

Climate Effects on Belowground Tea Litter Decomposition Depend on Ecosystem and Organic Matter Types in Global Wetlands

Stacey M. Trevathan-Tackett,* Sebastian Kepfer-Rojas, Martino Malerba, Peter I. Macreadie, Ika Djukic, Junbin Zhao, Erica B. Young, Paul H. York, Shin-Cheng Yeh, Yanmei Xiong, Gidon Winters, Danielle Whitlock, Carolyn A. Weaver, Anne Watson, Inger Visby, Jacek Tylkowski, Allison Trethowan, Scott Tiegs, Ben Taylor, Jozef Szpikowski, Grażyna Szpikowska, Victoria L. Strickland, Normunds Stivrins, Ana I. Sousa, Sutinee Sinutok, Whitney A. Scheffel, Rui Santos, Jonathan Sanderman, Salvador Sánchez-Carrillo, Joan-Albert Sanchez-Cabeza, Krzysztof G. Rymer, Ana Carolina Ruiz-Fernandez, Bjorn J. M. Robroek, Tessa Roberts, Aurora M. Ricart, Laura K. Reynolds, Grzegorz Rachlewicz, Anchana Prathap, Andrew J. Pinsonneault, Elise Pendall, Richard Payne, Ilze Ozola, Cody Onufrock, Anne Ola, Steven F. Oberbauer, Aroloye O. Numbere, Alyssa B. Novak, Joanna Norkko, Alf Norkko, Thomas J. Mozdzer, Pam Morgan, Diana I. Montemayor, Charles W. Martin, Sparkle L. Malone, Maciej Major, Mikołaj Majewski, Carolyn J. Lundquist, Catherine E. Lovelock, Songlin Liu, Hsing-Juh Lin, Ana Lillebo, Jinquan Li, John S. Kominoski, Anzar Ahmad Khuroo, Jeffrey J. Kelleway, Kristin I. Jinks, Daniel Jerónimo, Christopher Janousek, Emma L. Jackson, Oscar Iribarne, Torrance Hanley, Maroof Hamid, Arjun Gupta, Rafael D. Guariento, Ieva Grudzinska, Anderson da Rocha Gripp, María A. González Sagrario, Laura M. Garrison, Karine Gagnon, Esperança Gacia, Marco Fusi, Lachlan Farrington, Jenny Farmer, Francisco de Assis Esteves, Mauricio Escapa, Monika Domańska, André T. C. Dias, Carmen B. de los Santos, Daniele Daffonchio, Paweł M. Czyryca, Rod M. Connolly, Alexander Cobb, Maria Chudzińska, Bart Christiaen, Peter Chiffard, Sara Castelar, Luciana S. Carneiro, José Gilberto Cardoso-Mohedano, Megan Camden, Adriano Caliman, Richard H. Bulmer, Jennifer Bowen, Christoffer Boström, Susana Bernal, John A. Berges, Juan C. Benavides, Savanna C. Barry, Juha M. Alatalo, Alia N. Al-Haj, and Maria Fernanda Adame



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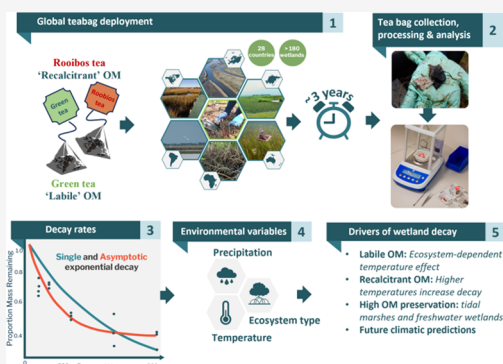


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ABSTRACT: Patchy global data on belowground litter decomposition dynamics limit our capacity to discern the drivers of carbon preservation and storage across inland and coastal wetlands. We performed a global, multiyear study in over 180 wetlands across 28 countries and 8 macroclimates using standardized litter as measures of “recalcitrant” (rooibos tea) and “labile” (green tea) organic matter (OM) decomposition. Freshwater wetlands and tidal marshes had the highest tea mass remaining, indicating a greater potential for carbon preservation in these ecosystems. Recalcitrant OM decomposition increased with elevated temperatures throughout the decay period, e.g., increase from 10 to 20 °C corresponded to a 1.46-fold increase in the recalcitrant OM decay rate constant. The effect of elevated temperature on labile OM breakdown continued...



