- <sup>1</sup> Title: Carbon cycling in mature and regrowth forests globally: a macroecological synthesis based on the
- 2 Global Forest Carbon (ForC) database

### 3 Authors:

- 4 Kristina J. Anderson-Teixeira<sup>1,2</sup>\*
- <sup>5</sup> Valentine Herrmann<sup>1</sup>
- 6 Becky Banbury Morgan<sup>1,3</sup>
- <sup>7</sup> Ben Bond-Lamberty<sup>4</sup>
- 8 Susan C. Cook-Patton<sup>5</sup>
- Abigail E. Ferson<sup>1,6</sup>
- 10 Helene C. Muller-Landau<sup>2</sup>
- Maria M. H. Wang<sup>1,7</sup>

### 12 Author Affiliations:

- Conservation Ecology Center; Smithsonian Conservation Biology Institute; National Zoological Park,
   Front Royal, VA 22630, USA
- Center for Tropical Forest Science-Forest Global Earth Observatory; Smithsonian Tropical Research
   Institute; Panama, Republic of Panama
- 3. School of Geography, University of Leeds, Leeds, UK
- Joint Global Change Research Institute, Pacific Northwest National Laboratory, College Park
   Maryland 20740, USA
- 5. The Nature Conservancy; Arlington VA 22203, USA
- 6. College of Natural Resources, University of Idaho; Moscow, Idaho 83843, USA
- 7. Grantham Centre for Sustainable Futures and Department of Animal and Plant Sciences, University of Sheffield, Western Bank, Sheffield, South Yorkshire S10 2TN, UK
- \*corresponding author: teixeirak@si.edu; +1 540 635 6546

# 25 Summary

- Background. Forests are major components of the global carbon (C) cycle and thereby strongly influence
- 27 atmospheric carbon dioxide (CO2) and climate. However, efforts to incorporate forests into climate models
- <sup>28</sup> and CO<sub>2</sub> accounting frameworks have been constrained by a lack of accessible, global-scale synthesis on how
- <sup>29</sup> C cycling varies across forest types and stand ages.
- 30 Methods/Design. Here, we draw from the Global Forest Carbon Database, ForC, to provide a macroscopic
- overview of C cycling in the world's forests, giving special attention to stand age-related variation.
- <sup>32</sup> Specifically, we use 11923 ForC records for 34 C cycle variables from 865 geographic locations to characterize
- ensemble C budgets for four broad forest types tropical broadleaf evergreen, temperate broadleaf,
- temperate conifer, and taiga. We include estimates for both mature and regrowth (age <100 years) forests,
- 35 and quantify trends with stand age in regrowth forests for all variables with sufficient data.
- 36 Review Results/Synthesis. The rate of C cycling generally increased from boreal to tropical regions in both
- mature and regrowth forests, whereas C stocks showed less directional variation. Net ecosystem production
- of mature forests was indistinguishable across biomes. The majority of flux variables, together with most live
- biomass pools, increased significantly with stand age when fit with logarithmic functions.
- 40 Discussion. As climate change accelerates, understanding and managing the carbon dynamics of forests is
- 41 critical to forecasting, mitigation, and adaptation. This comprehensive and synthetic global overview of C
- stocks and fluxes across biomes and stand ages will help to advance these efforts.
- 43 Key words: forest ecosystems; carbon cycle; stand age; productivity; respiration; biomass; global

# 44 Background

Forest ecosystems are shaping the course of climate change through their influence on atmospheric carbon dioxide (CO<sub>2</sub>; Bonan 2008, Friedlingstein *et al* 2019, IPCC 2018). Despite the centrality of forest C cycling in regulating atmospheric CO<sub>2</sub>, important uncertainties in climate models (Friedlingstein *et al* 2006, Krause *et al* 2018, Bonan *et al* 2019, Di Vittorio *et al* 2020) and CO<sub>2</sub> accounting frameworks (Pan *et al* 2011) can be traced to lack of understanding on how C cycling varies across forest types and in relation to stand history. This requires accessible, comprehensive, and large-scale databases with global coverage, which runs contrary to the traditional way forest C stocks and fluxes have been measured and published. Large-scale synthesis is critical to benchmarking model performance with global data (Clark et al 2017, Luo et al 2012), quantifying the the role of forests in the global C cycle (*e.g.*, Pan *et al* 2011), and using book-keeping methods to quantify actual or potential exchanges of CO<sub>2</sub> between forests and the atmosphere (Griscom *et al* 2017, Houghton 2020).

### 56 Forests in the global C cycle: current and future

A robust understanding of forest impacts on global C cycling is essential. Total annual photosynthesis in forests (gross primary productivity, GPP) is estimated at >69 Gt C yr<sup>-1</sup> (Badgley et al 2019), more than seven times the average annual fossil fuel emissions during 2009-2018 (9.5  $\pm$  0.5 Gt C yr<sup>-1</sup>; Friedlingstein et al 2019). Most of this enormous C uptake is counterbalanced by releases to the atmosphere through ecosystem respiration  $(R_{eco})$  and fire, with forests globally dominant as sources of both soil respiration (Warner et al 2019) and fire (van der Werf et al 2017). In recent years, total forest C uptake has exceeded 62 releases, such that globally forests have been a C sink. Considering only areas remaining in forest, this C sink 63 has averaged  $3.2 \pm 0.6$  Gt C yr<sup>-1</sup> for 2009-2018, offsetting 29% of anthropogenic fossil fuel emissions (Friedlingstein et al 2019). However, deforestation, estimated at ~1 Gt C yr<sup>-1</sup> in recent decades (Pan et al 2011, Tubiello et al 2020), reduces the net forest sink to ~1.1-2.2 Gt C yr<sup>-1</sup> (Friedlingstein et al 2019). The future of the current forest C sink is dependent both upon forest responses to climate change itself and human land use decisions, which will feedback and strongly influence the course of climate change. Regrowing forests in particular will play an important role (Pugh et al 2019), as almost two-thirds of the 69 world's forests were secondary as of 2010 (FAO 2010). As anthropogenic and climate-driven disturbances impact an growing proportion of Earth's forests (Andela et al 2017, McDowell et al 2020), understanding the 71 carbon dynamics of regrowth forests is increasingly important (Anderson-Teixeira et al 2013). Although age trends in aboveground biomass have been relatively well-studied and synthesized globally (Cook-Patton et al 2020), a relative dearth of data and synthesis on other C stocks and fluxes in secondary forests points to an 74 under-filled need to characterize age-related trends in forest C cycling. Such understanding is particularly 75 critical for reducing uncertainty regarding the potential for carbon uptake and climate change mitigation by regrowth forests (Krause et al 2018, Cook-Patton et al 2020). Understanding, modeling, and managing forest-atmosphere CO<sub>2</sub> exchange is thus central to efforts to mitigate climate change (Grassi et al 2017, Griscom et al 2017, Cavaleri et al 2015).

## 80 Evolution of forest C cycle research

- For more than half a century, researchers have sought to understand how forest carbon cycling varies across stands, including those of different biomes (e.g., Lieth 1973, Luyssaert *et al* 2007) and stand ages (e.g.,
- Odum 1969, Luyssaert et al 2008). Over this time, an increasingly refined conceptual understanding of the

elements of ecosystem C cycles has developed, as a growing number of variables have been defined (e.g., Chapin et al 2006), along with appropriate measurement methods (e.g., Clark et al 2001). New technology has also enabled researchers to directly measure an expanding set of variables, notably including the development of continuous measurements of soil CO<sub>2</sub> efflux (Kuzyakov 2006) and ecosystem-atmosphere CO<sub>2</sub> 87 exchange (Baldocchi et al 2001). Measurement techniques have been increasingly standardized; for example, of the biomass allometries that strongly influence estimates of most C cycle variables (e.g., Chojnacky et al 2014, Chave et al 2014). Further standardization has been made possible through research networks such as ForestGEO (Anderson-Teixeira et al 2015, Davies et al 2021), NEON (Schimel et al 2007), and FLUXNET (Baldocchi et al 2001, Novick et al 2018). Remote sensing technology has become increasingly useful for global- or regional-scale estimates of a few critical variables (e.g., aboveground biomass,  $B_{aq}$ : Saatchi et al 93 2011, Hu et al 2016, Spawn et al 2020, gross primary productivity, GPP: Li and Xiao 2019), yet measurement and validation of most forest C stocks and fluxes necessarily requires intensive on-the-ground data collection. Alongside these conceptual and methodological developments, there has been a proliferation of measurements 97 across the world's forests. The result of decades of research on forest C cycling is tens of thousands of records 98 distributed across thousands of scientific articles, varying in data formats, units, measurement methods, etc. 99 To address global-scale questions, researchers began synthesizing data into increasingly large databases (e.g., 100 Lieth 1973, Luyssaert et al 2007, Bond-Lamberty and Thomson 2010, Anderson-Teixeira et al 2016, 2018, 101 Cook-Patton et al 2020). The current largest, most comprehensive database on forest C cycling is ForC 102 (Anderson-Teixeira et al 2016, 2018), which contains published estimates of forest ecosystem C stocks and 103 annual fluxes (>50 variables), with different variables capturing distinct ecosystem pools (e.g., woody, foliage, and root biomass; dead wood) and flux types (e.g., gross and net primary productivity; soil, root, and 105 ecosystem respiration). These data are ground-based measurements, and ForC contains associated data 106 required for interpretation (e.g., stand history, measurement methods). Since its most recent publication 107 (Anderson-Teixeira et al 2018), For C has grown 129% through the incorporation of two additional large 108 databases that also synthesized published forest C data: the Global Soil Respiration Database (SRDB; 109 Bond-Lamberty and Thomson 2010, Jian et al 2020) and the Global Reforestation Opportunity Assessment database (GROA; Cook-Patton et al 2020). Following these additions, For C currently contains 39762 records 111

