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3 **Amazon deforestation alters small stream structure, nitrogen biogeochemistry and**  
4 **connectivity to larger rivers**

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7 Linda A. Deegan<sup>1\*</sup>, Christopher Neill<sup>1</sup>, Christie L. Haupt<sup>1,4</sup>, M. Victoria R. Ballester<sup>2</sup>,

8 Alex V. Krusche<sup>2</sup>, Reynaldo L. Victoria<sup>2</sup>, Suzanne M. Thomas<sup>1</sup>, and Emily de Moor<sup>3,5</sup>

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10 <sup>1</sup>The Ecosystems Center, Marine Biological Laboratory, 7 MBL Street, Woods Hole, MA 02543,

11 USA

12 <sup>2</sup>Laboratório de Análise Ambiental e Geoprocessamento, Centro de Energia Nuclear na

13 Agricultura, Universidade de São Paulo, Caixa Postal 1341 6000, Piracicaba, SP, Brazil

14 <sup>3</sup>Department of Ecology and Evolutionary Biology, Brown University, 80 Waterman Street,

15 Providence, RI 02912, USA

16 <sup>4</sup>CH2M Hill Polar Services, 2325 King Road, Fairbanks, AK 99709, USA

17 <sup>5</sup>Current address: Department of Geography, University of California Santa Barbara, 1832

18 Ellison Hall, Santa Barbara, CA 93106, USA

19 \*Corresponding author: ldeegan@mbl.edu

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24

## 25 Abstract

26 Human activities that modify land cover can alter the structure and biogeochemistry of  
27 small streams but these effects are poorly known over large regions of the humid tropics where  
28 rates of forest clearing are high. We examined how conversion of Amazon lowland tropical  
29 forest to cattle pasture influenced the physical and chemical structure, organic matter stocks and  
30 N cycling of small streams. We combined a regional ground survey of small streams with an  
31 intensive study of nutrient cycling using  $^{15}\text{N}$  additions in three representative streams: a second-  
32 order forest stream, a second-order pasture stream and a third-order pasture stream that were  
33 within several km of each other and on similar soils and landscape positions. Replacement of  
34 forest with pasture decreased stream habitat complexity by changing streams from run and pool  
35 channels with forest leaf detritus (50% cover) to grass-filled (63% cover) channel with runs of  
36 slow-moving water. In the survey, pasture streams consistently had lower concentrations of  
37 dissolved oxygen and nitrate ( $\text{NO}_3^-$ ) compared with similar-sized forest streams. Stable isotope  
38 additions revealed that second-order pasture stream had a shorter  $\text{NH}_4^+$  uptake length, higher  
39 uptake rates into organic matter components and a shorter  $^{15}\text{NH}_4^+$  residence time than the  
40 second-order forest stream or the third-order pasture stream. Nitrification was significant in the  
41 forest stream (19% of the added  $^{15}\text{NH}_4^+$ ) but not in the second-order pasture (0%) or third-order  
42 (6%) pasture stream. The forest stream retained 7% of added  $^{15}\text{N}$  in organic matter compartments  
43 and exported 53% ( $^{15}\text{NH}_4^+ = 34\%$ ;  $^{15}\text{NO}_3^- = 19\%$ ). In contrast, the second-order pasture stream  
44 retained 75% of added  $^{15}\text{N}$ , predominantly in grasses (69%) and exported only 4% as  $^{15}\text{NH}_4^+$ .  
45 The fate of tracer  $^{15}\text{N}$  in the third-order pasture stream more closely resembled that in the forest  
46 stream, with 5% of added N retained and 26% exported ( $^{15}\text{NH}_4^+ = 9\%$ ;  $^{15}\text{NO}_3^- = 6\%$ ). These  
47 findings indicate that the widespread infilling by grass in small streams in areas deforested for

48 pasture greatly increases the retention of inorganic N in the first- and second-order streams,  
49 which make up roughly three-fourths of total stream channel length in Amazon basin watersheds.  
50 The importance of this phenomenon and its effect on N transport to larger rivers across the larger  
51 areas of the Amazon Basin will depend on better evaluation of both the extent and the scale at  
52 which stream infilling by grass occurs, but our analysis suggests the phenomenon is widespread.

### 53 **Introduction**

54 Human activities that modify land cover have the potential to alter the physical structure  
55 of small streams and the manner in which nitrogen (N) is transformed and transported in stream  
56 networks (Sweeney et al. 2004; Mulholland et al. 2008). The Amazon Basin contains more than  
57 4 million km<sup>2</sup> of tropical forest and the earth's largest river network. It also has the world's  
58 highest rate of rainforest clearing, primarily to pasture for cattle ranching (Skole and Tucker  
59 1993; Lepers et al. 2005). Approximately 697,830 km<sup>2</sup> of forest have been cleared in Brazil since  
60 1970 (INPE 2010). This clearing has the potential to alter the structure and chemistry of the  
61 thousands of km of small streams that constitute three-fourths of stream channel lengths in the  
62 Amazon Basin (McClain and Elsenbeer 2001; Goulding et al. 2003).

63 Small streams are the primary receptors for nutrients and organic matter inputs from  
64 terrestrial ecosystems and play a key role as regulators of downstream flows of materials to  
65 larger rivers (Vannote et al. 1980; Alexander et al. 2000; Peterson et al. 2001). In the Amazon,  
66 streams act as key connections with terrestrial ecosystems (McClain and Elsenbeer 2005) and  
67 also are important habitat for fishes and aquatic organisms (Agostinho et al. 2005). Previous  
68 work conducted in the Large-Scale Biosphere-Atmosphere (LBA) Experiment in Amazonia  
69 showed that clearing of Amazon forest for pasture can change the flowpaths of water reaching  
70 streams (Germer et al. 2009), the structure and hydrology of stream channels (Neill et al. 2006)

71 and the chemistry of streamwater (Thomas et al. 2004; Biggs et al. 2004; Neill et al. 2006).  
72 Establishment of cattle pasture has also been shown to alter terrestrial N cycling by reducing the  
73 production of nitrate ( $\text{NO}_3^-$ ) in soils, resulting in low concentrations of  $\text{NO}_3^-$  in soil solution and  
74 reducing the potential for  $\text{NO}_3^-$  movement from land into small streams (Markewitz et al. 2004;  
75 Neill et al. 1996, 1997; Chaves et al. 2009).

76         There are several reasons why the dynamics of N transport and retention in Amazon  
77 streams may be different from streams in temperate regions where streams are recognized as  
78 important locations of N uptake and transformation (Peterson et al. 2001). High concentrations  
79 of N and low ratios of nitrogen to phosphorous (N:P) in tropical forest foliage and litter suggest  
80 that N is not limiting to forest vegetation (Vitousek 1984; Davidson et al. 2007). Relatively high  
81 rates of soil N mineralization and nitrification (Neill et al. 1995), abundant soil solution  $\text{NO}_3^-$   
82 (Markewitz et al. 2004; Neill et al. 2006) and high soil emissions of  $\text{N}_2\text{O}$  (Verchot et al. 1999;  
83 Melillo et al. 2001) all indicate natural N “saturation” (Hall and Matson 1999) of the forest,  
84 which could lead to high transport of inorganic N in soil and ground water to forest streams.  
85 Under these conditions, we would expect tropical forest streams to be less retentive of N than  
86 similar-sized streams in temperate regions. Because clearing of Amazon forest for pasture  
87 introduces conditions that indicate greater N limitation of vegetation, including lower N  
88 concentrations in foliage and litter, lower rates of soil nitrification and less  $\text{NO}_3^-$  in soil solution  
89 (Davidson et al. 2007; Chaves et al. 2009), we would also expect pasture streams to be more  
90 retentive of N than forest streams.

91         Much of our recent understanding of stream N dynamics has come from experiments in  
92 which low levels of dissolved inorganic N enriched with  $^{15}\text{N}$ , either as nitrate ( $^{15}\text{NO}_3^-$ ) or  
93 ammonium ( $^{15}\text{NH}_4^+$ ), have been added to streams for days to weeks. The fate of the added  $^{15}\text{N}$  is

94 then followed as fluvial export or into different inorganic and organic matter compartments in  
95 the stream channel (Peterson et al. 2001). Most experiments have introduced  $^{15}\text{NH}_4^+$ , which  
96 allows quantification of nitrification and subsequent uptake of produced  $\text{NO}_3^-$  as well as uptake  
97 of  $\text{NH}_4^+$  into biomass compartments (Mulholland et al. 2000). Because they have been conducted  
98 in a number of streams ranging from Alaska to Puerto Rico using similar methodology, these  
99 experiments can now be used to compare N dynamics across different sites (Webster et al. 2003).

100 We quantified how clearing moist tropical forest for cattle pasture along small, lowland  
101 streams in one of the Amazon's deforestation hotspots, central Rondônia, influenced stream  
102 physical and chemical structure, and stream channel N uptake, transformation and retention. We  
103 combined a regional ground survey of small streams with an intensive study of nutrient cycling  
104 using  $^{15}\text{N}$  additions in three representative streams: a second-order forest stream, a second-order  
105 pasture stream and a third-order pasture stream that were within several km of each other and on  
106 similar soils and landscape positions. We used the results to infer how forest clearing influenced  
107 stream structure, the dynamics of N biogeochemistry in the stream channel and the role that  
108 small streams play in controlling the movement of N in stream networks in forested and  
109 deforested regions of the Amazon.