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was ecosystem-dependent, with tidally influenced wetlands showing limited effects of temperature compared with freshwater wetlands. Based on climatic projections, by 2050 wetland decay constants will increase by 1.8% for labile and 3.1% for recalcitrant OM. Our study highlights the potential for reduction in belowground OM in coastal and inland wetlands under increased warming, but the extent and direction of this effect at a large scale is dependent on ecosystem and OM characteristics. Understanding local versus global drivers is necessary to resolve ecosystem influences on carbon preservation in wetlands.

KEYWORDS: blue carbon, macroclimate, *TeaComposition* H₂O, tea bags, teal carbon

1. INTRODUCTION

Inland and coastal wetland ecosystems, including a range of freshwater and saline ecosystems with different inundation regimes,¹ have a high capacity for soil carbon storage due to anaerobic waterlogged soils and the accumulation of internal and external carbon sources.^{2,3} Within these ecosystems, the vast majority of carbon is stored belowground (e.g., 62–99% in coastal wetlands and 50–93% in freshwater wetlands).^{4–6} The development of belowground carbon pools is largely dependent on inputs of carbon into the system and the microbial processes that moderate the decomposition of carbon-rich organic matter (OM). In addition to being fundamentally important for soil fertility and supporting biodiversity,⁷ the decomposition process in wetland soils is vital for soil carbon (trans)formation and sequestration in wetlands,⁸ and thus the global carbon cycle itself.^{9,10}

Identifying the drivers of wetland litter decomposition processes is essential for predicting feedbacks to environmental and climate change. Elevated temperature can enhance OM breakdown through increased microbial metabolism, which in turn might increase the decay rate for a range of wetland types.^{10–12} Changes in soil water content (e.g., through inundation patterns, wetland drainage, tidal cycles, and interannual variability in precipitation) can control much of the decay process in wetland soils, influencing initial mass loss through leaching and providing access to nutrients that support microbial metabolism, while also influencing oxygen availability, potentially limiting microbial mineralization under anaerobic conditions.^{2,9,13–15} Anaerobic conditions caused by waterlogging can dampen the enhanced decomposition caused by elevated temperature during the later stages of decay when decay processes are dominated by enzymatic breakdown of plant structural compounds,^{2,16} thereby complicating the impacts of warming temperatures in the long term. Further, variability in litter chemical composition (e.g., nitrogen concentration, tissue type, phylogenetic history) can be a major influence on the decomposition dynamics in wetlands and can result in litter-specific responses to external factors like salinity, inundation, and warming.^{10,17–22}

Although there is greater potential for belowground OM deposits (e.g., roots, rhizomes) to contribute to wetland soil carbon, decomposition studies in wetlands often focus on aboveground litter decomposition,^{10,23,24} despite high litter export rates and potential for herbivory.^{25–27} Additionally, existing global models of long-term decomposition are primarily based on terrestrial ecosystems and do not represent belowground wetland decay well due to different decomposition dynamics (e.g., wet conditions).²⁸ Long-term field datasets on belowground OM decomposition may help develop paradigms on the controls of wetland decay, including sea-level rise,^{29,30} soil water content, climate, and decomposers.^{31,32} Global wetland decay datasets may also help parametrize global carbon cycling through earth system models^{33,34} and by means of satellite-based models.³⁵