# 114 Biome differences

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Forest C cycling varies enormously across biomes, which cateogrize the world's forests according to major 115 differences in climate, vegetation, etc. Since the early 19th century, it has been recognized that climate plays 116 a dominant role in shaping differences among forests on a global scale (Humboldt and Bonpland 1807, 117 Holdridge 1947). Global scale data syntheses have shown that C fluxes including GPP, net primary 118 productivity (NPP), and soil respiration  $(R_{soil})$  decrease with latitude or, correspondingly, increase with 119 mean annual temperature and, to a lesser extent, precipitation (Fig. 1a; e.g., Lieth 1973, Luyssaert et al 120 2007, Hursh et al 2017, Banbury Morgan et al n.d.). C stocks of mature forests show less directional 121 variation (Fig. 1a). On average, above ground biomass  $(B_{ag})$  tends to decrease with latitude, but not as 122 dramatically as fluxes, and with the highest  $B_{ag}$  forests in relatively cool, moist temperate regions (Keith et 123 al 2009, Smithwick et al 2002, Hu et al 2016). In contrast, standing and downed dead wood (DW<sub>standing</sub>

from 10608 plots and 1532 distinct geographic areas representing all forested biogeographic and climate

zones, making it ideal for assessing how forest C cycling varies across biomes and with respect to stand age.

and  $DW_{down}$ , respectively) and the organic layer (OL) tend to accumulate more in colder climates where decomposition is slow relative to NPP (Harmon *et al* 1986, Allen *et al* 2002).

Correlative analyses relating C cycle variables to climate and other environmental variables have recently been taken to a new level through use of machine-learning algorithms that relate ground-based C cycle data 128 to global maps of environmental covariates, making it possible to create fine-scale global maps of C cycling 129 (e.g., Warner et al 2019, Cook-Patton et al 2020). This approach can be particularly effective when paired with satellite measurements that correlate to C cycle variables of interest; for example, solar-induced 131 chlorophyll fluorescence is useful for fine-scale mapping of gross primary productivity (GPP); Li and Xiao 132 2019), while LiDAR, radar, and optical imagery is being used to estimate  $B_{aq}$  at regional to global scales 133 (e.g., Saatchi et al 2011, Hu et al 2016). Any such analysis is however constrained by the quality and 134 coverage of ground-based estimates of forest C fluxes or stocks (e.g., Schepaschenko et al 2019). While 135 estimates of some variables (e.g.,  $B_{aq}$ , GPP, NPP,  $R_{soil}$ ) are widely available, many remain poorly characterized (e.g., DW; autotrophic respiration,  $R_{auto}$ ) –even at the coarse resolution of biomes. This is a 137 critical limitation not only for understanding forest C cycling, but also for quantifying forest-based climate 138 change mitigation across forest biomes or ecozones (e.g., IPCC 2019).

### 140 Age trends and their variation across biomes

Stand age is another important axis of variation in forest C cycling (Fig. 1b,d). In 1969, E.P. Odum's "The 141 Strategy of Ecosystem Development" laid out predictions as to how forest energy flows and organic matter 142 stocks vary with stand age (Odum 1969). Although the conceptualization of the C cycle in this paper is 143 simplistic by current standards, the paper was foundational in framing the theory around which research on the subject still revolves (Corman et al 2019), and the basic framework still holds, albeit with modest 145 modifications (Anderson-Teixeira et al 2013). Following stand-clearing disturbance, GPP, NPP, and 146 biomass of leaves  $(B_{foliage})$  and fine roots  $(B_{root-fine})$  increase rapidly and thereafter remain relatively stable  $(B_{foliage}, B_{root-fine}, \text{ sometimes } GPP)$  or decline slightly (NPP, sometimes GPP; e.g., ???, Pregitzer)148 and Euskirchen 2004, Amiro et al 2010, Goulden et al 2011). The decline in NPP occurs because R<sub>auto</sub> 149 increases relative to GPP as forests age, corresponding to declining carbon use efficiency with stand age (DeLucia et al 2007, Collalti et al 2020). Heterotrophic respiration, most of which originates from the soil 151  $(R_{het-soil})$  remains relatively constant with stand age (???, Pregitzer and Euskirchen 2004, Goulden et al 152 2011), with the result that net ecosystem production  $(NEP = GPP - R_{eco})$ , where  $R_{eco}$  is total ecosystem respiration) is initially negative, increases to a maximum at intermediate ages, and declines thereafter (???. 154 Pregitzer and Euskirchen 2004, Amiro et al 2010, Goulden et al 2011, Luyssaert et al 2008). The result is 155 that biomass accumulates rapidly in young forests, followed by a slow decline to near zero in old forests (e.g., 156 Lichstein et al 2009, Yang et al 2011). While these trends have been subject of fairly recent qualitative 157 review (Anderson-Teixeira et al 2013), there is need for a synthetic, quantitative review taking advantage of 158 the greatly expanded data now available.

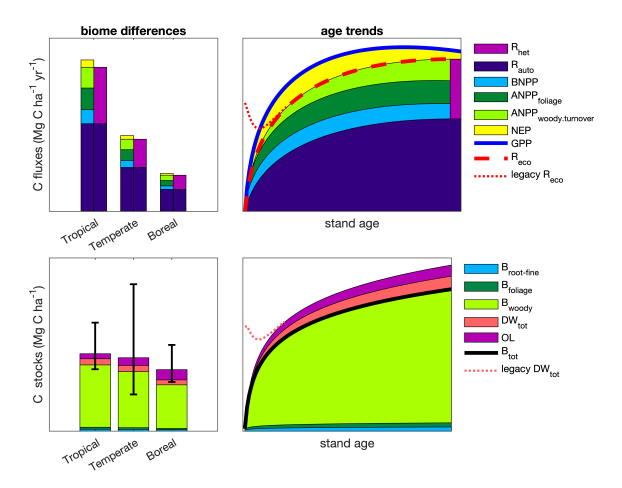


Figure 1 | Schematic diagram summarizing current understanding of biome differences and age trends in forest C cycling. Variables are defined in Table 1. Age trends, which represent idealized dynamics following a disturbance that removes all living vegetation, are an updated version of the classic figure from Odum (1969), with heavy lines corresponding to those in Odum's figure and NEP corresponding to Odum's 'net production'. Here, NEP conists primarily of woody aboveground net primary production  $(ANPP_{woody})$ , while  $ANPP_{woody,turnover}$  is the sum of woody mortality and branch turnover. Dotted lines refer to decomposition of potential 'legacy' organic material produced prior to the disturbance and remaining at the site (e.g., standing and fallen dead wood,  $DW_{tot}$ ; soil organic matter). Error bars on C stocks plot represent within-biome variability, wherein mean biomass is highest in the tropics, but maximum biomass is highest in temperate regions.

In the past several decades, researchers have started asking how age trends—mostly in  $B_{ag}$  or total biomass  $(B_{tot})$  accumulation—vary across biomes. Early research on this theme showed that biomass accumulation rates during secondary succession increase with temperature on a global scale (Johnson et al 2000, Anderson et al 2006) and with water availability in the neotropics (Poorter et al 2016). Most recently, Cook-Patton et al (2020) reinforced these earlier findings with a much larger dataset and crated a high-resolution global map of estimated potential C accumulation rates. However, there has been little synthesis of cross-biome differences in variables other than biomass and its accumulation rate (but see Cook-Patton et al (2020) for DW, OL, and soil C accumulation in young stands). Given the important role of secondary forests in the current and future global C cycle, concrete understanding of age trends in C fluxes and stocks and how these vary across biomes is critical to better understanding of the global C cycle. Accurate estimates of C sequestration rates by regrowth forests are also critical for national greenhouse gas accounting under the IPCC framework (IPCC 2019, Requena Suarez et al 2019) and to quantifying the value of regrowth forests for climate change mitigation (Anderson-Teixeira and DeLucia 2011, Goldstein et al 2020).

Here, we conduct a data-based review of carbon cycling from a stand to global level, and by biome and stand age, using the largest global compilation of forest carbon data, which is available in our open source Global Carbon Forest database (ForC; Fig. 2). Our goal is to provide a comprehensive synthesis on broad trends in forest C cycling that can serve as a foundation for improved understanding of global forest C cycling and highlight where key sources of uncertainty still reside.

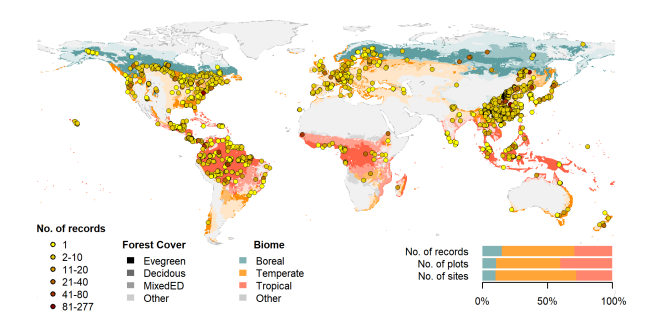


Figure 2 | Map of sites included in this analysis. Symbols are colored according to the number of records at each site. Underlying map shows coverage of evergreen, deciduous, and mixed forests (shading differences; data from Jung et al. 2006) and biomes (color differences). Distribution of sites, plots, and records among biomes is shown in the inset.

