## Young organic matter as a source of carbon dioxide outgassing from Amazonian rivers

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Rivers are generally supersaturated with respect to carbon dioxide, resulting in large gas evasion fluxes that can be a significant component of regional net carbon budgets<sup>1,2</sup>. Amazonian rivers were recently shown to outgas more than ten times the amount of carbon exported to the ocean in the form of total organic carbon or dissolved inorganic carbon<sup>1</sup>. High carbon dioxide concentrations in rivers originate largely from in situ respiration of organic carbon<sup>1-3</sup>, but little agreement exists about the sources or turnover times of this carbon<sup>2,4,5</sup>. Here we present results of an extensive survey of the carbon isotope composition  $({}^{13}C \text{ and } {}^{14}C)$ of dissolved inorganic carbon and three size-fractions of organic carbon across the Amazonian river system. We find that respiration of contemporary organic matter (less than five years old) originating on land and near rivers is the dominant source of excess carbon dioxide that drives outgassing in medium to large rivers, although we find that bulk organic carbon fractions transported by these rivers range from tens to thousands of years in age. We therefore suggest that a small, rapidly cycling pool of organic carbon is responsible for the large carbon fluxes from land to water to atmosphere in the humid tropics.

Riverine  $CO_2$  concentrations in Amazonian lowlands are 5–30 times supersaturated with respect to atmospheric equilibrium<sup>1</sup>; such conditions may be prevalent throughout the humid tropics. *In situ* 

respiration is the primary source of CO<sub>2</sub> sustaining supersaturation in rivers, although inputs from groundwater supersaturated by soil respiration can be important in small systems and from submerged root respiration in floodplain-influenced systems1-3,6-8. Although air-water gas exchange is a bi-directional process, atmospheric  $CO_2$  invasion has a negligible role compared to the large  $CO_2$  evasion fluxes, except at low supersaturation<sup>2,3,6,7</sup>. <sup>13</sup>C and <sup>14</sup>C isotopes can provide constraints on sources and turnover times of organic carbon fuelling river respiration, but no previous tropical study has used a dual-isotope approach to address these questions. Studies in temperate eastern USA provide contrasting findings. In the Hudson River, up to 70% of the centuries-old terrestrial organic carbon entering the river is respired in transit, and the average age of riverine organic carbon decreases downstream<sup>2</sup>. However, the youngest components of dissolved organic carbon (DOC) are preferentially respired in the York River<sup>5</sup>, and modern dissolved inorganic carbon (DIC) in the Parker River may be explained by respiration of young DOC produced within the estuary<sup>4</sup>. Documenting key patterns and controls on CO<sub>2</sub> sources in diverse ecosystems is critical to advance our understanding of CO2 outgassing from rivers and its contribution to regional net carbon budgets.

To identify dominant sources and turnover times of riverine carbon throughout the Amazon basin, we analysed <sup>14</sup>C and <sup>13</sup>C of



Figure 1 | Amazon basin and river sites sampled for carbon isotopes. We used GTOPO30 elevation<sup>28</sup> and a regional river network data set<sup>29</sup> to categorize each site by topography according to the percentage of the drainage area above 1,000 m elevation: mountain (diamond),  $\geq$  50% (16 sites); mixed (square),  $\geq 10\%$  (11 sites); and lowland (circle), <10% (33 sites). Mountain sites are found only in the Andean Cordillera, while mixed sites are large rivers draining both mountain and lowland areas. Site numbers are displayed. Distribution of sites by drainage area (river size) and mean basin elevation is shown. Additional information is in Supplementary Table S1.