Since litter chemical composition strongly affects the decomposition process, it is difficult to draw broader conclusions when a range of litter types are used over regional and global scales.^{36,37} A promising approach to advance our knowledge of belowground decomposition across ecosystems and climates is to use standardized substrates *in lieu* of local plant litter.^{38,39} Standardized green tea and rooibos tea “litters” have water-soluble-dominant (labile, rapid leaching) and lignin-dominant (recalcitrant, stable) compositions, respectively.^{39,40} Tea litter has been valuable at revealing short-term, 3 month drivers of belowground litter decay at regional and global scales, although a limited collection of studies for wetland and aquatic ecosystems exists.^{41,42} There are limitations and challenges in extrapolating the drivers of short-term incubations to inform longer-term predictions and processes of decomposition, i.e., linked to carbon storage and sequestration. Longer standardized tea litter incubations (e.g., ~1 year) show how differences in inundation and oxygen conditions and temperature may be driving wetland OM decay, particularly for the more recalcitrant rooibos tea litter,^{43,44} suggesting different ecosystem responses to anthropogenic changes in the future.

To improve our understanding of belowground litter decomposition dynamics in wetlands, we performed a global, up to four-year decomposition experiment using standardized tea litter. By burying the litter bags, we are subjecting the proxy substrates to the conditions and processes that influence belowground OM turnover processes. Labile and recalcitrant OM sources, represented by green and rooibos tea litters, respectively, were incubated in the soils of inland and coastal wetlands, as well as lotic (e.g., stream) and lentic (e.g., pond/lake) ecosystems for approximately three years across eight macroclimates. First, we compared short-term (3 months) and long-term (≥ 24 months) decomposition across ecosystem types. Next, we applied asymptotic and single exponential decay rate models⁴⁵ to the entire time series at each site to produce site-level decay parameters that describe an early leaching-influenced decay rate constant, the proportion of stable mass remaining at the end of the incubation, and an overall decay rate constant. We used these parameters to explore the impacts of climatic and ecosystem properties on different stages of standardized litter decomposition. We hypothesize that warmer and higher precipitation climates will enhance decomposition but have litter-specific impacts.⁴⁶ We also expect that ecosystem type, comprised of a range of varying factors including inundation, would significantly influence longer-term decay dynamics for both litter types.^{43,44} Lastly, we used the decay parameters to project decay responses under future climatic conditions, expecting temperature to be a key driver of decomposition.¹⁰ Together, this study aims to expand our knowledge of wetland OM decay and soil carbon preservation in global wetlands to improve global climate and carbon cycle modeling, as well as to provide

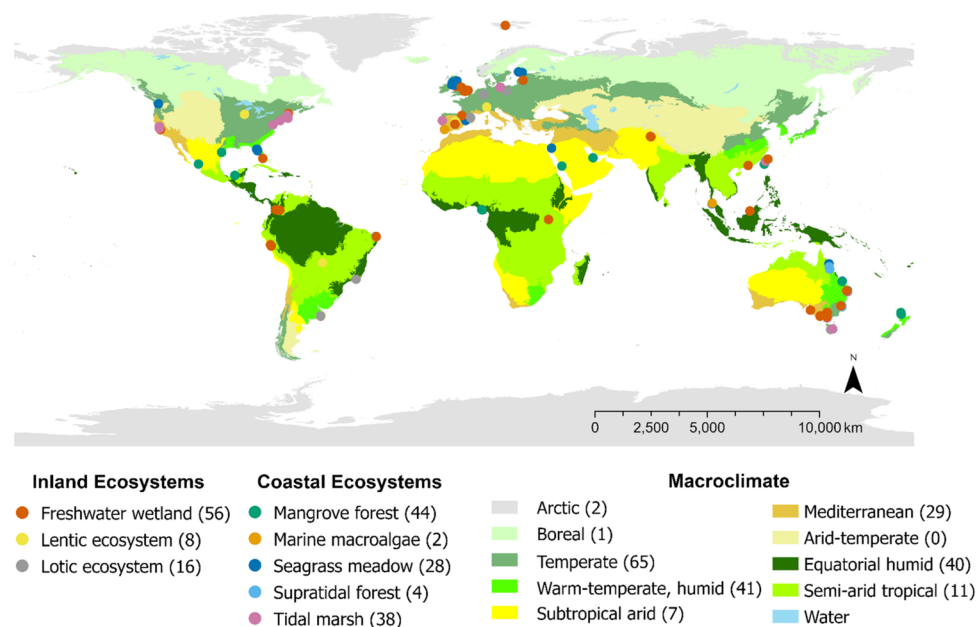


Figure 1. Map of TeaComposition H₂O sites across eight macroclimatic zones. Numbers next to ecosystem and macroclimate types indicate the site number. See Table S1 for more details on the sites. Climatic zones are from Walter and Breckle.⁵⁵

a catalyst for future long-term wetland decomposition studies using both natural and standardized litter.