## $^{1/8}$ Methods/ Design

This review synthesizes data from the ForC database (Fig. 2; https://github.com/forc-db/ForC; 179 Anderson-Teixeira et al 2016, 2018). For amalgamentes numerous intermediary data sets (e.g., Luyssaert et 180 al 2007, Bond-Lamberty and Thomson 2010, Cook-Patton et al 2020) and original studies. Original 181 publications were referenced to check values and obtain information not contained in intermediary data sets, 182 although this process has not been completed for all records. The database was developed with goals of 183 understanding how C cycling in forests varies across broad geographic scales and as a function of stand age. 184 As such, there has been a focus on incorporating data from regrowth forests (e.g., Anderson et al 2006, 185 Martin et al 2013, Bonner et al 2013) and obtaining stand age data when possible (83% of records in v.2.0; 186 Anderson-Teixeira et al 2018). Particular attention was given to developing the database for tropical forests 187 (Anderson-Teixeira et al 2016), which represented roughly one-third of records in ForC v2.0 188 (Anderson-Teixeira et al 2018). Since publication of ForC v2.0, we imported three large additional databases into ForC via a combination of R scripts and manual edits. First, we imported (via R script) the Global 190 Database of Soil Respiration Database (SRDB v4, 9488 records; Bond-Lamberty and Thomson 2010), and 191 corrections and improvements to SRDB arising from this process were incorporated in SRDB v5 (Jian et al

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(GROA v1.0, 10116 records; Cook-Patton et al 2020, Anderson-Teixeira et al 2020), which itself had drawn
    on an earlier version of ForC. Because all records in GROA were checked against original publications, these
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    records were given priority over duplicates in ForC. Third, we manually incorporated records of annual
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    NEP, GPP, and R_{eco} from the FLUXNET2015 dataset (Pastorello et al 2020), treating these records as
    authoritative when they duplicated earlier records (Appendix S1). We have also added data from individual
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    publications, with a particular focus on productivity (e.g., Taylor et al 2017), dead wood, and ForestGEO
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    sites (e.g., Lutz et al 2018, Johnson et al 2018). A record of data sets added to ForC over the course of its
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    development is available at https://github.com/forc-
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    db/ForC/blob/master/database management records/ForC data additions log.csv. The database
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    version used for this analysis has been tagged as a new release on Github (v3.0) and assigned a DOI through
    Zenodo (DOI: TBD).
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    All measurements originally expressed in units of dry organic matter (OM) were converted to units of C
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    using the IPCC default of C = 0.47 * OM (IPCC 2018). Duplicate or otherwise conflicting records were
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    purged as described in Appendix S1, resulting in a total of 22265 records (56% size of total database).
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    Records were filtered to remove plots that had undergone significant anthropogenic management or major
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    disturbance since the most recent stand initiation event. Specifically, we removed plots with any record of
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    managements manipulating CO<sub>2</sub>, temperature, hydrology, nutrients, or biota, as well as any plots whose site
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    or plot name contained the terms "plantation", "planted", "managed", "irrigated", or "fertilized" (13.9% of
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    duplicate-purged records). We also removed stands that had undergone any notable anthropogenic thinning
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    or partial harvest (5.6% of duplicate-purged records). We retained sites that were grazed or had undergone
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    low severity natural disturbances (<10\% mortality) including droughts, major storms, fires, and floods. We
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    removed all plots for which no stand history information had been retrieved (5.7%). In total, this resulted in
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    17349 records (43.6% of the records in the database) being eligible for inclusion in the analysis.
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    We selected 23 annual flux and 11 C stock variables for inclusion in the analysis (Table 1). These different
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    flux and stock variables represent different pools (e.g., aboveground biomass, root biomass, dead wood) and
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    levels of combination (e.g., total net primary productivity, NPP, versus the individual elements of NPP
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    such as foliage, roots, and branches). We did not analyze soil carbon, which is not a focus of the ForC
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    database. Note that two flux variables, aboveground heterotropic respiration (R_{het-ag}) and total respiration
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    (R_{het}), were included for conceptual completeness but had no records in ForC (Table 1). Records for our
    focal variables represented 90.3% of the total records eligible for inclusion. For this analysis, we combined
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    some of ForC's specific variables into more broadly defined variables. Specifically, net ecosystem exchange
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    (measured by eddy-covariance; Baldocchi et al 2001) and biometric estimates of NEP were combined into
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    the single variable NEP (Table 1). Furthermore, for NPP, aboveground NPP (ANPP), and the litterfall
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    component of ANPP (ANPP<sub>litter fall</sub>), ForC variables specifying inclusion of different components were
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    combined (e.g., measurements including or excluding fruit and flower production and herbivory).
    Throughout ForC, for all measurements drawing from tree census data (e.g., biomass, productivity), trees
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    were censused down to a minimum diameter breast height (DBH) threshold of 10 cm or less. All records were
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    measured directly or derived from field measurements.
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    We grouped forests into four broad biome types based on climate zones and dominant vegetation type
    (tropical broadleaf, temperate broadleaf, temperate needleleaf, and boreal needleleaf) and two age
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    classifications (young and mature). Climate zones (Fig. 2) were defined based on site geographic coordinates
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2020). Second, we imported (via R script) the Global Reforestation Opportunity Assessment database

Table 1. Carbon cycle variables included in this analysis, their sample sizes, and summary of biome differences and age trends.

Variable	Description	N records				
		records	plots	geographic areas	biome differences*	age trend $^\dagger$
Annual fluxes						
NEP	net ecosystem production or net ecosystem exchange (+ indicates C sink)	329	146	88	n.s.	+; xB
GPP	gross primary production $(NPP + R_{auto} \text{ or } R_{eco} + NEP)$	303	115	84	$TrB > TeB \ge TeN \ge BoN$	+; xB
NPP	net primary production $(ANPP + BNPP)$	214	112	74	$TrB > TeB \ge TeN > BoN$	n.s.
ANPP	aboveground NPP	343	236	131	$TrB > TeB \ge TeN > BoN$	+; xB
$ANPP_{woody}$	woody production $(ANPP_{stem} + ANPP_{branch})$	64	53	37	n.s.	+
$ANPP_{stem}$	woody stem production	217	190	117	$TrB > TeN \ge TeB \ge BoN$	n.s.
$ANPP_{branch}$	branch turnover	69	59	42	TrB > TeB > TeN	n.s.
$ANPP_{foliage}$	foliage production, typically estimated as annual leaf litterfall	162	132	88	$TrB > TeB \ge TeN > BoN$	+
$ANPP_{litterfall}$	litterfall, including leaves, reproductive structures, twigs, and sometimes branches	82	70	55	n.s.	+
$ANPP_{repro}$	production of reproductive structures (flowers, fruits, seeds)	51	44	34	n.t.	n.t.
$ANPP_{folivory}$	foliar biomass consumed by folivores	20	12	11	n.t.	n.t.
$M_{woody}$	woody mortality–i.e., $B_{ag}$ of trees that die	18	18		n.t.	n.t.
BNPP	below ground NPP ( $BNPP_{coarse} + BNPP_{fine}$ )	148	116	79	$\mathrm{TrB} > \mathrm{TeN} \geq \mathrm{TeB} \geq \mathrm{BoN}$	+
$BNPP_{coarse}$	coarse root production	77	56	36	$TeN \ge TrB$	n.s.
$BNPP_{fine}$	fine root production	123	99	66	n.s.	+
$R_{eco}$	ecosystem respiration $(R_{auto} + R_{het})$	213	98	70	$TrB > TeB \ge TeN$	+
$R_{auto}$	autotrophic respiration $(R_{auto-ag} + R_{root})$	24	23	15	n.t.	n.t.
$R_{auto-ag}$	aboveground autotrophic respiration (i.e., leaves and stems)	2	2	1	n.t.	n.t.
$R_{root}$	root respiration	181	139	95	$TrB \ge TeB$	+
$R_{soil}$	soil respiration $(R_{het-soil} + R_{root})$	627	411	229	$TrB > TeB > TeN \ge BoN$	n.s.
$R_{het-soil}$	soil heterotrophic respiration	197	156	100	$TrB > TeB \ge TeN$	n.s.
$R_{het-ag}$	aboveground heterotrophic respiration	0	0	0	-	-
$R_{het}$	heterotrophic respiration $(R_{het-ag} + R_{het-soil})$	0	0	0	-	-
Stocks						
$B_{tot}$	total live biomass $(B_{ag} + B_{root})$	188	157	87	$TrB \ge TeB > BoN$	+; xB
$B_{ag}$	aboveground live biomass $(B_{ag-wood} + B_{foliage})$	4466	4072	621	$\operatorname{TrB} \stackrel{-}{\geq} \operatorname{TeN} \geq \operatorname{TeB} > \operatorname{BoN}$	+; xB
$B_{ag-wood}$	woody component of aboveground biomass	115	102	64	$TeN > TrB \ge BoN$	+; xB
$B_{foliage}$	foliage biomass	134	115	72	${\rm TeN} > {\rm TrB} \ge {\rm BoN} \ge {\rm TeB}$	+; xB
$B_{root}$	total root biomass	2329	2298	360	n.s.	+; xB
R	$(B_{root-coarse} + B_{root-fine})$ coarse root biomass	134	120	73	$\text{TeN} > \text{TeB} \ge \text{BoN}$	⊥· vD
$B_{root-coarse}$	fine root biomass	226	180	109		+; xB →: vB
$B_{root-fine}$	deadwood $(DW_{standing} + DW_{down})$	79	73	42	n.s. n.t.	+; xB +; xB
$DW_{tot}$	standing dead wood	79 36	35	22	n.t.	n.t.
$DW_{standing}$ $DW_{down}$	fallen dead wood, including coarse and	278	265	37	n.t. n.t.	n.t. +; xB
OL	sometimes fine woody debris organic layer / litter/ forest floor	474	413	115	n.s.	+; xB

<sup>\*</sup> TrB: Tropical, TeB: Temperate Broadleaf, TeN: Temperate Needleleaf, BoN: Boreal, n.s.: no significant differences, n.t.: not tested  $^{\dagger}$  + or -: significant positive or negative trend, xB: significant age x biome interaction, n.s.: no significant age trend, n.t.: not tested

