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Cross-biome synthesis of source versus sink limits to tree growth

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3	Title: Cross-biome synthesis of source versus sink limits to tree growth
4	Authors: Antoine Cabon ¹ *, Steven A. Kannenberg ¹ , Altaf Arain ^{2,3} , Flurin Babst ^{4,5} , Dennis
5	Baldocchi ⁶ , Soumaya Belmecheri ⁵ , Nicolas Delpierre ^{7,8} , Rossella Guerrieri ⁹ , Justin T.
6	Maxwell ¹⁰ , Shawn McKenzie ^{2,3} , Frederick C. Meinzer ¹¹ , David J. Moore ⁴ , Christoforos
7	Pappas ^{12,13} , Adrian V. Rocha ¹⁴ , Paul Szejner ¹⁵ , Masahito Ueyama ¹⁶ , Danielle Ulrich ¹⁷ , Caroline
8	Vincke ¹⁸ , Steven L. Voelker ¹⁹ , Jingshu Wei ²⁰ , David Woodruff ¹¹ , William R. L. Anderegg ¹
9	Affiliations:
10	¹ University of Utah, School of Biological Sciences, Salt Lake City UT, USA
11	² McMaster University, McMaster Centre for Climate Change, Hamilton, ON L8S4K1,
12	Canada
13	³ McMaster University, School of Earth, Environment and Society, Hamilton, ON L8S4K1,
14	Canada
15	⁴ School of Natural Resources and the Environment, University of Arizona, Tucson, AZ,
16	USA
17	⁵ Laboratory of Tree-Ring Research, University of Arizona, Tucson, AZ, USA
18	⁶ Department of Environmental Science, Policy and Management, University of California,
19	Berkeley, CA, USA
20	⁷ Université Paris-Saclay, CNRS, AgroParisTech, Ecologie Systématique et Evolution,
21	91405, Orsay, France.
22	⁸ Institut Universitaire de France (IUF), France
23	⁹ DISTAL, Alma Mater Studiorum, University of Bologna, Bologna, Italy
24	¹⁰ Department of Geography, Indiana University, Bloomington, IN, USA
25	¹¹ USDA Forest Service, Pacific Northwest Research Station, Corvallis, OR, USA

26 27	¹² Centre d'étude de la forêt, Université du Québec à Montréal, C.P. 8888, Succursale Centre- ville, Montréal, QC, H3C 3P8, Canada
28 29	¹³ Département Science et Technologie, Téluq, Université du Québec, 5800 rue Saint-Denis, Bureau 1105, Montréal, OC, H2S 3L5, Canada
30	¹⁴ University of Notre Dame, Department of Biological Sciences, Notre Dame, IN, USA
31 32	¹⁵ Geology Institute, National Autonomous University of Mexico, Coyoacán, CDMX, Mexico
33 34	¹⁶ Graduate School of Life and Environmental Sciences, Osaka Prefecture University, 599- 8531, Japan
35	¹⁷ Department of Ecology, Montana State University, Bozeman, MT, USA
36	¹⁸ Earth and Life Institute, Université Catholique de Louvain, Louvain-la-Neuve, Belgium
37 38	¹⁹ College of Forest Resources and Environmental Science, Michigan Technological University, Houghton, MI, United States
39 40	²⁰ Department of Ecology, W. Szafer Institute of Botany, Polish Academy of Sciences, ul. Lubicz 46, 31-512 Kraków, Poland
41	*Corresponding author. Email: antoine.cabon@utah.edu
42 43	Abstract. Uncertainties surrounding tree carbon allocation to growth are a major limitation to
44	projections of forest carbon sequestration and response to climate change. The prevalence and
45	extent to which carbon assimilation (source) or cambial activity (sink) mediate wood production
46	is fundamentally important and remains elusive. Here we quantify source-sink relations across
47	biomes by combining eddy-covariance gross primary production with extensive on-site and
48	regional tree-ring observations. We find widespread temporal decoupling between carbon
49	assimilation and tree growth, underpinned by contrasting climatic sensitivities of these two
50	processes. We elucidate substantial differences in assimilation-growth decoupling between
51	angiosperms and gymnosperms and stronger decoupling with canopy closure, aridity, and
52 53	to be increasingly prominent under global change.

- **One-Sentence Summary**: Cross-biome correlation analysis shows pervasive decoupling
- 55 between carbon assimilation and tree growth.