## 110 **Methods**

### 111 *Regional stream survey*

112 We surveyed physical and chemical characteristics of first- and second-order streams that  
113 drained watersheds (0.1 to 17.8 km<sup>2</sup>) dominated by either forest (6 streams) or pasture (12  
114 streams) near Nova Vida, Rancho Grande and Ji Paraná in central Rondônia in August 2005  
115 (Fig. 1, Table 1). Target streams were identified from Landsat 7-ETM+ images (Ballester et al.  
116 2003). We then combined an analysis of land cover and stream networks derived from Landsat

117 7-ETM+ and IKONOS remote sensing in the 74,057 km<sup>2</sup> Ji-Paraná River Basin (Fig. 1) to  
118 estimate the extent to which deforestation has altered streams at the river-basin scale. Aerial  
119 images analysis was used to determine the extent of *Paspalum repens* infilling of stream  
120 channels by combining the Landsat ETM image with low level videography data (obtained from  
121 INPE 2010) and IKONOS data (obtained from EOS-WEBSTER 2009). IKONOS images are  
122 color composites of the multispectral bands of 2000-2002 IKONOS with a spatial resolution of 4  
123 ×4 m acquired and licensed to NASA LBA-Ecology Program, the NASA Scientific Data Buy  
124 Project and The University of New Hampshire Earth Science Information Partner (EOS-  
125 WEBSTER). Selected images were ground-truthed during the regional stream survey (above)  
126 and used to classify the full array of IKONOS and videography images.

127         Streams were classified as “forest” streams if >75% of the catchment was in native forest  
128 and “pasture” streams if >65% of the catchment was in pasture. Native forest vegetation  
129 consisted of perennial evergreen broadleaf trees with a high number of palms (Pires and Prance  
130 1986). Forest in the region is typically cleared for pasture to the stream edge to allow access by  
131 cattle (approximately 1 to 1.5 animal per ha) and converted directly to pasture by planting with  
132 introduced forage grasses of the genus *Brachiaria* in upland areas. Riparian areas are often  
133 dominated by the native grass *P. repens* (Neill et al. 2006). Pasture in this region is never  
134 fertilized with N and only rarely is P added. The climate of central Rondônia is humid tropical,  
135 with a mean annual temperature of 26°C, mean annual precipitation of about 2200 mm y<sup>-1</sup> and a  
136 dry season that last approximately 5 months (June to October). (Bastos and Diniz 1982). All  
137 sites were in areas of gently rolling topography underlain by Pre-Cambrian granite with well-  
138 drained Kandiodults and Paleodults soils that cover approximately 22% of the Brazilian Amazon  
139 (Moraes et al. 1995).

140 We surveyed stream width, depth and benthic substrate type (classified as tree leaf pack,  
141 living riparian grass, sand, fine organic, wood, or other) with 11 cross-sectional point transects in  
142 100 m (first-order streams) or 800 m (second-order streams) stream reaches. Conductivity and  
143 dissolved oxygen (YSI Model 58) and pH (Orion 290A+ meter) were measured in the field.  
144 Discharge was determined from cross sectional area and flow velocity (FP-201 Global Water  
145 flow meter). Water samples for  $\text{NH}_4^+$ ,  $\text{NO}_3^-$  and soluble reactive phosphate (SRP) were filtered  
146 in the field through ashed GFF filters and frozen. Whole water samples were returned to the lab  
147 for alkalinity and suspended particulate organic matter (SPOM) analysis.

148 We analyzed differences in mean physical, structural and chemical characteristics of forest  
149 and pasture streams using one-way analysis of variance (SAS Institute 2002). Tests for percent  
150 cover were performed after arcsin-square root transformation. Other environmental variables  
151 were tested on non-transformed data.

#### 152 *Stream $^{15}\text{N}$ additions*

153 We selected three streams typical of the region located on Nova Vida Ranch (Fig. 1,  
154  $10^{\circ}09'\text{S}$ ,  $62^{\circ}49'\text{W}$ ) for whole-stream additions of  $^{15}\text{NH}_4^+$ : (1) a second-order forest stream; (2) a  
155 second-order pasture stream and; (3) a third-order pasture stream located approximately 2 km  
156 downstream of the confluence of the second-order forest and pasture streams. Pastures were  
157 cleared in 1989 (second-order stream) or 1983 (third-order stream) and planted directly to  
158 *Brachiaria brizantha*. These streams were in similar landscape positions and on similar soils  
159 (Kandiudults) (Moraes et al. 1996). Our interpretation uses the “space for time” substitution for  
160 land use (Allan 2004). This approach assumes that for locations that differ in land use but are  
161 similar in other respects, differences between sites can be viewed as equivalent to changes over

162 time in a single location. There was no third-order stream in a forested watershed available for  
163 study in this region of expanding deforestation and fragmented forests.

164 The second-order forest stream (watershed area 17.8 km<sup>2</sup>) was shaded, had a pool and run  
165 stream channel with a sandy bottom, a mean wetted width of 4 m, a maximum depth of 42 cm  
166 and dry season discharges of 15-40 L s<sup>-1</sup>. The second-order pasture stream (watershed area of 8.4  
167 km<sup>2</sup>) was bordered by the C4 grass *P. repens* (Medina et al. 1976) along its entire length, had  
168 slow-moving deep runs with extensive channel infilling by grass, deep (> 15 cm in some places)  
169 organic sediments underlain by sand with a mean wetted channel width of 5.2 m, an open water  
170 channel width of 1.4 m and a mean depth of 42 cm. Dry season discharges range from 14 to 90 L  
171 s<sup>-1</sup>. The third-order pasture stream (watershed area of 27 km<sup>2</sup>) had high sandy banks, a pool and  
172 run channel structure with a generally sandy bottom and lacked extensive growth of grass in the  
173 stream channel. Although this stream had some riparian trees, the canopy was relatively open  
174 and most of the stream was exposed to the sun. It had a wetted channel width of 3.6 m, a nearly  
175 identical open water channel width of 3.5 m and a mean depth of 34 cm. It had dry season  
176 discharges of 45 to 100 L s<sup>-1</sup>. We selected representative reaches of 800 m (forest stream), 500 m  
177 (second-order pasture stream) and 760 m (third-order pasture stream) that had no obvious  
178 tributaries or other obvious surface water inputs. Although the third-order stream was  
179 downstream of the second-order pasture stream and the <sup>15</sup>N additions were done simultaneously,  
180 <sup>15</sup>N values at the start of the third-order stream reach were never elevated above baseline.  
181 Additionally, solute additions demonstrated that N added to the second order stream did not  
182 reach third-order pasture stream (Neill et al. 2006).

183 Our <sup>15</sup>N addition methodology, sample analysis and calculations followed the protocols  
184 in LINX (Lotic Intersite Nitrogen eXperiment) (Mullholland et al. 2000, Peterson et al. 2001).

185 We determined physical characteristics, organic matter standing stocks and background natural  
186 abundance levels of  $^{15}\text{N}$  for each stream reach. We then added 98.5% enriched  $^{15}\text{N}$ -labeled  
187  $\text{NH}_4\text{Cl}$  at a constant rate to each stream for 21 d during the dry season to achieve an expected  
188  $\delta^{15}\text{N}$  of  $\text{NH}_4^+$  of 500 ‰ ( $0.28 \text{ g d}^{-1}$ , totaling 5.90 g in the forest stream,  $0.98 \text{ g d}^{-1}$ , totaling 21.50  
189 g in the second-order pasture stream and  $0.86 \text{ g d}^{-1}$ , totaling 18.90 g in the third-order pasture  
190 stream). We measured  $^{15}\text{N}$  in major ecosystem compartments upstream and downstream of the  
191 addition site before, during and for 14 d after the addition. The forest addition began on 12  
192 August 2001 and the two pasture stream additions began on 15 August 2003.

193 Water level, dissolved oxygen concentration, and conductivity were recorded every 15  
194 min throughout the  $^{15}\text{N}$  additions (YSI 600XL multi-parameter sondes). Streamwater samples  
195 were collected every 2 to 3 days and discharge determined every 3 to 7 days or when rains  
196 caused notable changes to stream flow (see Regional Survey for field methods). Multiple sodium  
197 chloride additions following standard solute injection methods (Stream Solute Workshop 1990)  
198 were used in our previous work (Neill et al. 2006) and during the  $^{15}\text{N}$  additions at characteristic  
199 discharges to determine the extent of groundwater and surface water inputs in each reach. During  
200 the  $^{15}\text{N}$  additions, we completed two  $\text{Cl}^-$  additions (discharges of 30 and  $45 \text{ L s}^{-1}$ ) in the forest, one  
201 in the second-order pasture (discharge  $45 \text{ L s}^{-1}$ ) and one in the third-order (discharge  $105 \text{ L s}^{-1}$ )  
202 stream. In all three streams, no surface water channels were observed flowing into the stream  
203 reaches and no measurable gains or losses of  $\text{Cl}^-$  tracer were detected over the study reaches.  
204 These measurements suggest that these stream reaches were neither measurably gaining nor  
205 losing water during the experiments.

206 We combined measurements of organic matter biomass with estimates of the area of each  
207 benthic habitat type to calculate standing stocks of organic matter and N for the stream reaches.

208 Stream depth and benthic substrate area and type (classified as grass, leaf pack, sand, detritus,  
209 thin layer of fine organic matter over sand, clay, gravel, and woody debris) were determined  
210 using cross-sectional point transects in each reach (N=75 cross sections in forest, 45 in second-  
211 order and 39 in third-order pasture streams). Leaf pack, detritus and fine organic matter were  
212 layers of varying thickness (from a few cm to mm) over a sandy substrate. Very small patches of  
213 filamentous algae were found but their area (< 0.01% of stream bottom) did not warrant a  
214 separate habitat class.

215 Biomass of leaf pack (nearly entire, identifiable tree leaves), coarse benthic organic  
216 matter (CBOM; unidentifiable leaf fragments and organic material > 1 mm) and fine benthic  
217 organic matter (FBOM, < 1 mm) were quantified by pressing an open-ended cylinder (0.0573  
218 m<sup>2</sup>) 5 cm into the sediment, collecting and sieving all organic matter in the cylinder. The riparian  
219 *P. repens* grass habitat was divided into: (1) channel edge (submersed or emergent in the stream  
220 channel), and (2) floodplain (2 m from the open channel but out of direct contact). Both channel  
221 edge and floodplain grass habitats were found in the second-order pasture, while only the  
222 channel edge grass habitat was found in the third-order pasture stream. No riparian grass was  
223 present in the forest stream. Leaves (above-water green tissue) and roots and rhizomes (below-  
224 water) biomass were estimated using 0.25 m<sup>2</sup> quadrates. All organic matter samples were rinsed  
225 with tap water and dried at 60 °C.