<sup>&</sup>lt;sup>235</sup> according to Köppen-Geiger zones (Rubel and Kottek 2010). We defined the tropical biome as including all equatorial (A) zones, temperate biomes as including all warm temperate (C) zones and warmer snow

climates (Dsa, Dsb, Dwa, Dwb, Dfa, and Dfb), and the boreal biome as including the colder snow climates (Dsc, Dsd, Dwc, Dwd, Dfc, and Dfd). Any forests in dry (B) and polar (E) Köppen-Geiger zones were excluded from the analysis. We defined leaf type (broadleaf / needleleaf) based on descriptions in original 230 publications (prioritized) or values extracted from a global map based on satellite observations (SYNMAP; 240 Jung et al 2006). For young tropical forests imported from GROA but not yet classified by leaf type, we assumed that all were broadleaf, consistent with the rarity of naturally regenerating needleleaf forests in the 242 tropics. We also classified forests as "young" (< 100 years) or "mature" (> 100 years or classified as 243 "mature", "old growth", "intact", or "undisturbed" in original publication). Assigning stands to these groupings required the exclusion of records for which ForC lacked geographic coordinates (0.4% of sites in 245 full database) or records of stand age (5.7% of records in full database). We also excluded records of stand 246 age = 0 year (0.8% of records in full database). In total, our analysis retained 76.1 of the focal variable records for forests of known age. Numbers of records by biome and age class are given in Table S1. 248 Data were summarized to produce schematics of C cycling for mature forets of each biome. To obtain the 249 values reported in the C cycle schematics, we first averaged any repeated measurements within a plot. Values 250 were then averaged across geographically distinct areas, defined as plots clustered within 25 km of one 251 another (sensu Anderson-Teixeira et al 2018), weighting by area sampled if available for all records. This 252 step was taken to avoid pseudo-replication. 253 We tested whether the C budgets described above "closed"-i.e., whether they were internally consistent. 254 Specifically, we first defined relationships among variables: for example,  $NEP = GPP - R_{eco}$ , 255  $BNPP = BNPP_{coarse} + BNPP_{fine}, DW_{tot} = DW_{standing} + DW_{down}$ ). Henceforth, we refer to the 256 variables on the left side of the equation as "aggregate" fluxes or stocks, and those that are summed as 257 "component" fluxes or stocks, noting that the same variable can take both aggregate and component positions 258 in different relationships. We considered the C budget for a given relationship "closed" when component 259 variables summed to within one standard deviation of the aggregate variable. 260 To test for differences across mature forest biomes, we also examined how stand age impacted fluxes and 261 stocks, employing a mixed effects model ("lmer" function in "lme4" R package; Bates et al 2015) with biome 262 as fixed effect and plot nested within geographic area as random effects on the intercept. When Biome had a 263 significant effect, we looked at a Tukey's pairwise comparison to see which biomes were significantly different from one another. This analysis was run for variables with records for at least seven distinct geographic areas 265 in more than one biome, excluding any biomes that failed this criteria (Table 1). 266 To test for age trends in young (<100yrs) forests, we employed a mixed effects model with biome and 267 log10[stand.age] as fixed effects and plot nested within geographic.area as a random effect on the intercept. This analysis was run for variables with records for at least three distinct geographic areas in more than one 269 biome, excluding any biomes that failed this criteria (Table 1). When the effect of stand age was significant 270 at p  $\leq 0.05$  and when each biome had records for stands of at least 10 different ages, a biome  $\times$  stand.age interaction was included in the model. We note that the logarithmic function fit in this analysis does not 272 always correspond to theoretical expectations (Fig. 1); however, data limitations did not support fitting of 273 functions with more parameters or reliable comparison of different functional forms. Within the data constraints, we deemed a logarithmic function to be the most appropriate functional form 275 for the majority of variables.

To facilitate the accessibility of our results and data, and to allow for rapid updates as additional data

become available, we have automated all database manipulation, analyses, and figure production in R (Team 2020).

## 280 Review Results/ Synthesis

#### 281 Data Coverage

Of the 39762 records in For V3.0, 11923 met our strict criteria for inclusion in this study (Fig. 2). These 282 records were distributed across 5062 plots in 865 distinct geographic areas. Of the 23 flux and 11 stock 283 variables mapped in these diagrams, ForC contained sufficient mature forest data for inclusion in our 284 statistical analyses (i.e., records from  $\geq 7$  distinct geographic areas) for 20 fluxes and 9 stocks in tropical 285 broadleaf forests, 15 fluxes and 8 stocks in temperate broadleaf forests, 14 fluxes and 7 stocks in temperate 286 conifer forests, and 8 fluxes and 7 stocks in boreal forests. For regrowth forests (<100 yrs), ForC contained 287 sufficient data for inclusion in our statistical analyses (i.e., records from  $\geq 3$  distinct geographic areas) for 11 288 fluxes and 10 stocks in tropical broadleaf forests, 16 fluxes and 10 stocks in temperate broadleaf forests, 16 289 fluxes and 10 stocks in temperate conifer forests, and 14 fluxes and 9 stocks in boreal forests. 290

### 291 C cycling in mature forests

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high-biomass forests of the US Pacific Northwest.

Average C cycles for mature tropical broadleaf, temperate broadleaf, temperate conifer, and boreal forests  $\geq$  100 years old and with no known major natural or anthropogenic disturbance are presented in Figures 2-5 (and available in tabular format in the *ForC* release accompanying this publication:

ForC/numbers\_and\_facts/ForC\_variable\_averages\_per\_Biome.csv).

For variables with records from  $\geq 7$  distinct geographic areas, these ensemble C budgets met our criteria for budget "closure". That is, component variables summed to within one standard deviation of their respective aggregate variables in all but one instance. In the temperate conifer biome, the average composite measure of root biomass  $(B_{root})$  was less than the combined average value of coarse and fine root biomass  $(B_{root-coarse})$  and  $B_{root-fine}$ , respectively). This lack of closure was driven by very high estimates of  $B_{root-coarse}$  from

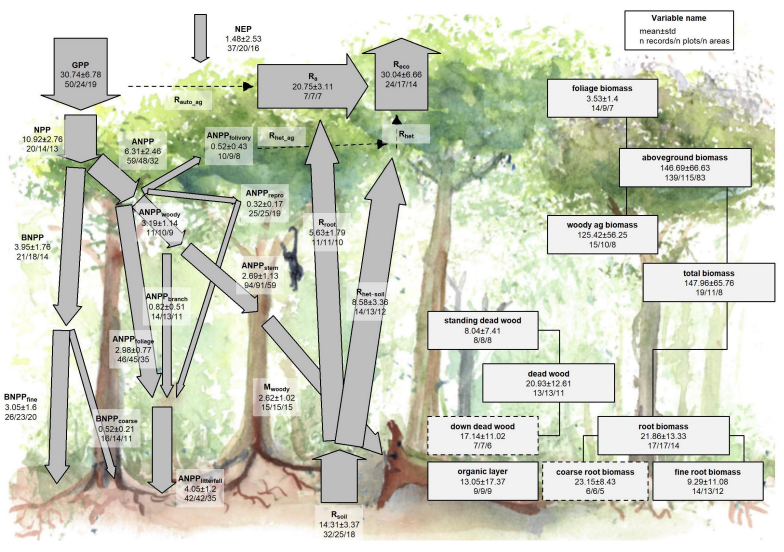


Figure 3 | C cycle diagram for mature tropical broadleaf forests. Arrows indicate fluxes (Mg C  $ha^{-1}$  yr<sup>-1</sup>); boxes indicate stocks (Mg C  $ha^{-1}$ ), with variables as defined in Table 1. Presented are mean  $\pm$  std, where geographically distinct areas are treated as the unit of replication. Note that variables differ in geographical representation, resulting in potential imbalances (Figs. S5-S30). Probability that estimates reflect true biome means scales with the number of distinct geographical areas represented. Dashed shape outlines indicate variables with records from <7 distinct geographic areas, and dashed arrows indicate fluxes with no data. To illustrate the magnitude of different fluxes, arrow width is proportional to the square root of the corresponding flux. An asterisk after a variable name indicates lack of C cycle closure.

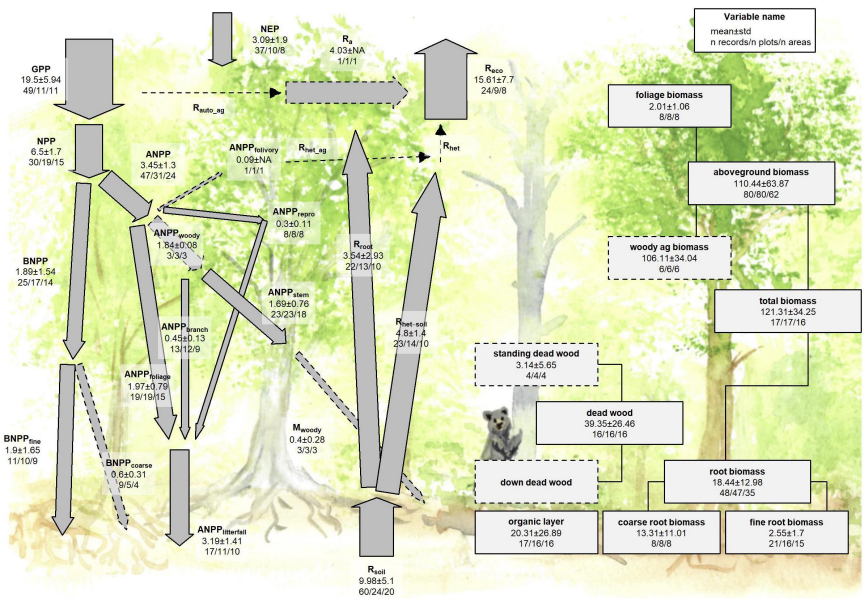


Figure 4 | C cycle diagram for mature temperate broadleaf forests. Arrows indicate fluxes (Mg C  $ha^{-1}$  yr<sup>-1</sup>); boxes indicate stocks (Mg C  $ha^{-1}$ ), with variables as defined in Table 1. Presented are mean  $\pm$  std, where geographically distinct areas are treated as the unit of replication. Note that variables differ in geographical representation, resulting in potential imbalances (Figs. S5-S30). Probability that estimates reflect true biome means scales with the number of distinct geographical areas represented. Dashed shape outlines indicate variables with records from <7 distinct geographic areas, and dashed arrows indicate fluxes with no data. To illustrate the magnitude of different fluxes, arrow width is proportional to the square root of the corresponding flux. An asterisk after a variable name indicates lack of C cycle closure.