57 Main Text:

58 Forest ecosystems currently constitute a net carbon (C) sink that offsets around 25% of 59 yearly anthropogenic C emissions, thus actively mitigating climate change (1). C allocation to 60 aboveground wood biomass is the largest contributor to vegetation C storage over climate-61 relevant time scales. However, wood C allocation is poorly understood and is a major uncertainty 62 for projections of future forests' C storage potential (2). The common representation of wood 63 growth as a linear function of C assimilation has been identified as a major structural limitation 64 of current vegetation models (3, 4). The development of improved C allocation schemes 65 currently lacks a solid empirical and mechanistic basis (5). Thus, there is an urgent need to 66 illuminate the relationship between C assimilation and tree growth.

67 A fundamental debate revolves around the degree to which C assimilation via 68 photosynthesis (source limitation) versus direct environmental limitations to cambial cell 69 development (sink limitation) controls wood growth (6). As reflected by C allocation schemes in 70 the vast majority of vegetation models, source limitation has been the dominant paradigm for 71 decades (4). Yet, a growing body of literature indicates that cambial activity is typically more 72 sensitive than photosynthesis to a range of environmental conditions, including low water 73 availability, temperature, and nutrient availability (7-11). The prevalence of source vs sink 74 limitations to tree growth has far-reaching implications for forest dynamics under climate 75 change, because these processes will likely respond differently to global change (6-9), potentially shifting C allocation away from the stem. Substantial indirect evidence supports the 76 77 hypothesis that C sink limitations may be particularly important in cold, dry, and late-78 successional forests. For example, elevated non-structural C (e.g., starch and sugars) 79 concentrations are frequently observed in colder environments or during drought (8, 12). 80 Additionally, Free Air CO₂ Enrichment (FACE) experiments tends to show that increasing [CO₂] 81 improves tree growth in early-stage forests but often not in mature forests, perhaps because of 82 stronger nutrient limitations (13-15). But the relatively small scale and replication of FACE 83 experiments, especially in mature forests, prevents general conclusions regarding the linkage 84 between C source and sink dynamics in trees.

Co-located assessments of gross primary productivity (GPP) and tree growth theoretically
 enable the evaluation of the coupling between tree C assimilation and growth increment. Past
 studies adopting such an approach were nevertheless limited by dataset size (site number ≤5) and

88 vielded contrasting findings, with no clear explanation of observed differences (16–21). The 89 advent of large-scale, long-term networks of flux towers measuring C exchange across a diverse 90 assemblage of biomes, in combination with a growing number of both on-site and global tree-91 ring datasets, opens new opportunities to characterize C source-sink relationships at larger 92 temporal and spatial scales. Here, we compile a new dataset comprising GPP records at 78 forest 93 flux sites (Table S1), together with on-site tree ring width chronologies at a subset of 31 sites 94 (RW_{on-site}), as well as 1800 nearby regional ring width chronologies (RW_{region}). GPP and RW 95 records were detrended, in order to remove low frequency signals (e.g., stand structure, tree age 96 and size), and aggregated such that records were representative of year-to year variations of 97 stand C assimilation and aboveground woody growth, respectively (22). This C assimilation and 98 tree growth dataset extends across most of Europe and North America, encompassing a variety 99 of forested biomes from semi-arid to boreal, and representing both angiosperm and gymnosperm 100 tree species (Fig. 1, Fig. S1, Table S2). Leveraging this dataset, we (i) quantify the strength, (ii) 101 identify the seasonality, and (iii) explore environmental drivers of tree C source–sink 102 relationships across biomes.

103 We first characterized C source and sink relationships at the regional scale by statistically 104 accounting for the decrease of the correlation between GPP and RW_{region} (r_{region}) with increasing 105 geographic and climatic distances, as well as with an index of species dissimilarity between sites 106 (22) (Fig. S2). As expected from reported tree growth synchrony over large distances (23) we 107 observed sustained correlations up to ~500 km. We thus built on this widespread ecological 108 feature to derive robust regional estimates of tree C assimilation and growth correlation, r_{D=0}, for 109 theoretical co-located sites of identical climate and species composition (i.e., spatial distance, 110 climatic distance and species dissimilarity of 0), integrating over multiple timescales. We then 111 complemented regional-scale analyses with paired GPP and on-site tree-ring correlations (ron-site, 112 see annual GPP and RW series in **Fig. S3**). The latter dataset has a lower sample size compared 113 to the regional network but is model-free and therefore reduces the risk of methodological 114 artifacts.

Both on-site and regional correlations showed an overall weak association between tree C
assimilation and growth, r_{on-site} and r_{D=0} reaching maxima of 0.26 and 0.38, respectively (Fig 2A–
B). The observed difference between on-site and regional estimates could be offset by setting
species dissimilarity to the average encountered for RW_{on-site}, resulting in a maximum regional

119 correlation of 0.27 (22). RW_{region} observations partially build on the international tree ring data 120 bank, where sampling is often biased towards dominant and climate sensitive trees (24).