226 We measured the <sup>15</sup>N content of NH<sub>4</sub><sup>+</sup> and NO<sub>3</sub><sup>-</sup> and the <sup>15</sup>N content in organic matter  
227 compartments (CBOM, FBOM, SPOM, filamentous algae and leaves) in longitudinal transects  
228 (7 to 8 stations in the reach) once before the start, 5 times during, and 4 times post <sup>15</sup>N addition.  
229 We sampled *P. repens* on day 20. We examined <sup>15</sup>N uptake by the microbial community on

230 ceramic tiles conditioned for 7-d prior to  $^{15}\text{N}$  addition. We simultaneously measured  $\delta^{13}\text{C}$  (‰) in  
231 the organic matter compartments to examine the origin of organic matter in the stream channel.

### 232 *Laboratory analyses*

233 Concentrations of  $\text{NO}_3^-$ ,  $\text{NH}_4^+$ , SRP and total dissolved nitrogen (TDN) were determined  
234 by ion chromatography (Dionex DX-500) or by autoanalyzer.  $\text{NH}_4^+$  was measured by the phenol-  
235 hypochlorite method (FIAstar method AN5220), nitrate (as  $\text{NO}_3^- + \text{NO}_2^-$ ) was measured by  
236 cadmium reduction (FIAstar method AN5201) and SRP was measured by the  
237 antimony/molybdate and ascorbic acid method (Alpkem method A303-S200-00). TDN was  
238 measured by alkaline persulfate digestion (Eaton et al. 1995). DON was calculated from the  
239 difference between total dissolved nitrogen (TDN) and ( $\text{NH}_4^+ + \text{NO}_3^-$ ). SPOM was estimated  
240 gravimetrically. Alkalinity was determined by two-point titration with  $\text{H}_2\text{SO}_4$  (Hach 16900-01  
241 digital titrator).

242 The  $\delta^{15}\text{N}$  content of organic matter compartments was determined with a Finnigan Delta-  
243 plus mass spectrometer. The  $\delta^{15}\text{N}$  content of  $\text{NH}_4^+$  and  $\text{NO}_3^-$  were determined by diffusion  
244 (Sigman et al. 1997; Holmes et al. 1998). All  $\delta^{15}\text{N}$  values were corrected for background  $^{15}\text{N}$  by  
245 subtracting the average of  $\delta^{15}\text{N}$  value of samples collected prior to the start of the  $^{15}\text{N}$  addition  
246 and from the upstream station (-90 m) during the addition. DON in these streams typically  
247 comprises about one-third to one-half total dissolved N. Because laboratory incubation  
248 experiments indicated that DON material was not reactive in the time scale of about 2 h it took  
249 for streamwater to move through the study reaches, we did not correct for breakdown of DON. In  
250 2 week laboratory incubations, we detected no consistent change in DON concentrations (Krushe  
251 pers. comm.)

### 252 *Calculations*

253 We calculated the mass of  $^{15}\text{N}$  in different compartments based on  $\text{NH}_4^+$  and  $\text{NO}_3^-$   
254 concentrations, discharge, estimates of the biomass of organic matter pools and  $^{15}\text{N}$  of the  
255 compartment. We used the rate of increase and final distribution of the  $^{15}\text{N}$  label in  $\text{NH}_4^+$  and  
256  $\text{NO}_3^-$  to calculate  $^{15}\text{N}$  uptake kinetics (Mulholland et al. 2000). We used the concentration of  $^{15}\text{N}$   
257 and the mass of different organic matter components to estimate the amount of the added  $^{15}\text{N}$   
258 retained in each compartment. Total export in water was the sum over the 21-d experiment of  
259 particulate and dissolved N transported out of the reach determined from water flux and the  
260 concentrations and  $^{15}\text{N}$  content of  $\text{NH}_4^+$ ,  $\text{NO}_3^-$  and SPON.

261 We used a simple mixing model (Phillips et al. 2005) to estimate the range of potential  
262 sources of organic matter (forest leaves, riparian grass, algae) to stream CBOM, FBOM and  
263 SPOM.

#### 264 *Regional river basin nitrogen budgets*

265 We combined an analysis of land cover and stream networks from the regional survey  
266 with the  $^{15}\text{N}$  uptake dynamics to estimate the extent to which deforestation has altered streams  
267 and nitrogen movement at the river-basin scale. We used the extent of streams in different land  
268 cover classes and the information on regional stream infilling by riparian grasses plus the uptake  
269 rates per  $\text{m}^2$  of stream bottom to calculate total N uptake in first- and second-order forest and  
270 pasture streams. We used budgets of the fate of  $^{15}\text{N}$  to calculate N retained in the first- and  
271 second-order stream reaches. The total N uptake and retention were determined for two  
272 scenarios: (1) entire Ji-Paraná basin covered by forest (pre-settlement scenario) and (2) under  
273 current land use (27% deforested in 1999; non-forest land use was 93% pasture).

## 274 **Results**

### 275 Regional survey

276 Geomorphological and ecological changes brought about by forest conversion to pasture  
277 produced widespread changes in physical structure and function of small streams. Across the  
278 region, forest and pasture streams had similar mean basin area, discharge and depth but forest  
279 streams had a narrower wetted width and more benthic cover of leaves (50%) and sand than  
280 pasture streams (Table 1). Pasture streams were wider, with slow moving water and deep  
281 accumulations of fine organic material (19% cover). Usually a small open water channel was  
282 present in pasture streams lined with floating mats of *P. repens* on both sides (63% riparian grass  
283 cover). Forest and pasture streams had similar pH, alkalinity and conductivity but pasture  
284 streams had lower concentrations of  $\text{NO}_3^-$  and dissolved oxygen (often near zero) and higher  
285 total suspended solids than forest streams (Table 1).

286 Landscape scale analysis of images of the Ji-Parana watershed (LANDSAT, videography  
287 and IKONOS) showed that deforestation of 27% of the Ji-Parana watershed altered 7,102 km  
288 (34% of total) first- and second-order channels. Infilling by *P. repens* was easily recognized in  
289 IKONOS and videography images (Fig. 1) and when combined with the on-the-ground survey  
290 and observations, indicated that almost all small pasture streams were dominated by riparian  
291 grass infilling.

292  $^{15}\text{N}$  additions

293  
294 *Benthic habitats.* The distribution of benthic habitats in the second-order forest stream  
295 used for  $^{15}\text{N}$  addition was similar to that of forest streams in the regional survey (Table 2). Leaf  
296 pack (51%), sand (28%) and areas of sand overlain with a thin (a few mm) layer of fine organic  
297 matter (16%) were the dominant habitats. There were no extensive or deep accumulations of fine  
298 or coarse organic matter even in pools. Filamentous macroalgae were rare. The benthic habitat of  
299 the second-order pasture stream used for  $^{15}\text{N}$  addition was also similar to second-order pasture

300 streams in the regional survey. Benthic habitat cover was dominated by *P. repens* (55%) with  
301 water flowing through the root zone (Table 2). Grass detritus, sand and fine organic matter made  
302 up the remainder of benthic habitat. In the third-order pasture stream, riparian grass was found  
303 only on point bars (6%) and much of the stream had a sandy bottom. Benthic habitat was  
304 dominated by sand (42%), leaf and grass detritus (23%) and areas with a thin layer of fine  
305 organic matter over sand (14%). Filamentous macroalgae occurred attached to hard substrate  
306 (which was rare) or riparian grass in both pasture streams but were not common.

307 *Water chemistry.* The chemical characteristics of the second-order forest and pasture  
308 streams were similar to those in the regional survey (Table 3, Fig. 2). All forest and pasture  
309 streams had similar mean pH, alkalinity, conductivity,  $\text{NH}_4^+$  concentrations. The second-order  
310 pasture stream had lower concentrations of  $\text{NO}_3^-$  and dissolved oxygen (range 0.1 to 2.8  $\text{mg L}^{-1}$ )  
311 and higher SRP than the forest stream or the third-order pasture stream. DON was generally one  
312 third to about one half of the total dissolved N (Fig 2).  $\text{NH}_4^+$  and  $\text{NO}_3^-$  increased as discharge  
313 decreased in the forest, but declined in proportion to discharge in the second-order pasture  
314 stream. There was no clear pattern with discharge in the third-order pasture stream. The ratio of  
315 dissolved inorganic N:P was highest in the forest stream (105), lowest in the second-order  
316 pasture stream (2) and intermediate (17) in the third-order pasture stream.

317 *Organic matter and N standing stocks.* The second-order pasture stream had 10 times  
318 more total N  $\text{m}^{-2}$  of stream reach (52.1  $\text{g N m}^{-2}$ ) than either the second-order forest (4.1  $\text{g N m}^{-2}$ )  
319 or the third-order pasture stream (2.7  $\text{g N m}^{-2}$ ) (Table 2). The location of N stocks in stream  
320 channels also differed. In the forest stream, leaf pack habitat had more than 25 times the N stock  
321 than depositional areas of fine organic sediments. In the second-order pasture stream, the large  
322 area of grass cover, high grass standing stock and abundance of CBOM and FBOM in grass

323 habitat led to very high N stocks. There were typically accumulations (10 to 50 cm thick) of dead  
324 organic material in some of the deeper areas of the open stream channel and under the riparian  
325 grasses in the small pasture stream. In the third-order pasture stream, grass cover and grass  
326 standing stock were low and the stream had total N stocks and a distribution of N stocks that  
327 were similar to the forest stream.