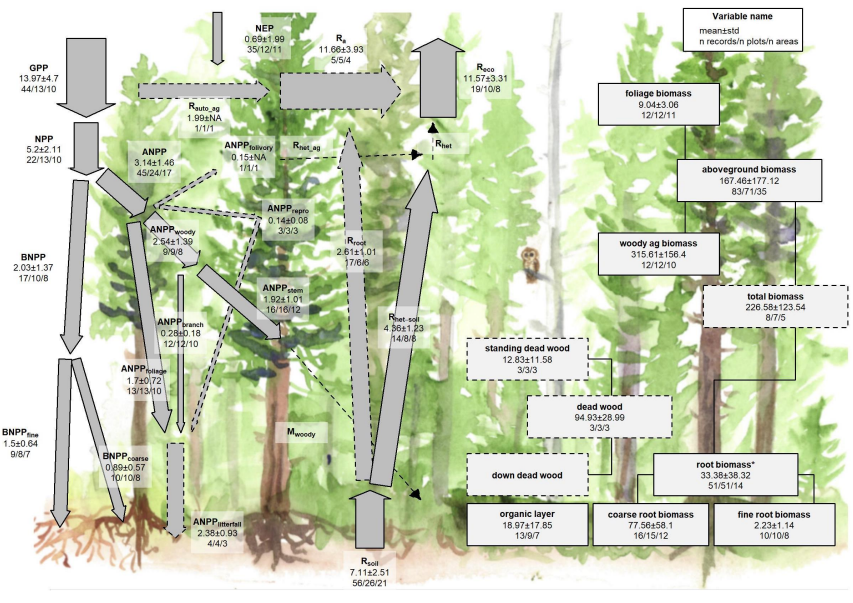


Figure 5 | C cycle diagram for mature temperate conifer forests. Arrows indicate fluxes (Mg C ha $^{-1}$  yr $^{-1}$ ); boxes indicate stocks (Mg C ha $^{-1}$ ), with variables as defined in Table 1. Presented are mean  $\pm$  std, where geographically distinct areas are treated as the unit of replication. Note that variables differ in geographical representation, resulting in potential imbalances (Figs. S5-S30). Probability that estimates reflect true biome means scales with the number of distinct geographical areas represented. Dashed shape outlines indicate variables with records from <7 distinct geographic areas, and dashed arrows indicate fluxes with no data. To illustrate the magnitude of different fluxes, arrow width is proportional to the square root of the corresponding flux. An asterisk after a variable name indicates lack of C cycle closure.

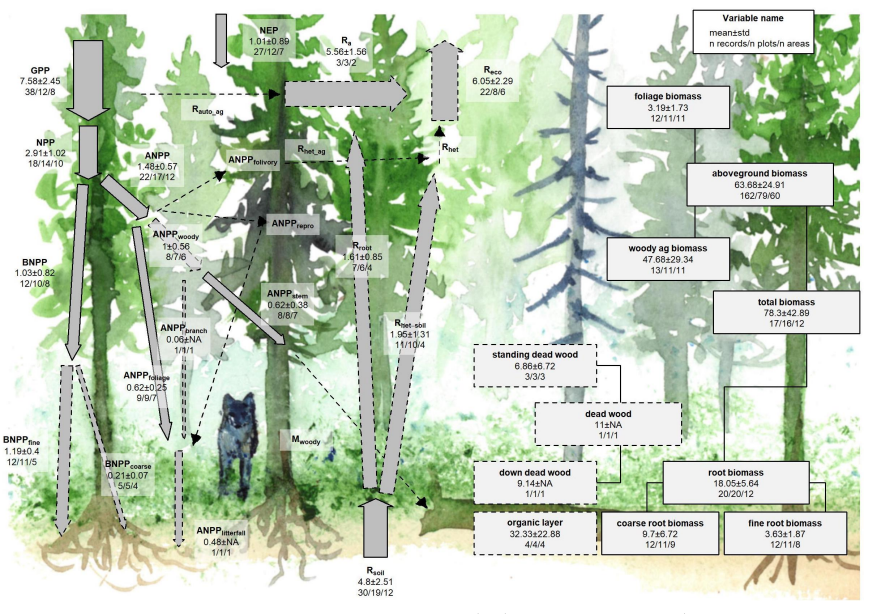


Figure 6 | C cycle diagram for mature boreal conifer forests. Arrows indicate fluxes (Mg C ha<sup>-1</sup> yr<sup>-1</sup>); boxes indicate stocks (Mg C ha<sup>-1</sup>), with variables as defined in Table 1. Presented are mean  $\pm$  std, where geographically distinct areas are treated as the unit of replication. Note that variables differ in geographical representation, resulting in potential imbalances (Figs. S5-S30). Probability that estimates reflect true biome means scales with the number of distinct geographical areas represented. Dashed shape outlines indicate variables with records from <7 distinct geographic areas, and dashed arrows indicate fluxes with no data. To illustrate the magnitude of different fluxes, arrow width is proportional to the square root of the corresponding flux. An asterisk after a variable name indicates lack of C cycle closure.

There were sufficient data to assess differences among biomes in mature forest values for 15 flux variables, and significant differences among biomes were detected for 12 variables (Table 1). In all of these cases-including C fluxes into, within, and out of the ecosystem-C fluxes were highest in tropical forests, intermediate in 304 temperate (broadleaf or conifer) forests, and lowest in boreal forests (Table 1, Figs. 7, S5-S19). Differences 305 between tropical and boreal forests were always significant, with temperate forests intermediate and significantly different from one or both. Fluxes tended to be numerically greater in temperate broadleaf than 307 temperate conifer forests, but the difference was never statistically significant. This pattern held for the 308 following variables: GPP, NPP, ANPP, ANPP,  $ANPP_{stem}$ ,  $ANPP_{branch}$ ,  $ANPP_{foliage}$ , BNPP,  $R_{eco}$ ,  $R_{root}$ , 309  $R_{soil}$ , and  $R_{het-soil}$ . For two of the variables without significant differences among biomes  $(ANPP_{litter\,fall})$ 310 and  $BNPP_{fine}$ ; Figs. S12 and S15, respectively), the same general trends applied but were not statistically 311 significant. Another exception was for  $BNPP_{root-coarse}$ , where all records came from high-biomass forests in 312 the US Pacific Northwest, resulting in marginally higher values for the temperate conifer biome (Table 1, Fig. 313 S14; differences significant in mixed effects model but not in post-hoc pairwise comparison). 314 The most notable exception to the pattern of decreasing flux per unit area from tropical to boreal biomes 315 was NEP, with no significant differences across biomes but with the largest average in temperate broadleaf 316

forests, followed by tropical, boreal, and temperate conifer forests (Figs. 7, S5).

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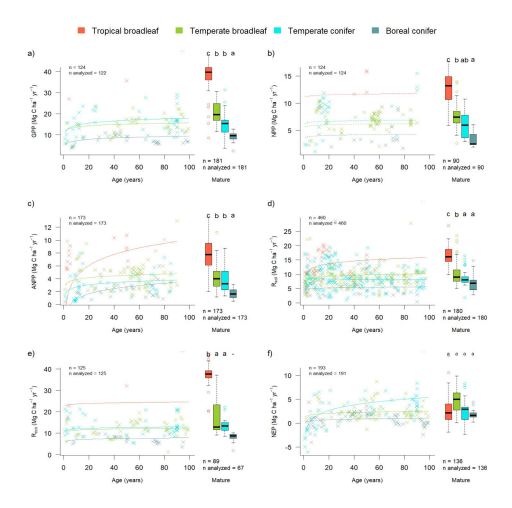


Figure 7 | Age trends and biome differences in some of the major C fluxes: (a) GPP, (b) NPP, (c) ANPP, (d)  $R_{soil}$ , (e)  $R_{eco}$ , and (f) NEP. In each panel, the left scatterplot shows age trends in forests up to 100 years old, as characterized by a linear mixed effects model with fixed effects of age and biome. The fitted line indicates the effect of age on flux (solid lines: significant at p<0.05, dashed lines: non-significant), and non-parallel lines indicate a significant age x biome interaction. The boxplot illustrates distribution across mature forests, with different letters indicating significant differences between biomes. Data from biomes that did not meet the sample size criteria (see Methods) are plotted, but lack regression lines (young forests) or test of differences across biomes (mature forests). Individual figures for each flux with sufficient data, along with maps showing geographic distribution of the data, are given in the Supplement (Figs. S4-S19).