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**Figure 2** | **Distribution of** <sup>14</sup>**C and** <sup>13</sup>**C isotopes.** Lower  $\Delta^{14}$ C values indicate older carbon. Symbol shapes are as in Fig. 1; Red (shaded), blue, orange, and green symbols represent CO<sub>2</sub>, DOC, FPOC and CPOC respectively. Isotopic ranges of carbon sources are shown by grey boxes. Plant  $\delta^{13}$ C ranges from -32% to -26% for lowland plant material fixed via the C<sub>3</sub> photosynthetic pathway dominant among plants, to -16% to -12% for tropical C<sub>4</sub> grasses<sup>24,30</sup>. Phytoplankton take up H<sub>2</sub>CO<sub>3</sub> and impose <sup>13</sup>C fractionations of  $\sim -20\%$ , leading to biomass  $\delta^{13}$ C values of -32 to -45% in mixed and lowland rivers<sup>8,10,30</sup>, beyond the range found in our lowland organic carbon and CO<sub>2</sub> observations. Carbonate rocks and CO<sub>2</sub> from solid earth degassing are <sup>14</sup>C-free ( $\Delta^{14}$ C = -1,000%; ref. 10).  $\delta^{13}$ C of carbonate rocks is  $\sim 0\%$ ; degassed CO<sub>2</sub> ranges from -6% for volcanic to 10‰ for metamorphic sources<sup>10</sup>. The carbonate weathering region represents CO<sub>2</sub> in equilibrium with HCO<sub>3</sub><sup>-</sup> resulting from weathering by CO<sub>2</sub> derived from respired, contemporary C<sub>3</sub> or C<sub>4</sub> plant material<sup>10</sup>.

DIC, DOC, and suspended fine and coarse particulate organic carbon fractions (FPOC and CPOC), grouping sites topographically (Fig. 1). This survey represents the most extensive dual carbon isotope inventory so far in a large, diverse basin, and the first <sup>14</sup>C analysis of DIC in Amazonian rivers. It complements but greatly exceeds previous carbon isotope surveys<sup>5,7,9</sup>, enabling an integrated assessment of carbon cycling.

DIC is composed of dissolved CO<sub>2</sub> and bicarbonate and carbonate ions in pH-dependent chemical and isotopic equilibrium<sup>10</sup>. In studies of marine and homogeneous river systems, where pH is nearly uniform, it has been the convention to report the isotopic composition of total DIC, which is directly measured. However, when the turnover of DIC by CO2 fluxes is as rapid as in many of these tropical rivers, a quasi-steady-state condition constrains the isotopic composition of outputs by CO2 outgassing to equal that of inputs by CO<sub>2</sub> production from respiration<sup>7,8</sup>. In response to this pre-eminence of CO<sub>2</sub> sources and the large range of observed pH (3.8 to 8.7), we base our analysis on the calculated isotopic values of  $CO_2$ gas in equilibrium with DIC, avoiding the need to discuss the isotopic variation in <sup>13</sup>C of DIC caused by changes in pH alone (see Methods and Supplementary Discussion). We report measured  $\delta^{13}$ C-DIC in Supplementary Table S3. For <sup>14</sup>C we use  $\Delta^{14}$ C notation, which includes a correction for <sup>13</sup>C such that  $\Delta^{14}$ C of CO<sub>2</sub> and DIC are equal at any pH (ref. 11).

 $CO_2$  in nearly all lowland rivers had a  $\Delta^{14}C$  signature bound within the range of local atmospheric  $\Delta^{14}C$ -CO<sub>2</sub> estimated for the lowland



Figure 3 | Temporal evolution of <sup>14</sup>C-CO<sub>2</sub> at four lowland sites from medium to large rivers in the Ji-Parana basin and Rio Negro. Legend is as in Fig. 1. These sites drain continental shields and were analysed for <sup>14</sup>C-DIC 2–3 times between 1996 and 2003.

DIC sampling period (1996-2003, Fig. 2; Methods). Testing of thermonuclear bombs resulted in a large increase in atmospheric <sup>14</sup>CO<sub>2</sub>, reaching a peak in the late 1960s and steadily decreasing afterwards<sup>12,13</sup>. Continual changes in atmospheric radiocarbon content means that we need to assess the mean age for modern carbon samples by the offset between riverine  $\Delta^{14}$ C and the sampling year's annual mean atmospheric  $\Delta^{14}$ C-CO<sub>2</sub> (ref. 14). From 1996 to 2003,  $\Delta^{14}\text{C-CO}_2$  at four supersaturated sites in medium to large lowland rivers decreased by 32-42‰, equivalent to the annually averaged atmospheric  $\Delta^{14}$ C-CO<sub>2</sub> decrease of 36‰ (-5.6 ± 2.3‰ yr<sup>-</sup> (Fig. 3). Thus, atmospheric offsets remained roughly constant, suggesting constant respiratory organic carbon turnover times for each basin. The range of these offsets suggests rapid turnover of photosynthetically sequestered atmospheric CO<sub>2</sub>; outgassed CO<sub>2</sub> is derived from atmospheric  $CO_2$  sequestered within <4 yr in the medium-sized Ji-Parana basin ( $\Delta^{14}$ C offsets: 14 ± 6‰, n = 8) and 4–7 yr previously at the Rio Negro mouth ( $\Delta^{14}$ C offsets: 29 ± 9‰, n = 3).