328 *Organic matter  $\delta^{13}C$ .* In the forest stream, the  $\delta^{13}C$  of organic matter compartments  
329 ranged from -26.2 ‰ in SPOM to -31.5 ‰ in biofilms (Table 4). The  $\delta^{13}C$  of riparian grass  
330 leaves and roots and rhizomes ranged from -12.7 to -17.6 ‰ (Table 4). All organic matter classes  
331 (CBOM, FBOM and SPOM) in the forest stream had isotopic values consistent with being  
332 derived 100% from inputs of terrestrial C3 vegetation. The  $\delta^{13}C$  of CBOM, FBOM and SPOM in  
333 the pasture streams was enriched compared with the forest stream, suggesting inputs of C4  
334 grasses or algae. In the second-order pasture stream, contributions to CBOM came 90 to 100%  
335 from C4 grass with the remainder derived from algae (10%). For FBOM, contributions were  
336 smaller (70%) from grass and larger (30%) from algae. In the third-order pasture stream,  
337 contributions to CBOM were from grass (50 to 80%), trees (30 to 55%) and algae (10 %) and  
338 contributions to FBOM were from grass (20 to 40%), trees (50 to 60%) and algae (2 to 30%). In  
339 both pasture streams, algae provided a larger contribution to FBOM (~20%) than to CBOM  
340 (~10%). SPOM was derived from similar sources as FBOM in all streams.

341  *$^{15}NH_4^+$  and  $^{15}NO_3^-$ .* In all streams,  $\delta^{15}NH_4^+$  decreased with distance downstream and  
342 indicated that  $NH_4^+$  uptake differed among the streams (Fig. 3). In the forest stream,  $\delta^{15}NH_4^+$   
343 decreased over the course of the experiment because despite decreasing discharge streamwater  
344  $NH_4^+$  concentration increased seven-fold (Fig. 2). This resulted in a relatively constant  $NH_4^+$   
345 during the experiment. In both pasture streams,  $^{15}NH_4^+$  enrichment increased during the

346 experiment because of declining discharge and either declining or stable  $\text{NH}_4^+$  concentrations.  
347 Declining discharge in these streams led to a small decrease in  $\text{NH}_4^+$  flux during the experiment.

348 The second-order pasture stream had a shorter  $\text{NH}_4^+$  uptake length and a shorter  $\text{NH}_4^+$   
349 residence time than either of the other streams (Table 5). The third-order pasture stream had a  
350  $\text{NH}_4^+$  uptake rate, a  $\text{NH}_4^+$  uptake length and a  $\text{NH}_4^+$  residence time similar to the forest stream  
351 (Table 5).  $\text{NH}_4^+$  uptake velocity showed a different pattern and was slightly higher in the forest  
352 stream compared with the second-order pasture stream and lowest in the third-order pasture  
353 stream (Table 5). We saw no significant increase in  $\delta^{15}\text{NH}_4^+$  in the post-addition phase of any of  
354 the addition experiments, indicating that regeneration of  $\text{NH}_4^+$  was minor in all of the streams.  
355 On day 22 (post-addition day 1),  $^{15}\text{NH}_4^+$  returned almost to background levels in all streams and  
356 remained constant for the remainder of the post-addition period (Fig. 3).

357 In the second-order forest stream, nitrification was significant and increased  $\delta^{15}\text{NO}_3^-$   
358 downstream of the  $^{15}\text{N}$  addition site (Fig. 4A). Mean  $\text{NO}_3^-$  uptake lengths in the forest stream  
359 ranged from 125 m to infinity and averaged 717 m (Table 5). Uptake by direct nitrification  
360 accounted for 22% of the whole-stream  $\text{NH}_4^+$  uptake rate (10.1 of 46.6  $\mu\text{gN m}^{-2} \text{min}^{-1}$ ) and was  
361 the most significant removal pathway of  $\text{NH}_4^+$  (Table 5). Very low concentrations of  $\text{NO}_3^-$  (range  
362 0.0 to 0.7  $\mu\text{mol L}^{-1}$ ) in the second-order pasture stream made it impossible to reliably measure  
363 the  $\delta^{15}\text{NO}_3^-$  and limited our ability to estimate the nitrification rate. Low dissolved oxygen (range  
364 0.1 to 2.8  $\text{mg L}^{-1}$ ) combined with the very low  $\text{NO}_3^-$  concentrations implied that nitrification did  
365 not occur in the second-order pasture stream. In contrast, in the third-order pasture stream  $^{15}\text{NO}_3^-$   
366 was absent on day 1, but clearly present by day 3 (Fig. 4B). The lack of  $^{15}\text{NO}_3^-$  on day 1  
367 indicated that direct nitrification ( $^{15}\text{NO}_3^-$  production from added  $^{15}\text{NH}_4^+$ ) was minimal and that  
368 indirect nitrification (the production of  $^{15}\text{NO}_3^-$  from  $^{15}\text{NH}_4^+$  mineralized from  $^{15}\text{N}$ -labeled organic

369 matter) was responsible for formation of  $^{15}\text{NO}_3^-$  on days 3 to 21. Nitrification accounted for none  
370 of the  $^{15}\text{NH}_4^+$  uptake in the second-order pasture stream and less than 1% of  $^{15}\text{NH}_4^+$  uptake in the  
371 third-order pasture stream (Table 6). No  $^{15}\text{NO}_3^-$  uptake was measured in the second- and third-  
372 order pasture streams.

373  *$^{15}\text{N}$  in organic matter.* In all streams, added  $^{15}\text{N}$  accumulated in the organic matter  
374 compartments during the addition and then was lost relatively quickly after the addition ended  
375 (Fig. 5). In the forest stream, algae and biofilm, had the highest enrichments (Fig. 5A) and fastest  
376 post addition declines. In the forest stream,  $^{15}\text{N}$  in FBOM increased sharply during the  
377 experiment and declined quickly post-addition. The forest stream CBOM accumulated and lost  
378  $^{15}\text{N}$  slower than FBOM, returning to background within 5 days (Fig 5A).

379 In both pasture streams, algae were more highly labeled than microbial biofilm, FBOM  
380 or CBOM (Fig. 5B, C). In the second-order pasture stream, FBOM was more enriched than  
381 CBOM after 7 d (Fig. 5B). In contrast, in the third-order pasture stream, the CBOM was more  
382 enriched than FBOM after 7 d (Fig.5C). The  $^{15}\text{N}$  enrichment in all of these compartments was  
383 still increasing at the end of 21 d and tracked the increase in  $\delta^{15}\text{NH}_4^+$  that occurred over the  
384 course of the addition.

385 In both pasture streams, the  $^{15}\text{N}$  in all biomass compartments was highest in the stations  
386 30 to 100 m downstream of the addition point and then declined downstream, while the forest  
387 stream had a much more uniform distribution of  $^{15}\text{N}$  over the reach (Fig. 6). The highest  
388 enrichments did not always occur at the station nearest the  $^{15}\text{N}$  addition point where  $\delta^{15}\text{NH}_4^+$  was  
389 highest. This may have been caused by influx of unlabeled organic matter from upstream of the  
390 addition point in the case of CBOM or FBOM, or by sloughing or loss of the most labeled  
391 portions of algae or biofilm.

392 Riparian grass was highly labeled and both stream channel edge and floodplain grass  
393 became labeled (Fig. 7). In the second-order pasture stream, stream channel edge leaves and  
394 roots and rhizomes became highly labeled 30 to 50 m downstream of the addition point and were  
395 still highly labeled (>50‰) at 500 m (Fig. 7A). The leaves became more highly labeled than the  
396 roots and rhizomes downstream of 100 m, suggesting prompt translocation of acquired N into  
397 newly-produced leaves. Grass in the floodplain also became labeled, with the highest label 120  
398 to 230 m below the addition point (Fig. 7A), indicating that water moving below the grass and  
399 carrying the  $^{15}\text{NH}_4^+$  spread out away from the stream channel downstream of the addition point.  
400 In the third-order pasture stream, the grass roots and rhizomes were more highly enriched (100 to  
401 130‰) than the leaves (10 to 50‰), indicating N uptake and storage in rhizomes (Fig. 7B).

402 The biomass compartment-specific uptake rates of  $\text{NH}_4^+$  were higher in the pasture  
403 streams than in the forest stream. Uptake of  $\text{NH}_4^+$  was 2 times higher into grass leaves (6.4  
404 versus 3.3  $\mu\text{g N m}^{-2} \text{min}^{-1}$ ) and 7 times higher into roots and rhizomes (67 versus 10.8  $\mu\text{g N m}^{-2}$   
405  $\text{min}^{-1}$ ) in the second-order pasture stream compared to the third order pasture stream. Uptake of  
406  $\text{NH}_4^+$  into CBOM in the second-order pasture stream (1.7  $\mu\text{g N m}^{-2} \text{min}^{-1}$ ) was 85 times higher  
407 than in the forest stream (0.02  $\mu\text{g N m}^{-2} \text{min}^{-1}$ ) and 2 times higher (1.0  $\mu\text{g N m}^{-2} \text{min}^{-1}$ ) than the  
408 third-order pasture stream. A similar pattern was seen for rates of uptake into FBOM, which  
409 were 2.7  $\mu\text{g N m}^{-2} \text{min}^{-1}$  in the second-order pasture stream, 0.01  $\mu\text{g N m}^{-2} \text{min}^{-1}$  in the forest  
410 stream and 2.4  $\mu\text{g N m}^{-2} \text{min}^{-1}$  in the third-order pasture stream.  $^{15}\text{N}$  uptake into leaf pack in the  
411 forest stream (0.64  $\mu\text{g N m}^{-2} \text{min}^{-1}$ ) was half the rate of uptake into forest CBOM.