There were sufficient data to assess mature forest biome differences for nine stock variables, and significant differences among biomes were detected for five variables ( $B_{tot}$ ,  $B_{ag}$ ,  $B_{ag-wood}$ ,  $B_{foliage}$ ,  $B_{root-coarse}$ ; Table 319 1). C stocks had less consistent patterns across biomes (Figs. 8, S20-S30). For  $B_{tot}$  and  $B_{ag}$ , tropical 320 broadleaf forests had the highest mean biomass and boreal forests the lowest, with temperate broadleaf and 321 needleleaf ( $B_{aq}$  only) intermediate. However, maximum values for stocks including live or standing woody 322 biomass  $(B_{tot}, B_{ag}, B_{ag-wood}, B_{root}, B_{root_{coarse}}, DW_{tot}, DW_{standing})$  consistently occurred in temperate 323 biomes (Figs. 1, 8, S20-S30). For variables that were disproportionately sampled in such high-biomass forests 324  $(B_{ag-wood}, B_{foliage}, \text{ and } B_{root-coarse}; \text{ disproportionately sampled in the US Pacific Northwest}), temperate$ 325 conifer forests had significantly higher stocks than the other biomes.

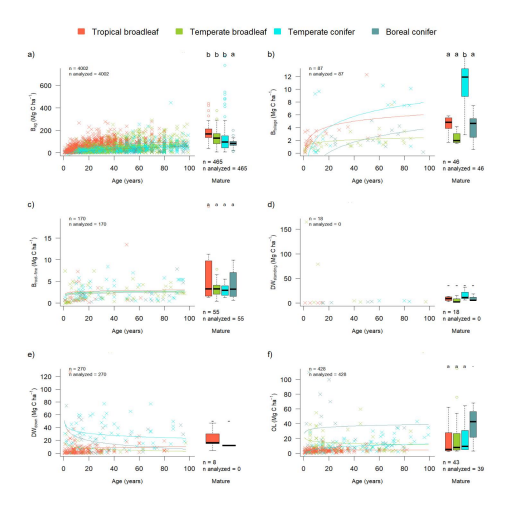


Figure 8 | Age trends and biome differences in some of the major forest C stocks: (a) aboveground biomass, (b) foliage, (c) fine roots, (d) dead wood. In each panel, the left scatterplot shows age trends in forests up to 100 years old, as characterized by a linear mixed effects model with fixed effects of age and biome. The fitted line indicates the effect of age on flux (solid lines: significant at p<0.05, dashed lines: non-significant), and non-parallel lines indicate a significant age x biome interaction. The boxplot illustrates distribution across mature forests, with different letters indicating signifiant differences between biomes. Data from biomes that did not meet the sample size criteria (see Methods) are plotted, but lack regression lines (young forests) or test of differences across biomes (mature forests). Individual figures for each stock with sufficient data, along with maps showing geographic distribution of the data, are given in the Supplement (Figs. S20-S30).

### C cycling in young forests

- <sup>328</sup> C fluxes commonly increased significantly with stand age (Tables 1, S2, Figs. 7, 9, S5-S30). For C contained
- <sup>329</sup> 16 C flux variables with sufficient data for analyses of age trends in young forests (see Methods). Of these,
- ten increased significantly with log10[age]: NEP, GPP, ANPP,  $ANPP_{woody}$ ,  $ANPP_{foliage}$ ,
- $ANPP_{litterfal}$ , BNPP,  $BNPP_{fine}$ ,  $R_{eco}$ , and  $R_{root}$ . The remaining six-NPP,  $ANPP_{stem}$ ,  $ANPP_{branch}$ ,
- $BNPP_{coarse}$ ,  $R_{soil}$ , and  $R_{het-soil}$ -displayed no significant relationship to stand age.

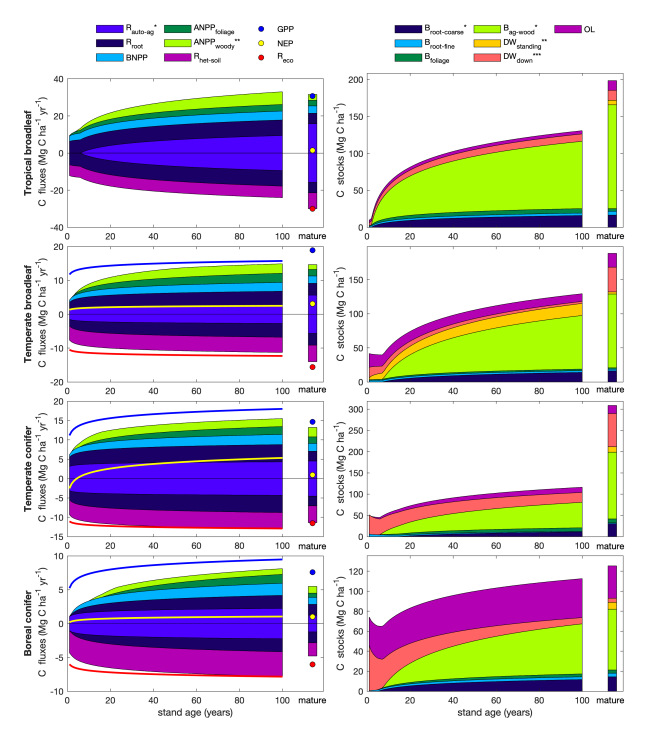


Figure 9 | Age trends in C cycling. Selection of variables for plotting seeks to maximize sample size and broad geographic representation while representing all elements of C cycle. Asterisks indicate variables whose age trends were calculated based on other variables (\* young and mature forests; \*\* young forests only; \*\*\* mature forests only), as follows. For all forests:  $B_{ag-wood} = max(0, B_{ag} - B_{foliage})$ ,  $B_{root-coarse} = max(0, B_{root} - B_{root-fine})$ ,  $DW_{standing} = max(0, DW_{tot} - DW_{down})$ . For tropical forests:  $ANPP_{woody} = max(0, ANPP - ANPP_{foliage})$ ,  $R_{auto-ag} = R_{auto} - Rroot$ , where  $R_{auto} = NPP(1/CUE - 1)$  and CUE = 0.46 (Collati et al. 2020). for non-tropical forests:  $ANPP_{woody} = min(ANPP_{stem}, ANPP_{woody})$ ,  $R_{auto-ag} = R_{eco} - Rsoil$ , . Note that there remain substantial uncertainties as to the functional form of age trends and discrepencies in closure among related variables.

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Differences in C fluxes across biomes typically paralleled those observed for mature forests, with C cycling
    generally most rapid in the tropics and slowest in boreal forests. The single exception was ANPP_{stem}, for
    which temperate broadleaf and conifer forests had flux rates similar to tropical forests. Notably, and in
335
    contrast to the lack of biome differences in NEP for mature forests (Fig. 7), the tendency for temperate
336
    forests to have greater fluxes than boreal forests held for NEP in regrowth forests (tropical forests excluded
337
    because of insufficient data).
338
    "Closure" and internal consistency of the C flux budget was less successful for young than mature forests
330
    (Figs. 9). Summed regression equations for R_{soil-het} and R_{root} were generally very close to R_{soil}. We
340
    calculated R_{auto-aq} as the difference between R_{eco} and R_{soil} (except for tropical forests, which had
    insufficient R_{eco} data), effectively guaranteeing near-closure of the CO<sub>2</sub> efflux (respiration) portion of the
342
    budget (negative values in Figs. 9). In contrast, the CO<sub>2</sub> influx portion of the budget generally did not
343
    "close": the sum of R_{auto} (R_{root} + R_{auto-ag}, as described above) and components of NPP consistently fell
    short of GPP, particularly in in young stands (range across forest types and ages: 0.9-7.6 Mg C ha<sup>-1</sup> yr<sup>-1</sup>).
345
    Moreover, there was not consistent budget closure among the components of NPP, and substantially
346
    different age trends resulting from the sum of components versus total NPP (Figs. 9). Although age trends
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    of young forests often converged towards mature forest averages, there were also some discrepancies between
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    young forest trends and mature forest averages (Figs. 7, 9, S5-S30), most notably including a tendency for
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    higher fluxes in regrowth boreal forests than in their mature counterparts (Fig. 9).
    In terms of C stocks, ten variables (all but standing deadwood, DW_{standing}) had sufficient data to test for
351
    age trends (Table 1, Figs. 8, S20-S30). All of these displayed a significant overall increase with with
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    log10[stand.age]. Age \times biome interactions were also significant for all ten of these C stock variables (Table
353
    S2), with living C stocks tending to accumulate more rapidly during the early stages of forest regrowth in
354
    tropical forests (Figs. 8, S20-S30). In the case of two non-living C stocks (DW_{down} and OL), age \times biome
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    interactions were such that age trends were positive in some biomes and negative in others. Specifically,
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    DW_{down} declined with age in temperate and boreal forests, compared to an increase with age in tropical
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    forests (Figs. 8, S29). Similarly, OL declined slightly with age in temperate broadleaf forests, contrasting an
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    increase in the other three biomes (Figs. 8, S30). Again, there were some discrepencies between young forest
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    trends and mature forests, most notably including generally higher C stocks in mature forests relative to
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    their 100-year counterparts, particularly for temperate conifer forests (again, likely a geographic
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    representation issue?) and, to a lesser extent, tropical broadleaf forests.
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### Discussion

For C v3.0 provided unprecedented coverage of most major variables, yielding a broad picture of C cycling in 364 the world's major forest biomes. Carbon cycling rates generally increased from boreal to tropical regions and 365 with stand age (Figs. 1, S5-S30). Specifically, most C fluxes were highest in tropical forests, intermediate in 366 temperate (broadleaf or conifer) forests, and lowest in boreal forests – a pattern that generally held for 367 regrowth as well as mature forests (Figs. 1, 7-8, 9). The notable exception was mature forest NEP, which, 368 as the difference between GPP and  $R_{eco}$ , was statistically indistinguishable across biomes (Fig. 7f). There was also little directional variation in mean mature forest C stocks (biomass, dead wood, and organic layer) 370 across biomes, although maximum values for the majority of stocks (all including live or standing woody 371 biomass) occurred in temperate biomes (Figs. 1, 3-6, 8). Consistent with theory and previous studies (Fig. 372 1), the majority of flux variables, together with most live biomass pools, increased significantly with stand 373 age (Table 1; Figs. 7-9, S5-S30). Together, these results indicate that, moving from cold to tropical climates and from young to old stands, there is a general acceleration of C cycling, whereas C stocks and NEP of 375 mature forests, which are defined by the differences between in- and out- fluxes, do not vary systematically 376 across biomes. Together, these results refine and expand out understanding of C cycling in mature forests, 377 while providing the first global-scale analysis of age trends in multiple forest C stocks and fluxes (Figs. 9). 378

