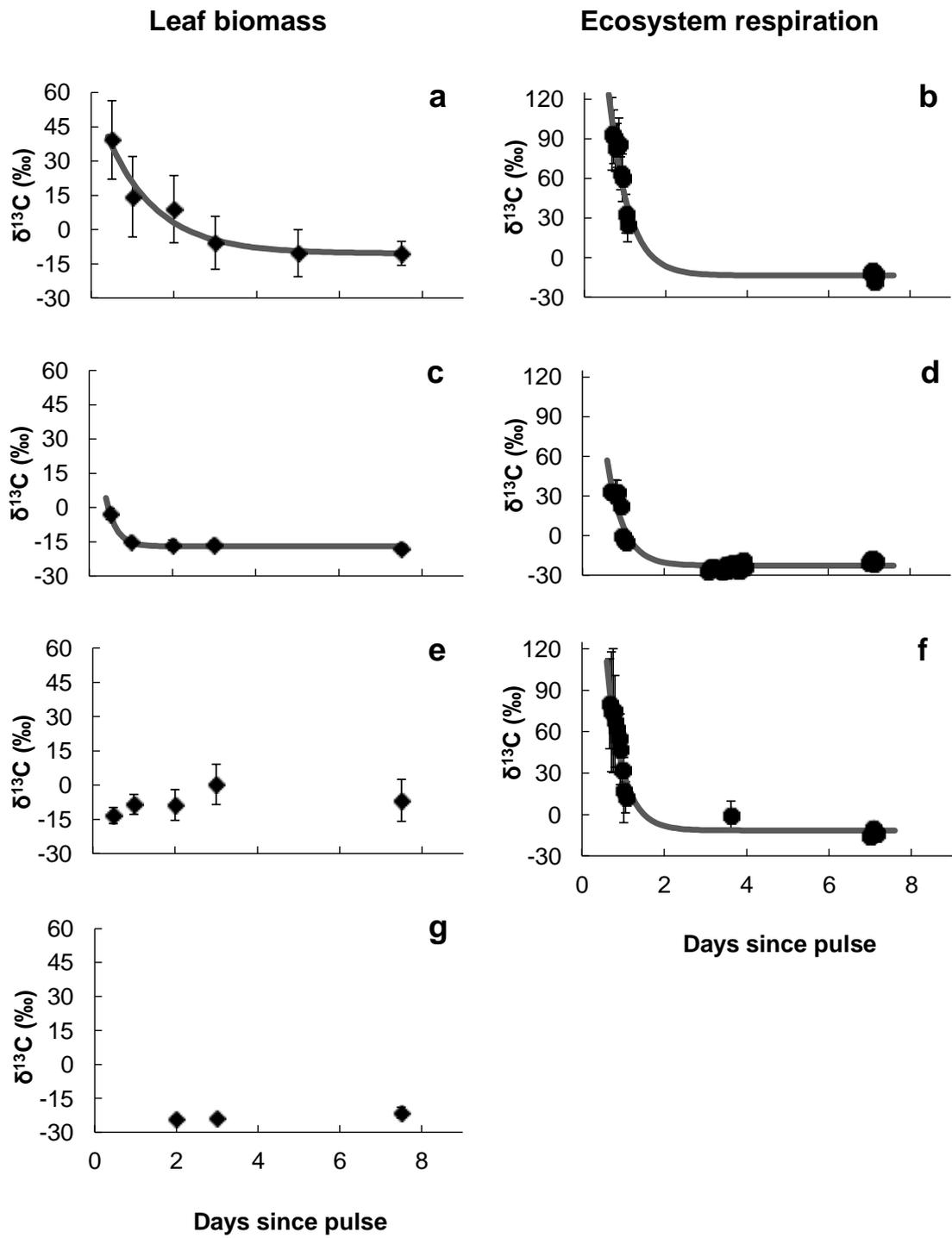


Figure 5