2. MATERIALS AND METHODS

2.1. TeaComposition H₂O Initiative and Decomposition Experiment. The standardized litters are Lipton® (Unilever) green and rooibos teas, packaged in the original nylon mesh. Green tea represents labile forms of OM (high water-soluble compound content) and the rooibos tea represents recalcitrant/stable (high fiber/lignin content) forms of OM (*Camellia sinensis*, EAN no.: 8 722700 055525; *Aspalathus linearis*, EAN no.: 8 722700 188438).³⁹ Generally, the definitions of labile and recalcitrant are context dependent and sit along a spectrum depending on substrate type and chemical characteristics, spatiotemporal frame of observation, and microbial characteristics.⁴⁷ For the purpose of this study in which initial chemical characteristics are well-known,^{39,40} we will refer to the tea litter OM in terms of its inherent chemical characteristics,⁴⁸ that is, labile and recalcitrant OM for green and rooibos teas, respectively.

This work was performed within TeaComposition H₂O, a global initiative to collect long-term decomposition data from wetlands and aquatic ecosystems using standardized litter methods. We defined wetlands as inclusive of freshwater/aquatic and coastal/marine marshes, peatlands, and waters that do not exceed 6 m depth at low tide.¹ We focused on vegetated sites that did not receive experimental manipulations, resulting in data from 196 sites (Figure 1). See Supporting Methods for details on information collected for each site (Figure 1 and Table S1). The “freshwater wetland” category comprised a diversity of wetland types, so we further categorized using the IUCN Ecosystem Typology 2.0 for additional statistical analysis (Table S1, Supporting Methods).⁴⁹ Monthly mean temperature and total precipitation from local weather stations were calculated for each month from deployment to the final sampling. Monthly mean temperature variation was calculated as the standard deviations of the monthly temperatures during the incubation period. Subtidal sites used in the final analyses

represented <10% of the sites, most of which were lagoonal/estuarine (e.g., near intertidal). Therefore, local air temperatures were used for all sites.⁵⁰

At each site, green and rooibos tea litter bags were buried 10–15 cm deep into the soil by hand in two plots at least 1 m apart. This depth generally captures the rhizosphere or surficial OM layers of our inland and coastal wetlands while at a depth that is easily retrievable. Within each plot, there were two replicates for each tea type (i.e., $n = 4$ for each tea type at each sampling time and site).⁴³ Deployment occurred in the summer of 2017 for the northern hemisphere (e.g., ~June to August) and summer of 2017/2018 for the southern hemisphere (e.g., ~December to February). Tea bags were collected 3, 6, 12, 24, and 36 months after deployment. As sites varied in accessibility and time constraints, some sites ended the experiment earlier at 12 or 24 months. Further, the 3-year sampling in 2020 occurred during the global coronavirus pandemic, resulting in 19 sites with delayed final samplings between 36 and 48 months. The initial mass was calculated by weighing the tea in the bag and then subtracting the mean bag mass of 0.20 g (± 0.002 g S.E.M, averaged over 40 empty bags). Post-incubation samples were cleaned of soil and dried at 60–70 °C until a constant weight. Contaminating root biomass (i.e., root in-growth) was removed before weighing the final dry tea mass without the bag.

2.2. Decay Modeling. As the study focused on quantifying the long-term tea litter decay, we calculated decay parameters for sites that had incubations for 1 year or longer. Only sites with at least two replicates across three sampling times (i.e., six data points over the first year) were included. After filtering, 181 sites remained for labile green tea, and 184 sites remained for recalcitrant rooibos tea. See Supporting Methods for details on data cleaning.