Our analysis reveals that carbon cycling is most rapid in the tropics and slowest in boreal regions, including

### 379 C cycling across biomes

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C fluxes into (GPP), within (e.g., NPP and its components), and out of (e.g.,  $R_{soil}$ ,  $R_{eco}$ ) the ecosystem. 381 For mature forests, this is consistent with a large body of previous work demonstrating that C fluxes generally decline with latitude (or increase with temperature) on a global scale (e.g., Luyssaert et al 2007, 383 Gillman et al 2015, Li and Xiao 2019, Banbury Morgan et al n.d.). This consistency is not not surprising, 384 particularly given commonality in the data analyzed or used for calibration. The finding that these patterns 385 hold consistently across across numerous fluxes, while unsurprising, is novel to this analysis (but see Banbury Morgan et al n.d. for nine autotrophic fluxes). 387 The notable exception to the pattern of fluxes decreasing from tropical to boreal regions is NEP, which 388 showed no significant differences across biomes, albeit with the highest mean in temperate broadleaf forests (Fig. 7f). Unlike the other C flux variables, NEP does not characterize the rate at which C cycles through 390 the ecosystem, but is the balance between C sequestration (GPP) and respiratory losses  $(R_{eco})$  and 391 represents net CO<sub>2</sub> sequestration (or release) by the ecosystem (Fig. 1). NEP tends to be relatively small in mature forest stands, which accumulate carbon slowly relative to younger stands (discussed further below), if 393 at all (Fig. 1; Luyssaert et al 2008, Amiro et al 2010, Besnard et al 2018). It is therefore unsurprising that 394 there are no pronounced differences across biomes (in agreement with Luyssaert et al 2007), suggesting that variation in NEP of mature forests is controlled less by climate and more by other factors including 396 moderate disturbances (Curtis and Gough 2018) or disequilibrium of  $R_{soil}$  relative to C inputs (e.g., in 397 peatlands where anoxic conditions inhibit decomposition; Wilson et al 2016). The fact that mature temperate broadleaf forests have a higher mean than the other biomes may reflect the fact that most of these 399 forests are older secondary forests that, while classified here as mature, are still accumulating carbon (Curtis 400 and Gough 2018). 401 In contrast to the patterns observed for NEP in mature stands, NEP of stands between 20 and 100 years of 402

age varied across biomes, being lowest in boreal forests, intermediate in temperate broadleaf forests, and

highest in temperate conifer forests (with insufficient data to assess tropical forests; Figs. 7, S5). This is 404 consistent with findings that live biomass accumulation rates (e.g.,  $\Delta B_{aq}$  or  $\Delta B_{tot}$ ) during early secondary succession decrease with latitude (Figs. 7a, S16-S22; Anderson et al 2006, Cook-Patton et al 2020). Note, 406 though, that NEP includes not only  $\Delta B_{tot}$ , but also changes in  $DW_{tot}$ , OL, and soil carbon (not analyzed 407 here), and biome differences in the accumulation rates of these variables have not been detected, in part because these variables do not consistently increase with stand age (Figs. 1, 8, S27-S30, and see discussion 409 below; Cook-Patton et al 2020). 410 For regrowth forests, little is known about cross-biome differences in carbon fluxes, and we are not aware of 411 any previous large-scale comparisons of C fluxes that have been limited to regrowth forests. Thus, this analysis was the first to examine flux trends in regrowth forests across biomes. The observed tendency for 413 young forest fluxes to decrease from tropical to boreal regions paralleled patterns in mature forests (Figs. 7, 414 S5-S19), suggesting that regrowth forests follow latitudinal trends in carbon cycling similar to those of mature forests (e.g., Banbury Morgan et al n.d.). 416 In contrast to C fluxes and biomass accumulation rates in regrowth forests, stocks showed less systematic 417 variation across biomes. For aboveground biomass, which is the variable in ForC with broadest geographical 418 representation, the modest trend of declining biomass from tropical to boreal regions mirrors observations 419 from spaceborne lidar that reveal a decline in aboveground biomass (for all forests, including secondary) with 420 latitude across the N hemisphere (Hu et al 2016). The highest-biomass forests on Earth are, however, found 421 in coastal temperate climates of both the southern and northern hemisphere (Figs. 1, 8a; Keith et al 2009, 422 Smithwick et al 2002, Hu et al 2016). Disproportionate representation of forests in one such region—the US 423 Pacific Northwest-inflated estimates of temperate conifer fluxes and stocks for some variables and was 424 responsible for all of the anomalous results described here (e.g., lack of complete C budget closure, anomalous 425 trend across biomes for  $BNPP_{coarse}$ ). Thus, biome differences should always be interpreted relative to the 426 geographic distribution of sampling, which only rarely covers the majority of forested area within a biome. 427 Whereas aboveground biomass can be remotely sensed (albeit with significant uncertainties; Ploton et al 428 2020) and receives significant research attention, far less is known about geographical variation in deadwood 429 and organic layer (OL) across biomes, which has proved a limitation for C accounting efforts (Pan et al 430 2011). Although these stocks can be important-exceeding 100 Mg C ha<sup>-1</sup> in some stands (Figs. 8, S27-S29). 431 this study is the first to synthesize deadwood data on a global scale (but see Cook-Patton et al 2020 for 432 young forests). Unfortunately, data remain too sparse for statistical comparison across biomes (Figs. 8, 433 S27-S29; but see below for age trends), pointing to a need for more widespread quantification of both standing and downed deadwood. For C coverage of OL stocks is more comprehensive, revealing no significant 435 differences across temperate and tropical biomes, but a tendency towards higher OL in boreal forests, 436 consistent with the idea that proportionally slower decomposition in colder climates results in more buildup of organic matter (Allen et al 2002, Anderson-Teixeira et al 2011). Further research on non-living C stocks in 438 the world's forests will be essential to completing the picture. 439

### 440 Age trends in C cycling

Our study reveals that most C fluxes quickly increase to a plateau as stands age (Figs. 7, 9), consistent with current understanding of age trends in forest C cycling (Fig. 1; e.g., Anderson-Teixeira et al 2013, Amiro et al 2010, Magnani et al 2007). While limited records in very young (i.e., <5 year old) stands resulted in poor resolution of the earliest phases of this increase for many variables (sometimes detecting no age trend; Table

1), any autotrophic C flux (e.g., GPP, NPP and its components,  $R_{auto}$ ) would be minimal immediately 445 following a stand-clearing disturbance. These would be expected to increase rapidly with the most metabolically active components of biomass, foliage and fine roots, which also increase rapidly with stand age 447 (Fig. 8). In contrast, soil heterotrophic respiration  $(R_{het-soil})$  and total soil respiration  $(R_{soil})$  are expected 448 to be non-zero following stand-clearing disturbance, although these may decrease with a reduction of root respiration ( $R_{soil}$  only) and C exudates or increase in response to an influx of dead roots and litter 450 (Ribeiro-Kumara et al 2020, Maurer et al 2016, Bond-Lamberty et al 2004). In this study, we detect no 451 significant age trends in either variable. Notably, net carbon sequestration (NEP) exhibits an overall increase with age across the first 100 years of stand development, with more pronounced patterns in temperate than boreal forests (Fig. 7). This finding is 454 largely consistent with, but built from a far larger dataset than, previous studies showing an increase in 455 NEP across relatively young stand ages (Pregitzer and Euskirchen 2004, Baldocchi et al 2001, Luyssaert et al 2008). However, NEP has been observed to decline from intermediate to old stands (???, Luyssaert et al 457 2008), and the NEP estimated by our model for 100-year-old temperate conifer stands (~5 Mg C ha<sup>-1</sup> yr<sup>-1</sup>) 458 exceeds the mean of mature forests in the same biome (0.7 Mg C ha<sup>-1</sup> yr<sup>-1</sup>; Fig. 5). A decrease in NEP is consistent with the observed deceleration of biomass accumulation as stands age, although both biomass and 460 non-living C stocks will often continue to increase well beyond the 100-yr threshold used here to delimit 461 young and mature stands (Luyssaert et al 2008, McGarvey et al 2014, Lichstein et al 2009). 462 In terms of stocks, our study reveals consistent increases in live biomass stocks with stand age-a pattern that 463 is well-known and expected (e.g., Lichstein et al 2009, Yang et al 2011)—and more variable age trends in 464 deadwood and OL. The latter are particularly sensitive to the type of disturbance, where disturbances that 465 remove most organic material (e.g., logging, agriculture) result in negligible deadwood in young stands, 466 followed by a buildup over time (tropical stands in Fig. 8; e.g., Vargas et al 2008). In contrast, natural 467 disturbances (e.g., fire, drought) can produce large amounts of deadwood (mostly  $DW_{standing}$ ) that slowly 468 decomposes as the stand recovers, resulting in declines across young stand ages (e.g., temperate and boreal 469 stands in Fig. 8; e.g., Carmona et al 2002). Again, further study and synthesis of non-living C stocks across 470 biomes and stand ages will be valuable to giving a more comprehensive picture. 471