412 Turnover times of N in CBOM and FBOM in the forest streams were 2.5 to 5 times faster  
413 than their turnover in both pasture streams. The turnover time for CBOM was  $5 \pm 1$  d in the  
414 forest stream,  $26 \pm 16$  d in the second-order pasture stream and  $16 \pm 5$  d in the third-order pasture

415 stream. The turnover time for FBOM was  $4 \pm 2$  d in the forest stream,  $15 \pm 5$  d in the second-  
416 order pasture stream and  $10 \pm 3$  d in the third-order pasture stream. The turnover time of N in  
417 forest leaf pack was  $7 \pm$  d. These turnover times assumed no uptake of regenerated N and were  
418 likely underestimates of actual turnover time.

#### 419 Nitrogen mass balance

420 A mass balance accounting of added  $^{15}\text{N}$  indicated a change from predominantly N pass-  
421 through and export in the forest stream to N retention in the second-order pasture stream. In the  
422 forest stream, the predominant fate of N was export as DIN (Table 6). Fifty- seven percent of  
423  $^{15}\text{NH}_4^+$  added was exported as  $^{15}\text{NH}_4^+$  and 8% was exported as  $\text{NO}_3^-$ . Seven percent of the added  
424  $\text{NH}_4^+$  entered organic matter pools and export of N as suspended particulate organic matter was  
425 minor. Forty percent of  $^{15}\text{N}$  added to the forest stream was not accounted for at the end of the  
426 addition.

427 In contrast to the forest, in the second-order pasture stream almost 75% of added  $^{15}\text{N}$  was  
428 retained in biomass compartments, predominately riparian grass. Export accounted for 11% of  
429 added  $^{15}\text{N}$ , either as  $^{15}\text{NH}_4^+$  or in particulate form, and no  $^{15}\text{N}$  was exported as  $^{15}\text{NO}_3^-$ . Most of  
430 the total  $^{15}\text{N}$  recovered in biomass compartments in the second-order pasture stream was retained  
431 by riparian grass. Fourteen percent of  $^{15}\text{N}$  added in the second-order pasture stream was not  
432 accounted for.

433 The pattern of export and retention in the third-order pasture stream differed from that in  
434 the second-order pasture stream, with only 5% of added N retained and 26% exported as  $\text{NH}_4^+$   
435 (9%),  $\text{NO}_3^-$  (6%) and particulate N (11%). Two percent or less of the added  $^{15}\text{N}$  was stored as  
436 CBOM, FBOM or riparian grass. Seventy percent of added  $^{15}\text{N}$  was not accounted for.

#### 437 Scaling to regional river basin N budgets

438 We estimate based on length of stream channel altered based on the remote sensing and  
439 changes to N cycling based on the  $^{15}\text{N}$  additions that current deforestation of the Ji-Paraná basin  
440 of 27% has resulted in a 7-fold increase in basin-wide  $\text{NH}_4^+$  uptake (20, 279  $\text{kg N d}^{-1}$  under  
441 current land use versus 2,817  $\text{kg N d}^{-1}$  in the fully forested watershed) and a 16-fold increase in  
442  $\text{NH}_4^+$  retention compared to the watershed in the fully forested state (15, 151  $\text{kgN d}^{-1}$  under  
443 current land use versus 929  $\text{kg N d}^{-1}$  in the fully forested watershed). The 4-fold greater uptake  
444 rates of ammonium, lowered nitrification and high retention in riparian grass in small pasture  
445 streams resulted in retention of more than 14,000  $\text{kg of N d}^{-1}$  that would have been delivered  
446 downstream to third-order streams, and then flowed further downstream to larger rivers, had  
447 deforestation not occurred.

## 448 **Discussion**

### 449 Stream structure and water chemistry

450 Forest conversion to pasture in the watershed of second-order streams transformed stream  
451 morphology by eliminating the pool and run structure of the forest and replacing it with a narrow  
452 run of open water and a wide, wet, marshy bordering area created by extensive infilling with  
453 riparian grass in the stream and adjacent pasture. This led to high biomass of grass, CBOM and  
454 FBOM, hypoxic conditions, lower concentrations of  $\text{NO}_3^-$  and higher concentration of SRP in the  
455 pasture stream. These characteristics are linked to slower water velocities and greater transient  
456 storage of water and higher inputs of organic matter in pasture streams (Neill et al. 2006). Low  
457 dissolved oxygen has been associated with low concentrations of  $\text{NO}_3^-$ , low DIN:DIP and  
458 inducement of N limitation of pasture stream periphyton (Neill et al. 2001). The high ratio of  
459 dissolved inorganic N:P of 105 in the forest stream and 2.4 in the second-order pasture stream  
460 also indicated P limitation of algal and bacterial growth in the forest stream but N limitation in

461 the pasture stream (Redfield 1958; Elser et al. 2007). The source of CBOM and FBOM in the  
462 second-order pasture stream also shifted to C4 grass sources from the C3 vegetation in the forest.  
463 Because of the large biomass and proximity of the streamside C4 grass *P. repens* it was the most  
464 likely source of C4-derived CBOM and FBOM in the second-order pasture stream.

465         The extensive infilling of first- and second-order streams by riparian grasses was  
466 widespread based on our ground survey and image analysis, and led to similar physical and  
467 chemical conditions across a wide range of locations. While the stabilization of banks by grasses  
468 and narrowing of stream channels occurs in meadow and pasture streams in different  
469 environments (Trimble 1997; Hession et al. 2003; Sweeney et al. 2004), the development of  
470 wide, marshy flooded streambanks in pasture represents a structural change not reported for  
471 other areas of the world. The condition in these altered streams were similar to large Amazon  
472 floodplain lakes, where dense mats of floating aquatic vegetation are widespread and high  
473 organic matter inputs lead to hypoxic conditions (Junk 1973; Wellcome 1985). Similar pasture  
474 stream infilling by grasses occurs elsewhere in deforested regions of the lowland Amazon, such  
475 as in eastern Pará state (R. Figueiredo, personal comm.), but is poorly documented.

476         In contrast to the second-order pasture stream, forest conversion to pasture in the  
477 watershed of the third-order stream resulted in less infilling of the channel by grass, lower  
478 streamside grass biomass and associated lower biomass of CBOM and FBOM. Third-order  
479 pasture streams also did not have the hypoxic conditions, low  $\text{NO}_3^-$  and higher concentrations of  
480 SRP found in the second-order pasture stream. In these ways, both physical and chemical  
481 conditions of the third-order stream more closely resembled conditions in the forest stream. The  
482 extent of forest clearing to the stream edge in the third-order pasture stream was also less than in  
483 the small streams in the regional survey. This may be related to the difficulty of clearing trees

484 from the larger and wetter floodplains of larger streams. There was still a substantial shift in the  
485 origin of CBOM and FBOM from forest vegetation to C4 grasses in the third-order pasture  
486 stream. We could not distinguish between potential C4 sources in *P. repens* near the stream  
487 channel or the *B. brizantha* that occupied most of the upland portion of the watershed and some  
488 fraction of CBOM and FBOM may reflect the upland grass source.

489         These finding suggests that many of the most important structural and chemical changes  
490 to streams following deforestation depend on stream scale and that the largest changes occur at  
491 the smallest stream orders. The differences in channel structure between the second- and third-  
492 order pasture streams are potentially explained by an increase in erosive power per unit of  
493 streambed area with increasing discharge (Bagnold 1966). This higher erosive power may  
494 contribute to the lack of grass infilling if stream edge riparian vegetation provides little effective  
495 bank protection during channel-forming flows (Davies-Colley 1997). While we found virtually  
496 no  $\text{NO}_3^-$  in the second-order pasture streams, in a survey of mostly larger streams and rivers in  
497 Rondônia Biggs et al. (2004) found a positive relationship between  $\text{NO}_3^-$  concentration and  
498 percentage of the watershed in pasture, indicating a change in this relationship with stream size.  
499 Land-use practices may also play a role in the scale-dependence of stream structural changes.  
500 Ranchers typically clear trees from land immediately adjacent to small perennial streams to  
501 provide cattle with access to water and encourage grass infilling of stream courses to provide a  
502 source of actively-growing forage during the dry season, but clearing the wider riparian forest  
503 adjacent to larger streams is more difficult and sometimes not done. By these practices, ranchers  
504 may be encouraging the differences we measured.

505         We found that the change in land use from forest to pasture surrounding second-order  
506 streams led to a dramatic shift in the sources of both coarse, fine and suspended organic matter

507 from forest-derived leaves to riparian grass and that this shift took place rapidly (within a few  
508 years) following land-use conversion. This was consistent with the more enriched  $\delta^{13}\text{C}$  values of  
509 POC in tributaries of the Ji-Paraná River in Rondônia that had higher proportions of pasture in  
510 their watersheds (Bernardes et al. 2004). The  $\delta^{13}\text{C}$  values we found in second-order pasture  
511 streams were much more enriched in  $^{13}\text{C}$  and had a higher C4 grass-derived fraction than those in  
512 larger rivers (> third-order), most likely because larger rivers still contain substantial fractions of  
513 forest in their watersheds and because riparian and floodplain forests can disproportionately  
514 deliver fresh particulate organic C to larger rivers (Mayorga et al. 2005).

#### 515 Forest stream $\text{NH}_4^+$ uptake and transformations

516 In the forest stream, uptake into CBOM, FBOM and POM was low,  $\text{NH}_4^+$  uptake velocity  
517 was low, the uptake lengths of both  $\text{NH}_4^+$  and  $\text{NO}_3^-$  were long and 53% of  $^{15}\text{N}$  added was  
518 exported from the reach as dissolved inorganic N. The fate of the 40% of added  $^{15}\text{N}$  that was not  
519 recovered was not clear. While denitrification was possible in anoxic locations in the channel  
520 such as the hyporheic zone, we did not observe the required  $\text{NO}_3^-$  uptake. We did not measure  
521  $\text{DO}^{15}\text{N}$  export, but production of  $\text{DO}^{15}\text{N}$  presumably would have been low given that we did not  
522 have strong uptake of  $^{15}\text{N}$  into any abundant rapid turnover compartments, such as algae (rare in  
523 forest). Fluvial fluxes were well quantified and there were no major discharge events or  
524 measurable loss of water to hyporheic flows that might have resulted in large amounts of  
525 unquantified  $^{15}\text{N}$  export.