For our decay modeling approach, we fit the site-level data with single exponential and asymptotic decay functions following Gill et al.⁴⁵

Asymptotic exponential decay

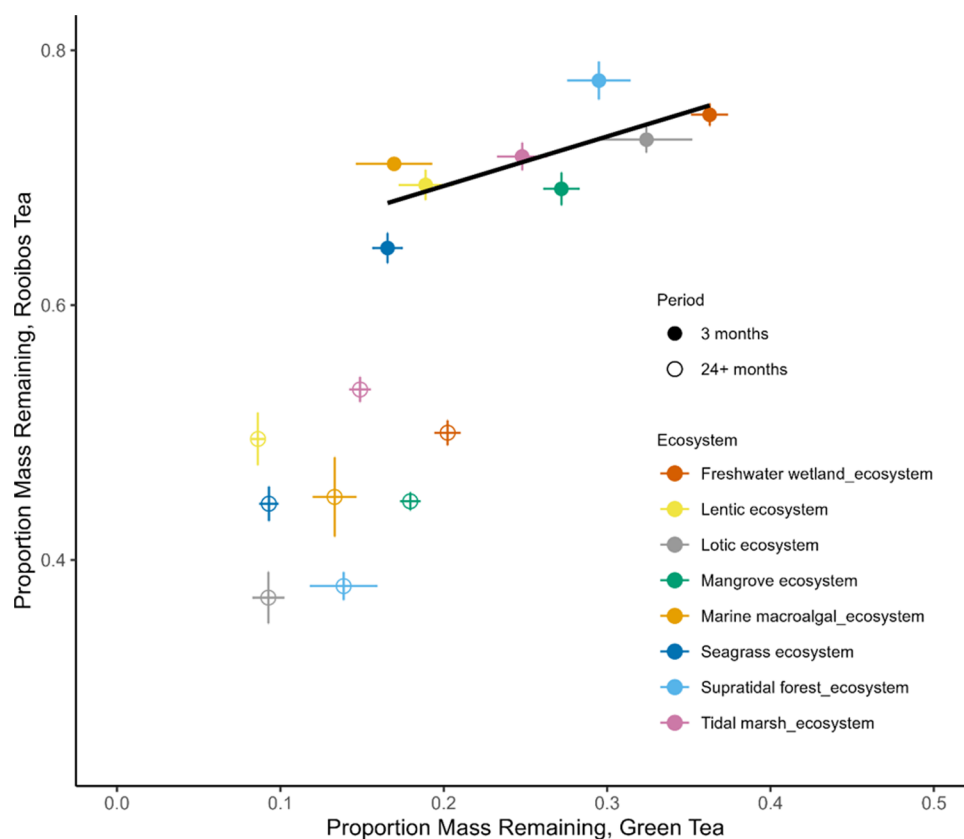


Figure 2. Total proportion of mass remaining of labile (green tea) and recalcitrant (rooibos tea) organic matter (OM) in each ecosystem type at early (3 months) and later (≥ 24 months) stages of decay. Values are means $\pm$ standard errors. Ecosystem-level values for each sampling period can be found in Table S4.

$$\text{proportion mass remaining} = A + (1 - A)e^{-tk_a} \quad (1)$$

where A is the asymptote (A), t is the time (days), and k_a is the early decay rate constant (day^{-1}). The asymptotic decay function uses a negative exponential function approaching a nonzero horizontal asymptote. This formulation partitions the tea litter between early- and late-stage decay. The early stage is characterized by the initial rapid decay ($1 - A$, as a proportion) at rate k_a (d^{-1}). The later stage is characterized by very slow or negligible decay after reaching the asymptote, i.e., the proportion of stable OM (A).

Single exponential decay

$$\text{proportion mass remaining} = e^{-tk_s} \quad (2)$$

The single exponential decay function describes the tea litter as a single pool that decomposes at a constant rate (k_s , d^{-1}) over time (t , days).