Rapid carbon turnover is probably widespread across Amazonian rivers. However, DIC from carbonate mineral weathering, with its typically enriched  $\delta^{13}$ C and highly depleted  $\Delta^{14}$ C (Fig. 2), commonly obscures the influence of respiration. To focus on sites where DIC originates largely as respired CO<sub>2</sub>, we identified sites draining substantial carbonate lithologies through their inorganic solute composition<sup>15</sup> (Methods and Supplementary Discussion). Only lowland sites had little potential for direct, substantial carbonate contributions to DIC (38 samples in 25 lowland sites, Supplementary Fig. S1). Observations in carbonate-free lowlands contained supersaturated, predominantly contemporary CO<sub>2</sub> (Table 1, Fig. 2) and could be divided into two groups based on atmospheric  $\Delta^{14}$ C-CO<sub>2</sub> offsets. The largest group (32 samples from 21 sites) encompassed first-order streams and large rivers carrying contemporary CO<sub>2</sub> with atmospheric  $\Delta^{14}$ C-CO<sub>2</sub> offsets ranging from -3 to 38‰  $(14 \pm 11\%)$ , indicating a mean CO<sub>2</sub> age of approximately 2 yr (and <5 yr in 87% of the observations); enriched  $\delta^{13}$ C-CO<sub>2</sub> in the Ji-Parana region (-17.5  $\pm$  2.2‰, n = 21) suggest a C<sub>4</sub> plant influence. The second group of carbonate-free observations consisted of six samples in three small and two medium-sized rivers in the Ji-Parana region having considerable negative  $\Delta^{14}$ C offsets  $(-32 \pm 21\%)$ , indicating a mean source age of several decades. Relatively depleted  $\delta^{13}$ C-CO<sub>2</sub> values (-20.0 ± 1.7‰) compared to Ji-Parana rivers from the first group suggest an influence from groundwater influx of aged soil CO2 with a significant terrestrial C<sub>3</sub> plant source relative to other rivers in that highly deforested region. In both groups, respiration of submerged tree and grass roots can be excluded as important CO<sub>2</sub> sources because our data set was predominantly collected during low water.

Isotopic signatures in carbonate-free lowland rivers demonstrate

Table 1 | <sup>14</sup>C and <sup>13</sup>C isotopic composition for each carbon fraction

| Site             | CO <sub>2</sub>                   |                                    | DIC                                | DOC                           |                                    | FPOC                                |                                    | CPOC                              |                                    |
|------------------|-----------------------------------|------------------------------------|------------------------------------|-------------------------------|------------------------------------|-------------------------------------|------------------------------------|-----------------------------------|------------------------------------|
| Category         | $\Delta^{14}C$                    | $\delta^{13}C$                     | $\delta^{13}C$                     | $\Delta^{14}C$                | $\delta^{13}C$                     | $\Delta^{14}C$                      | $\delta^{13}C$                     | $\Delta^{14}C$                    | $\delta^{13}C$                     |
| Mountain         | -240 ± 233 (14)                   | $-12.9 \pm 2.2$                    | $-4.9 \pm 2.7$                     | 94 ± 176 (6)                  | $-26.0 \pm 3.0$                    | -202 ± 198 (8)                      | $-25.7 \pm 1.7$                    | -39 ± 146 (9)                     | -27.0 ± 1.6                        |
| Mixed<br>Lowland | $-14 \pm 99$ (11)<br>89 ± 44 (43) | $-19.4 \pm 1.2$<br>$-20.1 \pm 3.6$ | $-14.2 \pm 2.9$<br>$-17.0 \pm 5.9$ | 196 ± 59 (9)<br>177 ± 64 (15) | $-29.0 \pm 0.6$<br>$-29.0 \pm 0.9$ | $-135 \pm 141 (10)$<br>90 ± 55 (10) | $-28.2 \pm 0.9$<br>$-29.8 \pm 1.8$ | $-124 \pm 66 (9)$<br>112 ± 83 (4) | $-27.7 \pm 0.9$<br>$-28.5 \pm 0.7$ |
| Carbonate-free   | 98 ± 20 (38)                      | $-19.8 \pm 3.7$                    | $-17.1 \pm 6.2$                    | 175 ± 67 (11)                 | $-29.1 \pm 0.7$                    | 129 ± 10 (6)                        | $-29.2 \pm 2.1$                    | 121 ± 100 (3)                     | $-28.4 \pm 0.9$                    |