526 The forest stream had lower  $\text{NH}_4^+$  uptake velocity and a roughly 10-fold longer uptake  
527 length than forested temperate and tropical North American streams with similar discharge and  
528  $\text{NH}_4^+$  concentrations (Mulholland et al. 2000; Peterson et al. 2001; Webster et al. 2003). The  
529 North American streams with the most similar uptake lengths (475 m to 1350 m) were in

530 agricultural watersheds very high N concentrations, suggesting that  $\text{NH}_4$  was not tightly cycled  
531 or limiting in these agricultural streams (Hamilton et al. 2001; Webster et al. 2003). The  
532 nitrification rate in our forest stream was comparable to temperate North American streams  
533 (Webster et al. 2003), but substantially lower than the rate (50% to 60%) for a tropical forest  
534 stream in Puerto Rico (Merriam et al. 2002). Among temperate streams, no consistent control of  
535 nitrification has been identified. Higher temperatures have been shown to increase nitrification  
536 (Warwick 1986), suggesting that our streams should have had higher rates of nitrification than  
537 the temperate streams and rates more comparable to the tropical Puerto Rican forest stream.  
538 Nitrification generally occurs in sediment and biofilms (Stream Solute Workshop, 1990) and  
539 requires oxygen and  $\text{NH}_4^+$ . Tank et al. (2000) found almost no nitrification in a North American  
540 forest stream and attributed the low nitrification rate to low  $\text{NH}_4^+$  concentrations and poor stream  
541 sediment quality for colonization by nitrifying bacteria. In contrast, in other temperate streams  
542 with low  $\text{NH}_4^+$  concentrations, 10% to 50% of the total  $\text{NH}_4^+$  uptake from streamwater was  
543 attributed to direct nitrification (Dodds et al. 2000; Mulholland et al. 2000; Ashkenas et al.  
544 2004), similar to our forest stream. The export of inorganic N in the second-order forest (57%  
545  $\text{NH}_4^+$  and 8%  $\text{NO}_3^-$ ) was on the high end of export as  $\text{NH}_4^+$  (1 to 65%) and the low end of export  
546 as  $\text{NO}_3^-$  (1% to 50%) and export of SPON (1% to 12%) reported for temperate streams (Hall et  
547 al. 1998; Dodds et al. 2000; Mulholland et al. 2000; Tank et al. 2000; Hamilton et al. 2001;  
548 Ashkenas et al. 2004).

549 Long uptake lengths, moderate concentrations of DIN and high N:P all suggested algal  
550 and bacterial production in the forest stream was not limited by nutritional demand for N. Uptake  
551 of  $^{15}\text{NH}_4$  by CBOM, FBOM and leaf biomass was slower than  $^{15}\text{NH}_4$  uptake by nitrification.  
552 Combined, the biomass compartments accounted for only 1% of the whole stream  $\text{NH}_4$  uptake

553 and 6% of the total  $^{15}\text{NH}_4$  uptake. The low  $^{15}\text{NH}_4$  uptake rates by biomass compartments coupled  
554 with the high nitrification rates indicated low demand for N as a nutrient but use of  $\text{NH}_4^+$  as an  
555 energy source by nitrifying bacteria in the stream channel. Low rates of  $\text{NH}_4^+$  uptake were  
556 consistent with shaded conditions that limit algal productivity in forested lowland Amazon  
557 headwater streams (Fittkau 1967; Lowe-McConnell 1987).  $\text{NH}_4^+$  uptake but no  $\text{NO}_3^-$  uptake was  
558 also consistent with a general preference for  $\text{NH}_4^+$  assimilation by stream biota (Reynolds 1984;  
559 Fenchel et al. 1998) and by phytoplankton in Amazon River floodplain lakes (Fisher et al. 1988).  
560 Assimilatory uptake of  $\text{NO}_3^-$  has been found to contribute total inorganic N uptake in some  
561 temperate urban and agricultural streams (Arango and Tank 2008, Arango et al. 2008) but that  
562 was not an important process in any of our streams.

563         Several characteristics of mature lowland Amazon tropical forest suggest an open N cycle  
564 and the absence of strong N limitation of the forest ecosystem. These include high concentrations  
565 of N in leaves and litter, high concentrations of  $\text{NO}_3^-$  in forests soil solution and high ratio of  
566  $\text{NO}_3^-:\text{NH}_4^+$ , (Markewitz et al. 2004; Neill et al. 2006; Chaves et al. 2009), high soil emissions of  
567  $\text{N}_2\text{O}$  (Vitousek 1984; Davidson et al. 2007), and high rates of soil nitrification (Neill et al. 1997;  
568 Verchot et al. 1999). The result of the forest stream  $^{15}\text{N}$  addition indicated that the relatively  
569 open N cycle of tropical forests extends to small forest stream channels and further contributes to  
570 open N cycling and the absence of strong N retention at watershed and landscape scales in  
571 forested regions.

572 Pasture stream  $\text{NH}_4^+$  uptake and transformations

573         The second-order pasture stream had a much higher uptake of  $\text{NH}_4^+$  than the forest  
574 stream. This was evident in the shorter  $\text{NH}_4^+$  uptake lengths, faster rates of uptake into organic  
575 matter compartments, a shorter  $\text{NH}_4^+$  residence time and very low export of  $^{15}\text{NH}_4^+$ . The most

576 important reason for the higher  $\text{NH}_4^+$  uptake was the presence of riparian grasses in the stream  
577 channel. This grass accounted for 75% of recovered  $^{15}\text{N}$ . A slightly lower  $^{15}\text{NH}_4^+$  uptake velocity  
578 in the second-order pasture stream compared with the forest stream was unexpected and not  
579 easily explained given the high measured total uptake into riparian grass.

580         Low dissolved oxygen concentration inhibits nitrification and promotes denitrification of  
581  $\text{NO}_3^-$  (Seitzinger 1988; Christensen et al. 1990; Kemp and Dodds 2001). The hypoxic conditions  
582 in the second-order pasture stream channel limited nitrification and  $\text{NO}_3^-$  export was zero. The  
583 very high biomass of grass, CBOM and FBOM led to more export as PON (7%) than in the  
584 forest stream (0.1%). Floodplain grass, even 2 m away from the stream channel, was a  
585 significant sink for added  $^{15}\text{NH}_4^+$ . Increases in the amount of  $^{15}\text{N}$  detected in floodplain grass  
586 with distance downstream indicated that water spread away from the open stream channel and  
587 moved underneath the grass mat. This allowed a much larger area of grass to interact with  
588 streamwater and increase the total amount of N retained in the stream channel-floodplain system.  
589 Relatively little (14%) of  $^{15}\text{N}$  added to the second-order pasture stream was not accounted for.  
590 Because of the large grass biomass, variability in quantifying grass biomass and  $\delta^{15}\text{N}$  of the grass  
591 were the most likely sources of this discrepancy.

592         In the third-order pasture stream, most measures of N biogeochemistry were intermediate  
593 between the second-order forest and pasture streams and N export dominated over N retention.  
594 The third-order pasture stream had low uptake of  $^{15}\text{N}$  in CBOM, FBOM and streamside grasses  
595 and some  $\text{NH}_4^+$  export (9%). Some nitrification occurred in the well-oxygenated streamwater but  
596  $\text{NO}_3^-$  export was low (6%). Export of PON (11%) was also relatively high. The most puzzling  
597 aspect of the third-order addition was the low recovery (30%) of the added  $^{15}\text{N}$ . There were no  
598 major discharge events that were not measured or detectable loss of water to hyporheic flows

599 that might have resulted in large amounts of unquantified  $^{15}\text{N}$  export. Errors associated with  
600 uptake into organic compartments were relatively low as total organic biomass was relatively  
601 low. It is tempting to invoke denitrification, however, nitrification accounted for less than 1% of  
602  $^{15}\text{NH}_4^+$  uptake and no  $^{15}\text{NO}_3^-$  uptake was measured in the third-order pasture stream.

### 603 Landscape-scale implications for N retention

604 Changes in stream structure and biogeochemistry of N of small streams suggest that the  
605 connection to larger rivers via downstream transport of N typical of forested systems is  
606 weakened substantially by deforestation. Small forest streams function largely as transformers of  
607  $\text{NH}_4^+$  to  $\text{NO}_3^-$  and long-distance conduits for transport of inorganic N. Deforestation around  
608 small streams results in infilling of the stream and floodplain by grass and ultimately causes  
609 retention of large amount of N in grass biomass and preventing downstream N transport, at least  
610 over the time scales (weeks to months) measured by the  $^{15}\text{N}$  addition experiment. At the regional  
611 watershed level, changes to N cycling from clearing 25% of the watershed caused a 16-fold  
612 increase in N retention in first- and second-order pasture streams compared with forest streams.  
613 Larger (third-order) pasture streams were not as retentive of N as second-order pasture streams,  
614 but N retention as a percentage of recovered N is higher than small forest streams, suggesting  
615 deforestation for pasture creates a break in transport of  $\text{NO}_3^-$  and diminishes the flow of total N  
616 down small stream corridors to larger rivers.