#### 2 C variable coverage and budget closure

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confidence that our overall reported mature forest means provide useful baselines for analysis – with the 474 caveats that they are unlikely to be accurate representations of C cycling for any particular forest, and that 475 these sample means almost certainly do not represent true biome means (particularly for temperate conifer forests where high-biomass stands are over-represented in ForC). 477 In this analysis, the C cycle budgets for mature forests (Figs. 3-6) generally "close"—that is, the sums of 478 component variables do not differ from the larger fluxes by more than one standard deviation. On the one 479 hand, this reflects the general fact that ecosystem-scale measurements tend to close the C budget more easily and consistently than, for example, for energy balance (Stoy et al 2013). On the other, however, For C 481 derives data from multiple heterogeneous sources, and standard deviations within each biome are high; as a 482 result, the standard for C closure is relatively loose (c.f. Houghton 2020). The one instance where the C budgets doesn't close is likely due to differences in the representation of forest types (i.e., disproportionate 484 representation of US Pacific NW for  $B_{root-coarse}$  relative to  $B_{root}$ ; Fig. 5) rather than issues of 485

The large number of C cycle variables covered by ForC, and the general consistency among them, provide

consistent picture of C cycling within biomes for mature forests. This is an important and useful test, because it allows for consistency checks within the C cycle, for example leveraging separate and 488 independently-measured fluxes to constrain errors in another (Phillips et al 2017, Williams et al 2014, 489 Harmon et al 2011), or producing internally consistent global data products (Wang et al 2018). In contrast, age trends for young forests generally remain less clearly defined, in large part because their data records remain somewhat sparse for most variables (i.e., have low representation of different geographical 492 regions for any given age). While this analysis provides a first analysis of age trends in forest C cycling for 493 multiple variables at a global scale, improved resolution of these trends will require additional data. There are of course notable holes in the ForC variable coverage that limit the scope of our inferences here. 495 Notably, For C currently has sparse—if any—coverage of fluxes to herbivores and higher consumers, along with 496 the woody mortality  $(M_{woody})$  and dead wood stocks (Table 1, Figs. S27-S29). For C does not include soil 497 carbon, which is covered by other efforts (e.g., Köchy et al 2015). For C is not intended to replace databases that are specialized for particular parts of the C cycle analyses, e.g., aboveground biomass (Spawn et al 499 2020), land-atmosphere fluxes (Baldocchi et al 2001), soil respiration (Jian et al 2020), or the human 500 footprint in global forests (Magnani et al 2007). 501 Importantly, For C and the analyses presented here cover the forests that have received research attention, 502 which are not a representative sample of the world's existing forests-geographically or in terms of human 503 impacts (Martin et al 2012). Geographically, all variables are poorly covered in Africa and Siberia (Fig. 2), a 504 common problem in the carbon-cycle community (Xu and Shang 2016, Schimel et al 2015). In terms of 505 human impacts, research efforts tend to focus on interior forest ecosystems (Martin et al 2012), often in 506 permanently protected areas (e.g., Davies et al 2021). Studies of regrowth forests tend to focus on sites 507 where recurring anthropogenic disturbance is not a confounding factor. Yet, fragmentation and degradation 508 impact a large and growing proportion of Earth's forests (FAO and UNEP 2020). Fragmentation and the 509 creation of edges strongly impacts forest C cycling (e.g., Chaplin-Kramer et al 2015, Remy et al 2016, 510 Reinmann and Hutvra 2017, Smith et al 2019, Reinmann et al 2020, Ordway and Asner 2020). Partial 511 logging and other forms of non- stand clearing anthropogenic disturbance also strongly impact forest C 512 cycling (e.g., Huang and Asner 2010, Piponiot et al 2016) but are under-studied (Sist et al 2015) and excluded from this analysis. Fragmented and degraded forests do not fit the idealized conceptual framework 514 around which this review is structured (Fig. 1), yet their representation in models, sustainability assessments, 515 and C accounting systems is critical to accurate accounting of C cycling in Earth's forests (e.g., Huang and Asner 2010, Reinmann and Hutyra 2017, Smith et al 2019, Piponiot et al 2019). Finally, plantation forests 517 account for approximately 3% of Earth's forests (FAO and UNEP 2020) but are not included in this analysis. 518 While it is known that these tend to accumulate biomass faster than naturally regenerating forests (Anderson et al 2006, Bonner et al 2013), their global scale C cycling patterns remain less clearly understood (c.f. 520 Cook-Patton et al 2020). Additional research and synthesis are needed to fill these critical gaps in our 521 understanding of forest C cycling.

methodological accuracy. The overall high degree of closure implies that ForC gives an at least roughly

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## Relevance for climate change prediction and mitigation

The future of forest C cycling (Song *et al* 2019) will shape trends in atmospheric CO<sub>2</sub> and the course of climate change (Schimel *et al* 2015). Our findings, and more generally the data contained in *ForC* and summarized here, can help to meet two major challenges.

First, improved representation of forest C cycling in models is essential to improving predictions of the future course of climate change, for the simple reason that by definition future projections extend our existing observations and understanding to conditions that do not currently exist on Earth (McDowell et al 2018, 529 Bonan and Doney 2018, Gustafson et al 2018). To ensure that models are giving the right answers for the 530 right reasons (Sulman et al 2018), it is important to benchmark against multiple components of the C cycle that are internally consistent with each other (Collier et al 2018, Wang et al 2018). For C's tens of thousands 532 of records are readily available in a standardized format, and our analyses here indicate that their internal 533 consistency is reasonably high. Integration of ForC with models will be valuable to improving the accuracy and reliability of models (Fer et al 2021). 535 Second, ForC can serve as a pipeline through which information can flow efficiently from forest researchers to decision-makers working to implement forest conservation strategies at global, national, or landscape scales.

Second, For C can serve as a pipeline through which information can flow efficiently from forest researchers to
decision-makers working to implement forest conservation strategies at global, national, or landscape scales.
This is already happening: For C has contributed to updating the IPCC guidelines for carbon accounting in
forests (IPCC 2019, Requena Suarez et al 2019), mapping C accumulation potential from natural forest
regrowth globally (Cook-Patton et al 2020), and informing ecosystem conservation priorities (Goldstein et al
2020).

It is also interesting to consider the complementary utility of global-scale but spatially discontinuous 542 databases such as ForC and remote wall-to-wall remote sensing products. The latter provide unparalleled 543 insight into aboveground carbon stocks, but less constraint on belowground stocks or carbon fluxes in general (Bond-Lamberty et al 2016, Anav et al 2015). Combining observational data and remote observations may 545 provide a much more comprehensive and accurate picture of global forest C cycling, particularly when used 546 in formal data assimilation systems (Konings et al 2019, Liu et al 2018). Biomass is the largest C stock in 547 most forests, and most of the emphasis has traditionally been on this variable. Remote-sensing driven 548 aboveground biomass estimates (e.g., Saatchi et al 2011), calibrated based on high-quality ground-based data 549 (Schepaschenko et al 2019, Chave et al 2019), provide the most promising approach, but significant 550 uncertainties remain (Ploton et al 2020). Note, however, that factors such as stand age and disturbance 551 history are difficult, if not impossible, to detect remotely, and can only be characterized for very recent 552 decades (Hansen et al 2013, Song et al 2018, Curtis et al 2018). Ground-based data such as ForC are 553 therefore valuable in defining age-based trajectories in biomass, as in Cook-Patton  $et\ al\ (2020)$ , and thus 554 constraining variables such as carbon sink potential (Luyssaert et al 2008). 555

Efforts such as the Global Carbon Project (Friedlingstein et al 2019) and NASA's Carbon Monitoring

System (Liu et al 2018) typically compute respiration as residuals of all other terms (Bond-Lamberty et al

2016, Harmon et al 2011). This means that the errors on respiration outputs are likely to be large and

certainly poorly constrained, offering a unique opportunity for databases such as ForC and SRDB (Jian et al

2020) to provide observational benchmarks. For example, Konings et al (2019) produced a unique top-down

estimate of global heterotrophic respiration that can both be compared with extant bottom-up estimates

(Bond-Lamberty 2018) and used as an internal consistency check on other parts of the carbon cycle (Phillips

et al 2017).

#### Conclusions

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As climate change accelerates, understanding and managing the carbon dynamics of forests—notably including dynamics and fluxes that cannot be observed by satellites—is critical to forecasting, mitigation, and

adaptation. The C data in ForC, as summarized here, will be valuable to these efforts. Notably, the fact that tropical forests tend to have both the highest rates of C sequestration in young stands (Fig. 8; Cook-Patton et al 2020), fueled by their generally high C flux rates (Table 1; Fig. 7), and the highest mean biomass (Fig. 570 8; Table 1; Hu et al 2016, Jian et al 2020) reinforces the concept that conservation and restoration of these 571 forests is a priority for climate change mitigation, along with high-biomass old-growth temperate stands (Grassi et al 2017, Goldstein et al 2020). It is also important to note the trade-off in climate mitigation 573 potential of restoration of young forests, with high rates of CO<sub>2</sub> sequestration (NEP; Cook-Patton et al 574 2020), versus conservation and management of mature forests, with low NEP but high C stocks that could not be recovered on a time scale relevant to climate change mitigation (Goldstein et al 2020). Generally 576 speaking, the conservation of mature forests will yield greater climate benefits (Anderson-Teixeira and 577 DeLucia 2011), but both approaches are critical to avoiding catastrophic climate change (IPCC 2018).

#### 579 Citations to add

580 Harris *et al* (2021)

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# Data availability statement

Materials required to fully reproduce these analyses, including data, R scripts, and image files, are archived in Zenodo (DOI: TBD]. Data, scripts, and results presented here are also available through the open-access ForC GitHub repository (https://github.com/forc-db/ForC), where many will be updated as the database develops.

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