All carbonate-free sites are in the lowlands. Data are reported as mean  $\pm$  s.d. (number of samples), in per mil (‰). The number of samples for <sup>13</sup>C is the same as that shown for <sup>14</sup>C. The number of samples for DIC is the same as for CO<sub>2</sub>.

that CO<sub>2</sub> originated from heterotrophic respiration of contemporary C<sub>3</sub> and C<sub>4</sub> organic carbon (Fig. 2), yet CO<sub>2</sub> appears isotopically distinct from the associated bulk organic carbon load. All organic carbon fractions were considerably depleted in <sup>13</sup>C relative to CO<sub>2</sub> (Table 1, Fig. 2 and Supplementary Discussion). Furthermore, DOC was generally older than CO<sub>2</sub>, FPOC showed a bimodal distribution with ages similar to CO<sub>2</sub> in non-turbid rivers and older carbon in high-sediment rivers, and CPOC exhibited a wide range of ages. We conclude that *in situ* respiration is fuelled largely by an unmeasured organic subfraction that cycles over a period of less than about five years and typically makes up a small component of the riverine organic carbon load. <sup>13</sup>C enrichment of CO<sub>2</sub> relative to organic carbon indicates that this missing organic carbon source fuelling river respiration is disproportionately composed of river-corridor and floodplain C4 grasses, which may be intrinsically more biodegradable7,16.

Mountain and mixed rivers contain older dissolved CO<sub>2</sub>  $(\Delta^{14}C = -749 \text{ to } 96\%)$  with clear carbonate mineral dissolution signatures. However, observed CO<sub>2</sub> supersaturation in these rivers must be generated by CO<sub>2</sub> sources other than carbonates (Supplementary Discussion). These CO<sub>2</sub> fluxes gradually flush out geologically derived DIC, replacing its isotopic signature. Indeed, a  $CO_2$  trend of increasing  $\Delta^{14}C$  and decreasing  $\delta^{13}C$  is observed from the Peruvian Andes to the Amazon mainstem, with  $\Delta^{14}$ C-CO<sub>2</sub> reaching 30-76‰ in the central mainstem, still below atmospheric levels (Fig. 2). Along the Ucavali and western-central mainstem, inputs of probably young CO<sub>2</sub> from *in situ* respiration and lowland tributaries drive large evasion fluxes of <sup>14</sup>C-depleted CO<sub>2</sub> derived from carbonate dissolution. In the more arid Ucayali mountain headwaters, highly depleted  $\Delta^{14}$ C-CO<sub>2</sub> (<-500‰), enriched  $\delta^{13}\text{C-CO}_2~(\sim-11\text{\%}),$  and considerable CO\_2 supersaturation point to solid-earth degassing as a large dry-season CO<sub>2</sub> source, as documented in other tectonically active mountain ranges<sup>17</sup>.

Control of respiration by a small fraction of organic carbon does not imply that bulk organic carbon is unreactive. On the contrary, measured organic carbon fractions appear to be mineralized throughout the river system. The strongest evidence is that FPOC generally becomes younger and more depleted in <sup>13</sup>C downstream from mountain sites (Fig. 2, Table 1), where all organic carbon fractions mirror the high-altitude  ${}^{13}$ C enrichment in plants of  $\sim 1\%$ per 1,000 m elevation<sup>18,19</sup>. Within 1,000 km from mountain headwaters,  $\delta^{13}$ C-FPOC becomes nearly indistinguishable from lowland carbon. With FPOC tightly associated with mineral surfaces<sup>19,20</sup>, no downstream changes in the ratio of FPOC to fine suspended sediment concentration (0.8-1.5%, Supplementary Table S2), and >85% of the Amazon mainstem mineral load ultimately originating in the Andes<sup>21</sup>, this observation implies nearly complete mineralization of old Andean FPOC and replacement with new lowland organic carbon during transit within the river or during long-term floodplain storage<sup>7</sup>. Even if all FPOC leaving the Andes were mineralized within the river channel, the resulting  $CO_2$  flux would be 1/40 of total  $CO_2$ evasion fluxes<sup>1</sup>, hardly affecting the isotopic signature of CO<sub>2</sub>. CPOC and DOC descending from the Andes follow identical trends of gradual <sup>13</sup>C depletion to lowland values (Fig. 2), although without a conservative mineral carrier as for FPOC, simple dilution by lowland