617 Significant nitrification in streams indicated that  $\text{NO}_3^-$  forest and pasture streams was  
618 derived from nitrification of  $\text{NH}_4^+$  *in situ* rather than direct transfer from  $\text{NO}_3^-$ -rich soil solution  
619 to streams via groundwater pathways. This is consistent with the Chaves et al. (2009), who found  
620 removal of  $\text{NO}_3^-$  in deep pasture soils in Rondônia and low concentrations of  $\text{NO}_3^-$  in  
621 groundwater. It is also consistent with the finding of Brandes et al. (1996) who used natural

622 abundance of  $^{15}\text{NO}_3^-$  to infer that  $\text{NO}_3^-$  in the streamwater of a small primary forest watershed  
623 near Manaus originated from in-channel nitrification. Our finding of low  $\text{NO}_3^-$  concentrations in  
624 small pasture streams combined with observed low production of  $\text{NO}_3^-$  in soils (Neill et al. 1997)  
625 and low  $\text{NO}_3^-$  concentrations in all potential hydrologic flowpaths from pasture (Chaves et al.  
626 2009) indicate that  $\text{NO}_3^-$  in larger pasture streams is also likely derived from nitrification in the  
627 stream channel rather than  $\text{NO}_3^-$  inputs from the watershed. The second-order pasture stream had  
628 no measureable levels of  $\text{NO}_3^-$ .

629 High N retention in small pasture streams is very different than the result from the  
630 temperate zone in which streams running through pasture can have lower rates of N uptake than  
631 streams in forest because narrower streams in pasture leads to less total area of stream bottom per  
632 unit stream length (Sweeney et al. 2004). There is some evidence that riparian vegetation along  
633 temperate forest streams may play a role in stream channel N cycling (Ashkenas et al. 2004), but  
634 the role of riparian grasses on stream N processing is not well studied.

635 These results contradict the trend in temperate latitudes of generally higher N export from  
636 agricultural compared with forested watersheds (Peterson et al. 2001; Allan 2004; Green et al.  
637 2004) and the suggestion of Matson et al. (1999) that increased N additions to tropical terrestrial  
638 systems will result in large losses of N to the ocean in tropical locations because of the absence  
639 of strong N limitation to terrestrial vegetation. Higher export from temperate agricultural areas  
640 results from a combination of changes in stream structure and N processing to favor export and  
641 runoff of N added as fertilizer. Our agricultural pastures are not fertilized, however, the changes  
642 in N cycling favor retention of any N that reaches the streams from land. Our work suggests a  
643 more nuanced view than proposed by Matson, which implied that streams would simply pass N  
644 received from forests downstream unchanged. Stream N cycling changes in response to land-use

645 resulting in high N retention in deforested portions of stream networks (primarily through grass  
646 infilling of smaller streams) and less N retention where deforestation does not result in the same  
647 degree of changes to stream structure and N cycling.

648         There are several potential limitations to our interpretation of larger-scale changes within  
649 stream networks. We did not examine larger forest streams (third-order or higher) because forest  
650 clearing has severely reduced the extent of large forested watersheds in central Rondônia. We  
651 performed our <sup>15</sup>N additions in the dry season, which represents only half of the year (May to  
652 October) and on a limited number of streams. Further work to examine net N retention for long  
653 periods and during the wet season is needed to understand the role of stream channels in  
654 landscape N retention over a complete annual cycle. In the wet season, it is possible that N  
655 transport in both forest and pasture streams will increase because of more frequent and higher  
656 discharge events with faster current velocities. The effects of increased wet season discharge on  
657 downstream flux may be moderated in pasture streams because the grass in stream channels  
658 remains vigorous year-round, is not scoured out annually, and serves to slow water velocities and  
659 filter out particulates, all suggesting the pattern of high retention in the grass is likely to remain  
660 in the wet season. It is also possible that the higher current velocities associated with wet season  
661 floods might dislodge particulates trapped in the riparian vegetation and increase episodic  
662 particulate transport.

### 663 **Conclusions**

664         Conversion of land use change from forest to pasture resulted in a cascade of effects that  
665 altered stream structure and N biogeochemistry. Deforestation simplified second-order stream  
666 structure by promoting channel infilling by streamside grasses and eliminating tree leaf detritus,  
667 creating streams with continuous slow moving deep runs and thick deposits of coarse and fine

668 benthic organic matter. The infilling of small pasture streams with grass created high organic  
669 matter loading, hypoxia and a 16-fold increase in N retention. The same high N retention did not  
670 occur in larger pasture streams where grass infilling was much less. Our findings of widespread  
671 infilling of first- and second-order pasture stream channels by riparian grass and the  
672 accompanying of this phenomenon with lower dissolved oxygen and increased N uptake and  
673 retention suggest that deforestation now alters stream ecosystem structure and N retention over  
674 many thousands of km of small Amazon stream channels. These structural changes and high N  
675 retention are largely restricted to second-order pasture streams. Better estimates of the extent and  
676 spatial scales at which small stream infilling occurs and a better understanding of where N gets  
677 delivered to streams will further improve our understanding and prediction of how N is  
678 transported within Amazon stream networks.

679         These results have several potential implications for the ecological function and  
680 conservation management of the lowland Amazon's streams. Small streams constitute three-  
681 fourths of stream channel length in the Amazon (Junk 1997; Goulding et al. 2003; McClain and  
682 Elsenbeer 2001). Small streams are vital to the life cycles of many fish important in commercial  
683 and subsistence fisheries and they contain the world's richest diversity of freshwater fishes and  
684 aquatic organisms, many of them endemic (Goulding 1980; Barthem 2004; Agostinho et al.  
685 2005). Low dissolved oxygen and greatly altered habitat structure in second-order pasture  
686 streams are likely to have implications for a variety of stream organisms, including fishes.  
687 Because grass infilling in small streams is related to clearing of stream-edge forest, conserving  
688 streamside forest could reduce to grass infilling and preserve stream physical structure and N  
689 patterns of N uptake and retention more like those of the original forest.

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896  
897

898 **Table 1**

899 Mean physical and chemical characteristics and benthic habitat cover for first- and second-  
 900 order forest (n=6) and pasture (n=12) streams surveyed in central Rondônia ( $\pm 1$  se). One-way  
 901 analysis of variance (SAS Institute 2002) on percent cover was performed after arcsin-square  
 902 root transformation; other environmental variables were tested on non-transformed data.

	Forest	Pasture	F-value	P-value
<b>Physical characteristics</b>				
Basin area (km <sup>2</sup> )	3.6 $\pm$ 2.8	2.4 $\pm$ 0.9	0.27	0.612
Forest cover (%)	93 $\pm$ 4	7 $\pm$ 4	163.2	<0.0001
Discharge (L s <sup>-1</sup> )	22 $\pm$ 14	7 $\pm$ 3	0.15	0.705
Mean depth (cm)	9 $\pm$ 3	15 $\pm$ 3	3.11	0.097
Mean total wetted width (m)	2.1 $\pm$ 0.5	10.2 $\pm$ 3.2	6.65	0.020
<b>Benthic cover characteristics</b>				
Grass (%)	0 $\pm$ 0	63 $\pm$ 9	20.14	0.0004
Leaf pack (%)	50 $\pm$ 4	1 $\pm$ 1	194.03	<0.0001
Sand (%)	19 $\pm$ 5	2 $\pm$ 1	14.99	0.0014
Fine organic (%)	22 $\pm$ 5	19 $\pm$ 5	0.07	0.800
Wood (%)	5 $\pm$ 1	1 $\pm$ 0	14.29	0.0016
Other (%)	4 $\pm$ 3	13 $\pm$ 3	3.64	0.075
<b>Chemical characteristics</b>				
pH	6.3 $\pm$ 0.2	6.1 $\pm$ 0.1	1.76	0.201
Alkalinity (mg L <sup>-1</sup> )	15 $\pm$ 3	24 $\pm$ 3	3.63	0.075

Conductivity ( $\mu\text{S cm}^{-1}$ )	$34 \pm 5$	$56 \pm 8$	3.36	0.085
$\text{NH}_4^+$ ( $\mu\text{mol L}^{-1}$ )	$0.9 \pm 0.7$	$4.4 \pm 1.9$	1.06	0.320
$\text{NO}_3^-$ ( $\mu\text{mol L}^{-1}$ )	$6.1 \pm 1.7$	$0.7 \pm 0.3$	32.16	<0.0001
Total suspended solids ( $\text{mg L}^{-1}$ )	$6.9 \pm 1.2$	$26.3 \pm 9.5$	4.49	0.0500
Dissolved oxygen ( $\text{mg L}^{-1}$ )	$5.6 \pm 0.9$	$2.7 \pm 0.7$	6.74	0.0016

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904 **Table 2**

905 Benthic habitat cover and N content of organic matter components in the streams receiving  $^{15}\text{N}$   
 906 addition. Nitrogen content and biomass of wood was not measured.

Stream and habitat	Cover (%)	N content ( $\text{g N m}^{-2}$ of stream channel)			
Second-order forest					
Open channel		CBOM	FBOM	Leaf or grass	Total
Fine organic	16	0	0.003	0.134	0.137
Sand	28	0	0.001	0.000	0.001
Leaf pack	51	0.087	0.049	3.815	3.951
Wood	5				
Total	100	0.087	0.053	3.949	4.089
Second-order pasture					
Open channel		0.05	0.07		0.12
Fine organic	12				
Sand	12		0.01		0.01
Detritus (grass)	17	0.19	0.10		0.29
Wood	5				
Grass channel habitat	55	12.74	16.00		28.74
Leaves				2.73	2.73
Roots and rhizomes				20.20	20.20
Total	100	12.98	16.18	22.93	52.09

Third-order pasture

## Open channel

Fine organic	14		0.06		0.06
Sand	42		0.33		0.33
Detritus (grass, leaves)	23	0.05	0.36		0.41
Wood	5				
Gravel	3				
Clay	7				
Grass channel habitat	6				
Leaves				0.06	0.06
Roots and rhizomes		0.15	0.97	0.40	1.52
Total	100	0.20	1.73	0.46	2.38

---

908 **Table 3**

909 Chemical characteristics (mean and range) of the second-order forest stream and the second- and  
 910 third-order pasture streams during the 21-d  $^{15}\text{N}$  additions.