121 However, we find that this is unlikely an issue here, as dominant trees account for most of stand

122 GPP and we statistically corrected for differences in climate (22). Overall, similar regional and

on-site results show the suitability of regional RW data to quantify local GPP-RW correlations

124 and broad agreement between the two approaches, which both suggest a substantial decoupling

between C assimilation and tree growth across multiple biomes.

123

126 On-site and regional GPP-RW correlations exhibited a similar temporal structure (22), 127 with correlation magnitude increasing with the length of the GPP integration period and 128 maximum correlations being found at the 10- and 12-months scales for $r_{on-site}$ and r_{region} . 129 respectively (Fig. 2). This supports the often-implicit assumption that annual tree-ring 130 increments are most strongly related to annual carbon assimilation (21). Overall, RW was best 131 correlated to GPP integrated over the period spanning previous Sep or Nov to current Aug, 132 consistent with a previous study (20), indicating a short temporal lag between C assimilation and 133 tree growth. This result suggests that, despite estimated low C source limitation of tree growth 134 overall, excess photosynthates are stored over winter following radial growth cessation and 135 allocated to the next year's growth. This phenomenon is often cited as a potential explanation for 136 delayed climatic effects on tree growth and growth auto-correlation (25, 26). Analysis of multi-137 year trends (Table S4) nevertheless indicates weak association of RW and GPP at this scale, 138 contrary to the hypothesis that C storage might lead to the convergence of tree growth and C 139 assimilation over the long term (27).

140 We found large spatial variations in the strength of GPP-RW coupling (Fig. 3). 141 Weighted deciles of maximum on-site r ranged from -0.08 to 0.60, consistent with previously 142 reported values (16–21). These spatial variations imply a range of source vs. sink limitations. We 143 estimate that because of approximations and measurement errors, RW–GPP correlations between 144 0.7 and 0.9 would be expected under strong source control of tree growth (22). The high end of the observed correlation range ($0.6 \le r_{on-site} \le 0.9$: 10% of observations) thus appears reflective of 145 146 substantial source limitation of tree growth at the corresponding sites, whereas the majority of 147 sites display evidence consistent with sink limitations. We did not observe a biome effect on on-148 site correlations but regional-scale r was significantly related to several environmental factors 149 (Fig. 3B). Specifically, gymnosperm proportion had a positive effect on current year r_{region} but a

150 negative one on previous year r_{region}, suggesting that gymnosperm growth relies more directly on 151 current and less on previous year C assimilation than angiosperms, reflecting fundamental 152 physiological differences between these two clades. A small but positive effect of species 153 richness on r_{region} suggests a link between species diversity and C use efficiency (i.e., the ratio 154 between net and gross primary production), which may arise as a result of increased 155 complementarity with structural and functional heterogeneity (28). Decreasing r_{region} with 156 increasing leaf area index indicates that closed canopy forests, which under a given climate tend 157 to be older and more nutrient-limited than open canopy forests, are prone to stronger decoupling 158 between C source and sink activity. This result agrees with the observations that CO₂ growth 159 fertilization tends to fade in older, nutrient-limited forests (15). Last, r_{region} was found to be 160 positively related to site temperature and water availability, consistent with known biophysical 161 controls of cambial activity and the ensuing prediction that sink limitations are stronger under 162 colder and drier conditions (6-9). These combined results draw a clear picture that increasing 163 resource limitation, aridity, and low temperatures promote C source-sink decoupling across a 164 broad range of biomes.

165 Finally, decoupled C assimilation and tree growth was further revealed by diverging 166 climate sensitivities of these two processes (22) (Fig. 4). As anticipated from C assimilation and 167 wood formation literature, GPP and RW both responded positively to temperature and water 168 availability but were weakly correlated with photosynthetically active radiation (hereafter 169 radiation) (29, 30). However, their seasonal variability differed markedly, indicating that 170 fundamentally different physiological processes may limit C assimilation and tree growth. GPP 171 responded mostly to spring and fall temperatures, as well as to summer water availability, 172 suggesting an important role of temperature-triggered leaf phenology controlling annual GPP 173 (31). In contrast, RW appeared to be most strongly related to year-round water availability, with 174 a weak positive temperature effect peaking in summer. This agrees with previous observations 175 that tree growth is primarily and increasingly water-limited in the study regions (29) and is 176 consistent with the central role of cell turgor in controlling cambial cell division and expansion 177 (7, 11). Overall, this analysis shows the large but contrasting climate sensitivity of the tree 178 growth and photosynthesis proxies used here. This is contrary to the expectation that RW and 179 GPP would have weaker but similar climate sensitivity if low RW–GPP were due primarily to 180 large measurement errors. These results instead strongly suggest that weak control of C

assimilation over tree growth is underpinned by fundamentally contrasting source and sink
processes with diverging environmental sensitivities (6).