organic carbon cannot be ruled out. DOC is generally modern (<50 yr old) everywhere, demonstrating that old DOC does not escape from the basin.

Isotopic evidence for dominance of respiration fluxes by a rapidly cycling, typically small fraction of total organic carbon confirms the hypothesis previously posited for the Amazon mainstem from respiration<sup>8,16</sup>, <sup>13</sup>C (ref. 7), and mass balance studies<sup>1,7</sup>. It implies that gradual consumption or replacement of old fractions in bulk organic carbon can occur in parallel with high rates of respiration of a highly labile organic carbon subfraction. This paradigm has been advanced for tropical and temperate soils<sup>22</sup>, providing strong conceptual linkage between aquatic and terrestrial carbon dynamics<sup>23</sup>. It suggests that deforestation in the Amazon leads to immediate changes to the organic sources of riverine heterotrophic energy and argues that such impact is not inconsistent with apparent lag times observed in bulk organic carbon composition<sup>24</sup>. Although the mechanism proposed here may be widespread across the humid tropics<sup>1</sup> and appears consistent with radiocarbon observations from temperate rivers<sup>2,4,5</sup>, it is probable that certain river systems, such as those draining eroding peats, are fuelled by old organic carbon. Explicit accounting of isotopic signatures of CO<sub>2</sub> outgassing from different river types may be required to accurately interpret isotopebased regional tropospheric CO2 inversions.

## METHODS

Sample collection and analysis. Samples analysed for <sup>14</sup>C-DIC were collected between 1991 and 2003, whereas organic-carbon <sup>14</sup>C samples are from 1995–1996. All samples were preserved with mercuric chloride immediately after collection at mid-depth from the deepest section of the channel. DIC samples were prepared as described in ref. 7 and stored in tightly capped glass bottles for up to 24 months; in the lab, the top half of the bottle was drawn into a vacuum line (eliminating particles) and stripped of CO<sub>2</sub> after acidification<sup>7</sup>. CPOC (63–2,000  $\mu$ m) was isolated either by sieving or with a plankton net, FPOC (0.1–63  $\mu$ m) by tangential flow microfiltration, and DOC (1,000 atomic mass units to 0.1  $\mu$ m) by tangential flow ultrafiltration<sup>19</sup>. Final concentration and drying was achieved by centrifugal evaporation or freeze drying<sup>19</sup>, and the dried powder stored in the dark at ambient temperature for up to 6 by:. Ultrafiltration yields ranged from 40% in the Andes to 80% in the lowlands<sup>19,25</sup>. Organic samples were combusted as in ref. 7.

Cryogenically purified CO2 from organic carbon and DIC was analysed for stable isotope and radiocarbon by dual-inlet Isotope Ratio Mass Spectrometry and Accelerator Mass Spectrometry (AMS)<sup>26</sup>, respectively; >90% of <sup>14</sup>C analyses were carried out at the Lawrence Livermore National Laboratory's Center for Accelerator Mass Spectrometry, and the rest at the University of Arizona AMS Laboratory. CO2 extracted from DIC was stored in sealed glass ampoules for up to 8 yr.  $^{13}$ C is reported in  $\delta^{13}$ C notation versus the PDB standard<sup>10</sup>. Radiocarbon values are reported as age-corrected  $\Delta^{14}$ C adjusted for sample  $\delta^{13}$ C (ref. 11); carbon is defined as modern when it originates after 1890 (ref. 11). Absolute  $\Delta^{14}$ C and  $\delta^{13}$ C analysis errors (1 $\sigma$ ) are typically <6‰ and <0.2‰, respectively. Isotopes for all carbon fractions were not always analysed at each site. Additional analyses include pH, major ions, alkalinity and total carbon fraction concentrations (Supplementary Table S2). Major ions were quantified by ion chromatography. Alkalinity was measured by Gran titration, or estimated from temperature, pH and DIC when not measured. CO2 concentrations were either measured directly by headspace equilibration (A.K.A., manuscript in preparation) or estimated from temperature, pressure, pH, DIC, and alkalinity. Sampling sites are grouped by topographic characteristics (Fig. 1). In the discussion, grouped observations are reported as mean  $\pm$  standard deviation