	Second order		Third order
	Forest	Pasture	Pasture
pH	6.6 (5.2-7.4)	6.1 (4.1-6.5)	6.1 (6.8-9.7)
Alkalinity ( $\text{mg L}^{-1}$ )	18.1 (14.5-22.0)	22.6 (18.0-32.2)	17.6 (15.9-20.9)
Dissolved oxygen ( $\text{mg L}^{-1}$ )	6.7 (4.9-7.7)	1.4 (0.1-2.8)	7.5 (6.2-7.8)
Conductivity ( $\text{mS cm}^{-1}$ )	0.053 (0.044-0.068)	0.1 (0.1-0.2)	0.1 (0.0-0.1)
$\text{NH}_4^+$ ( $\mu\text{mol L}^{-1}$ )	7.3 (1.5-13.4)	3.2 (0.8-12.1)	1.5 (0.3-10.2)
$\text{NO}_3^-$ ( $\mu\text{mol L}^{-1}$ )	7.5 (1.7-15.1)	0.1 (0.0-0.7)	9.1 (3.6-12.1)
Dissolved organic N ( $\mu\text{mol L}^{-1}$ )	12.2 (5.1-16.2)	9.5 (1.4-25.8)	8.8 (1.2-20.7)
SRP ( $\mu\text{mol L}^{-1}$ )	0.2 (0.1-0.3)	2.1 (0.5-4.2)	1.0 (0.2-5.1)
DIN:DIP	105 (16-503)	2.4 (0.7-10.7)	17.4 (2.3-40.5)

911

912

913 **Table 4**

914 Natural abundance of  $\delta^{13}\text{C}$  in stream organic matter compartments. Isotope values were from  
 915 samples taken along the entire reach before the  $^{15}\text{N}$  addition. Values are  $\pm 1$  sd (sample size in  
 916 parentheses).

	$\delta^{13}\text{C}$ (‰)		
	Second-order forest	Second-order pasture	Third-order pasture
Leaf pack	-31.0 $\pm$ 0.8 (13)		
Biofilm	-31.5 $\pm$ 1.8 (19)	-21.4 $\pm$ 1.4 (15)	-26.4 $\pm$ 0.9 (15)
Algae	-28.3 $\pm$ 3.5 (26)	-21.3 $\pm$ 0.9 (8)	-19.4 $\pm$ 2.2 (5)
Riparian grass			
leaves		-12.7 $\pm$ 0.4 (15)	-12.9 $\pm$ 0.5 (6)
rhizomes		-13.6 $\pm$ 2.1 (16)	-17.6 $\pm$ 4.4 (19)
CBOM	-30.4 $\pm$ 0.9 (39)	-17.9 $\pm$ 4.2 (25)	-20.2 $\pm$ 4.7 (19)
FBOM	-29.2 $\pm$ 1.3 (25)	-19.8 $\pm$ 1.4 (22)	-22.4 $\pm$ 1.3 (20)
SPOM	-26.2 $\pm$ 1.3 (25)	-20.2 $\pm$ 0.9 (16)	-24.5 $\pm$ 0.5 (9)

917

918 **Table 5**

919 Whole stream nitrogen processes during the 21-day  $^{15}\text{N}$  addition. nd = not determine because of very low  $\text{NO}_3^-$  concentrations. There  
 920 was no measureable uptake of  $\text{NO}_3^-$  in the third-order pasture stream.

Attribute	Parameter	Units	Second order forest	Second order pasture	Third order pasture
$\text{NH}_4^+$ dynamics	Uptake rate	$\mu\text{g N m}^{-2} \text{min}^{-1}$	$46 \pm 4$	$64 \pm 65$	$49 \pm 18$
	Nitrification	$\mu\text{g N m}^{-2} \text{min}^{-1}$	10.1	0	$5.1 \times 10^{-5}$
	Uptake velocity ( $V_f$ )	$\text{mm s}^{-1}$	$0.089 \pm 0.015$	$0.060 \pm 0.014$	$0.018 \pm 0.011$
	Uptake length	m	$1623 \pm 327$	$410 \pm 127$	$1264 \pm 330$
	Residence time	min	$369 \pm 75$	$168 \pm 52$	$310 \pm 81$
$\text{NO}_3^-$ dynamics	Uptake length	m	$717 \pm 824$	nd	no uptake
	Nitrification ( $k_n$ )	$\text{m}^{-1}$	0.028	nd	0.00003
	$A_o \times K_n$		$0.130 \pm 0.051$	nd	$0.029 \pm 0.062$

921

922 **Table 6**923 Mass balance of  $^{15}\text{N}$  added to stream reaches.

	Second-order				Third-order	
	Forest		Pasture		Pasture	
	$^{15}\text{N}$	% of	$^{15}\text{N}$	% of	$^{15}\text{N}$	% of
	(g)	added	(g)	added	(g)	added
Total $^{15}\text{N}$ added	5.90		21.50		18.90	
Retained						
CBOM	0.01	0.2	0.4	2.0	0.1	0.5
FBOM	0.01	0.1	0.8	3.7	0.4	2.2
Leaf pack	0.42	7.1				
Grass - channel edge			11.2	52.2	0.3	1.8
Grass - floodplain			3.7	17.4		
Total retained	0.43	7.4	16.2	75.3	0.8	4.5
Exported in stream water						
$\text{NH}_4^+$	1.98	33.6	0.7	3.7	1.8	9.3
$\text{NO}_3^-$	1.13	19.2	0.0	0.0	1.0	5.6
SPON	0.01	0.1	1.6	7.4	2.1	11.0
Total exported	3.12	52.9	2.3	11.1	4.9	25.9
Total $^{15}\text{N}$ recovered	3.55	60.3	18.6	86.2	5.7	30.4

924

925

926 **Figure Legends**

927 **Figure 1.** Land use in the Ji-Parana River Basin and images of Rondônia forest and pasture  
928 streams and streams used for the  $^{15}\text{N}$  additions. The Ji-Paraná map (middle) is derived from a  
929 Landsat ETM composite from 1999. Detailed images (side panels) show the widespread infilling  
930 of stream channels with riparian grasses (primarily *Paspalum repens*) determined from Landsat  
931 (left side: A), IKONOS (left side: B, C, and D) and videography (right side: E, F, G, H). Panel A  
932 is a Landsat image of Nova Vida Ranch near the city of Ariquemes. The white circles indicate  
933 the second-order forest and pasture streams, while the third-order pasture stream runs along the  
934 forest boarder near the bottom left of the image. The IKONOS images show representative  
935 channel and riparian area infilling by grasses (lime green coloration), the extent of pasture (pink)  
936 and forest (dark green with texture). *Paspalum* in streams and adjacent floodplains are evident in  
937 the fine scale videography (right panel).

938

939 **Figure 2**

940 Environmental conditions during the course of the  $^{15}\text{N}$  addition experiments. Plots are mean  
941 concentrations of  $\text{NH}_4^+$  (closed circles),  $\text{NO}_3^-$  (open circles), SRP (open triangles), DON (open  
942 squares) and discharge (solid line) during the  $^{15}\text{N}$  addition experiment in the second-order forest  
943 stream (A), the second-order pasture stream (B) and the third-order pasture stream (C). Solute  
944 concentrations represent averages ( $\pm 1$  sd) of 3 to 7 stations over the experimental reach, except  
945 for soluble reactive phosphate in the forest stream, which was measured at only one station at the  
946 downstream end of the reach on each date. Shaded area represents the period of  $^{15}\text{NH}_4^+$  addition.

947

948 **Figure 3**

949 The  $\delta^{15}\text{N}$  of  $\text{NH}_4^+$  in the second-order forest stream (A), second-order pasture stream (B) and  
950 third-order pasture stream (C). Profiles are  $\delta^{15}\text{NH}_4^+$  in stream water over the study reach on one  
951 date prior to the start, 5 dates during and 4 dates after the  $^{15}\text{N}$  addition ended. All values were  
952 normalized to upstream  $\delta^{15}\text{NH}_4^+$  values on each date.

953

954 **Figure 4**

955 The  $\delta^{15}\text{N}$  of  $\text{NO}_3^-$  in the second-order forest stream (A) and the third-order pasture stream (B).  
956 Profiles are  $\delta^{15}\text{NO}_3^-$  in stream water over the study reach on one date prior to the start, 5 dates  
957 during and 4 dates after the  $^{15}\text{N}$  addition ended. The forest stream had the highest rate of  
958 nitrification as reflected in the increase in  $\delta^{15}\text{NO}_3^-$ . Nitrate concentrations in the second-order  
959 pasture stream were too low to reliably measure  $^{15}\text{NO}_3^-$ . Closed symbols were during the  
960 addition, open symbols were post-addition. The  $\delta^{15}\text{N}$  values were normalized to upstream  
961 reference  $\delta^{15}\text{NO}_3^-$  on each date.

962

963 **Figure 5**

964 Time course of  $\delta^{15}\text{N}$  in biofilm, fine benthic organic matter (FBOM) and coarse benthic organic  
965 matter (CBOM) (all left axis) and algae (right axis in pasture streams) at the 40 m station in the  
966 second-order forest stream (A), the 50 m station in the second-order pasture stream (B) and the  
967 40 m station in the third-order (C) pasture stream. The  $\delta^{15}\text{N}$  values were normalized to upstream  
968 values. Shaded area represents the period of  $^{15}\text{NH}_4^+$  addition.

969

970 **Figure 6**

971 Longitudinal transects of  $^{15}\text{N}$  values in organic matter compartments on the final day of the  $^{15}\text{N}$   
972 addition. Values are  $\delta^{15}\text{N}$  in algae (right axis in pasture streams), biofilm, fine benthic organic  
973 matter (FBOM) and coarse benthic organic matter (CBOM) in the second-order forest stream  
974 (A), second-order pasture stream (B) and third-order pasture stream (C).

975

976 **Figure 7**

977 The  $\delta^{15}\text{N}$  in the riparian grass *P. repens* on day 20 in the second-order pasture stream (A) and  
978 third-order pasture stream (B). Open symbols represent pre-addition values, closed symbols are  
979 day 20. All  $\delta^{15}\text{N}$  values were normalized to upstream values.