183 Taken together, our results provide consistent evidence for the pervasive influence of 184 non-photosynthetic processes on tree radial growth. This conclusion has major implications in 185 terms of projections of forest dynamics and feedbacks with the global C cycle and climate 186 change, as most global vegetation models essentially simulate forest productivity and C 187 sequestration as a linear function of C assimilation (3, 4). Because sink processes are relatively 188 more sensitive to water availability than temperature constrains compared to C assimilation (Fig. 189 4) and are not directly dependent on atmospheric [CO₂], unaccounted for and widespread sink 190 limitations could lead to overestimating the positive effect of warming and CO₂ fertilization 191 while underestimating the negative effect of increasing water stress on forest productivity. 192 Overall, accounting for sink limitations of tree growth may lower projections of future forest C 193 sequestration in many regions and could thus potentially compromise forests' potential for 194 climate change mitigation. Based on these considerations, our results underscore that 195 incorporation of sink-limited carbon allocation schemes in global vegetation models is urgently 196 needed (3, 4).

197 Our results nevertheless indicate a certain degree of interaction between C source and 198 sink activities, as suggested by the weak but significantly positive correlations observed between 199 GPP and RW, as well as their temporal and spatial variations. Such dynamic coupling between C 200 assimilation and tree growth potentially reconciles contrasting observations of the prevalence of 201 source vs sink limitations (15) and provides a bridge between current source-centered 202 representations of tree growth and sink-driven schemes. Variations in the prevalence of source vs 203 sink limitations to tree growth further highlights the importance of understanding their drivers 204 (5). Here we show that across biomes, the occurrence of sink limitations is highly consistent with 205 known biophysical controls of cambial cell division, notably turgor-driven growth. Because 206 turgor is a central mechanism of growth across scales and has a large potential for both 207 integration of several relevant processes and parameter-parsimonious upscaling (32), the turgor-208 driven growth framework appears to be a promising way forward to developing mechanistic 209 sink-limited schemes in vegetation models.

210 Key remaining uncertainties include whether our results can be generalized to other 211 biomes such as tropical forests, which are central to the global C cycle, and quantifying the

- 212 dynamic nature of source and sink interactions. Likewise, characterizing the degree of C source
- and sink decoupling at decadal to centennial scales is relevant regarding climate change but
- 214 currently remains elusive due to the temporal depth of C assimilation measurements. Source-
- sink decoupling over both short and longer timescales implies less C limitation of tree growth.
- 216 Weak C limitation of tree growth under certain conditions nonetheless raises the question of the
- 217 fate of excess C. Closing trees' C budget and elucidating drivers of C allocation to different
- sinks, specifically stem vs underground growth and C storage thus emerges as a critical way
- 219 forward (*14*).
- 220

Fig 1. Spatial distribution of gross primary production (GPP) and regional ring width (RW_{region}) sites used in this study. RW_{region} sites are indicated by crosses and GPP sites by circle. The number of RW_{region} site-year observations associated with each flux tower is denoted by circle size. GPP sites that further include on-site RW are colored in yellow, and in red otherwise.

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Fig 2. Temporal structure of gross primary productivity (GPP) vs. ring width (RW) correlations. (A) Seasonal on-site correlations ($r_{on-site}$). Each cell corresponds to the average correlation calculated between on-site RW and GPP summed over a time-period defined by a window onset (from previous year Jan to current Dec) and length (from 1 to 12 months). (B) Regional-based estimates of null distance correlations ($r_{D=0}$) modelled by Eq. S1 (see Fig. S2 for an illustration of the 12-months case from current year Jan). Significant correlation values are displayed on top of corresponding cells (lightface: p-value < 0.05; bold: p-value < 0.01).



climatic water deficit – MACWD, mean annual temperature – MAT) except for that of leaf area
index (LAI) on current year correlations (ns).

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Fig 4. Gross primary production (GPP) and regional ring width (RW) climatic sensitivity. Climate-corrected partial correlations between GPP and regional RW and three climate variables (from the top to the bottom: mean temperature – T_{mean} , Palmer's drought severity index – PDSI, and radiation – Rad) at the three-monthly scale, over the period 1990–2015 (to the extent of series span). Error bars correspond to SE. Statistical significance of estimated coefficients is denoted as: * – p<0.05; ** – p<0.01; *** – p<0.001.

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- 316 Methodology: AC, WA, SK
- 317 Data contributions: All co-authors
- 318 Investigation: AC, WA, SK
- 319 Visualization: AC
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- 324
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328 Supplementary Materials

- 329 Materials and Methods
- 330 Fig S1 to S5
- Tables S1 to S4
- 332 References (52–67)
- 333
- 334
- 335