Isotopic calculation of CO<sub>2</sub> gas in equilibrium with DIC. DIC is composed of dissolved carbonate species (H<sub>2</sub>CO<sub>3</sub>(aq), HCO<sub>3</sub><sup>-</sup>(aq), and CO<sub>3</sub><sup>2-</sup>(aq)) in temperature- and pH-dependent equilibrium with one another. Isotopic fractionation occurs during conversion from one species to another and dissolution of CO<sub>2</sub> gas<sup>10</sup>; CO<sub>2</sub> gas is hereafter referred to as simply CO<sub>2</sub>.  $\delta^{13}$ C of CO<sub>2</sub> gas in equilibrium with DIC is calculated from measured  $\delta^{13}$ C-DIC and pH, and from temperature-dependent isotopic equilibrium fractionations between CO<sub>2</sub> and DIC species<sup>10,27</sup>. pH can vary dramatically in a large basin and is largely a function of weathering lithologies (Supplementary Table S2).  $\Delta^{14}$ C is defined to be insensitive to mass-dependent fractionation<sup>11</sup>; as a result  $\Delta^{14}$ C-CO<sub>2</sub> =  $\Delta^{14}$ C-DIC. Although a focus on isotopes of CO<sub>2</sub> instead of DIC is unconventional, it yields more straightforward assessments of the impact of respiration and air-water gas exchange on DIC across geochemically diverse rivers. Radiocarbon trends in atmospheric CO2. Measurements from Schauinsland Station, Germany, were used to characterize atmospheric  $\Delta^{14}$ C-CO<sub>2</sub> from 1991 to 2003 (refs 12, 14). The uncharacterized effect of seasonal and short-term atmospheric variability is minimized by comparing river <sup>14</sup>C only against timeweighted annual means<sup>14</sup>. A constant +8‰ offset was added to Schauinsland annual means to account for a 5‰ depletion from regional fossil-fuel emissions at Schauinsland relative to the well-mixed, mid-latitude European troposphere (Jungfraujoch site14), and approximately 3% further depletion at the midlatitude troposphere relative to tropical South America<sup>12,13</sup>. Atmospheric  $\Delta^{14}$ C-CO2 composition within the Amazon basin is unknown, but seasonal and regional variability may be as large as 10‰ (ref. 13). Riverine  $\Delta^{14}$ C values within 5% of our estimated atmospheric annual average for the sampling year probably represent carbon turnover times of one year or less. Mid-1990s tropospheric CO2 can be characterized by a partial pressure  $(p_{CO_2})$  of 370 p.p.m. and  $\delta^{13}C$ composition of -8‰ (refs 10, 12, 13).

**Carbonate mineral dissolution and DIC.** Carbonate weathering is a source of high DIC concentrations enriched in  $\delta^{13}$ C and highly depleted in  $\Delta^{14}$ C when, as is most common, it involves the dissolution of <sup>14</sup>C-dead carbonate minerals (CaCO<sub>3</sub>(s)) by carbonic acid (H<sub>2</sub>CO<sub>3</sub>(aq)) from respired modern plant matter in soils (Fig. 2 and Supplementary Discussion). We used alkalinity and dissolved inorganic cation composition to identify sites draining substantial carbonate lithologies<sup>15</sup> (Supplementary Fig. S1). Although exchange with the atmosphere or input of respired CO<sub>2</sub> may erase the isotopic signature of carbonate dissolution, we identified and excluded carbonate-influenced sites in order to focus unambiguously on the isotopic relationship between organic carbon and DIC.

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**Supplementary Information** is linked to the online version of the paper at www.nature.com/nature.

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