By using the parameters from both the asymptotic and the single exponential decay models, we were able to describe tea litter decay in the following ways: (1) the negative exponential rate before reaching the asymptote quantifies the early decay rate constant (k_a) and is linked to abiotic leaching of water-soluble compounds (eq 1), (2) the asymptote (A) is the proportion of stable mass remaining under a long-term decay constant and has the potential to contribute to soil carbon stocks (eq 1), and (3) the overall negative exponential decay rate constant (k_s) quantifies the overall decay rate in each time series (eq 2).⁴⁵ See Supporting Methods for details on model fits.

2.3. Statistical Analyses and Prediction Modeling.

Using linear models, we tested the effects of ecosystem type and climate on litter decay parameters (i.e., A , k_a , and k_s) for each OM type (Table S2, Supporting Methods). Ecosystem type and climate were important factors in previous shorter-term tea litter decay studies (3–12 months).^{43,51} Therefore, the first model included the following terms: precipitation, temperature, temperature variability (as standard deviation), ecosystem type, and two-way interactions between ecosystem type and each of the three local climate terms to compare the sensitivity of the ecosystem types to the climatic factors. The macroalgal ecosystems were represented by only two sites each, so they were removed for this analysis. Model selection using the Bayesian information criterion (BIC) was performed for each OM type and decay parameter combination.⁵² The second and third models for macroclimate and freshwater wetland IUCN typologies were analyzed in single-factor models separately. For all models, significant interactions between factors were explored with Tukey posthoc pairwise comparisons using the emmeans package.⁵³ All analyses were performed using the `lm()` function in R version 4.1.3.⁵⁴

We generated worldwide spatial predictions of decay parameters (i.e., A , k_a , and k_s) based on linear models using only local climate without accounting for ecosystem type due to incomplete geospatial coverage of each ecosystem type in this study. We sourced from Copernicus Climate Data Store spatially explicit climate factors for temperature, precipitation, and temperature variability using eight IPCC global climate models from the IPCC's Fifth Assessment Report (i.e., Coupled Model Intercomparison Project Phase 5; CMIP5),

Table 1. Main and Interaction Effects from Linear Models of Decay Parameters and Environmental Variables^{a,b}

model	explanatory variables	labile OM (green tea)			recalcitrant OM (rooibos tea)		
		<i>A</i>	<i>k_a</i>	<i>k_s</i>	<i>A</i>	<i>k_a</i>	<i>k_s</i>
ecosystem & climate	ecosystem:precipitation						
	ecosystem:temperature	8.49***		8.57***			
	ecosystem:temperature variation						
	ecosystem	170.99***	319.6***	2404.6***	273.15***	449.65***	5444.0***
	precipitation						
	temperature mean	18.45***		18.34***	15.84***		37.34***
	temperature variation	0.47		0.908			
macroclimate	days of incubation	2.64			7.21**	7.10**	12.48***
	macroclimate	6.51***	3.21**	7.30***	4.02***	1.46	13.03***
	days of incubation	3.30*	2.51	3.05*	5.25**	8.32***	18.0***
freshwater wetland	IUCN Typologies 2.0	4.91***	0.48	5.06***	2.45*	1.02	8.16***
	days of incubation	0.291	2.26	1.31	3.01	2.42	0.77

^aEach model is run independently for the six OM type and decay parameter combinations. For the ecosystem and climate model, the starting model included all eight variables. We used the Bayesian information criterion (BIC) to identify the best-fitting model, for which we report *F*-values and significance for each variable. Missing *F*-values in the table indicate variables selected against by BIC model selection. The macroclimate categories are presented in Figure 1. The freshwater wetland model uses the IUCN Typologies 2.0 that incorporate climate, inundation, and vegetation characteristics. Temperature variation represents the mean monthly variation during the timeframe of incubation. Two decay equations were applied to site-level time series to provide information on different stages of decay. An asymptotic exponential model (eq 1) provides *A* (proportion of stable mass remaining) and *k_a* (rate constant of early decay, i.e., leaching). A single exponential decay model (eq 2) provides *k_s* values that represent the overall decay rate constant. Values represent *F*-values, and significant *p*-values are in bold. Precipitation is in mm units. ^b*p*-values: *** < 0.001, ** 0.001 ≤ *p* ≤ 0.01, * 0.01 < *p* ≤ 0.05.

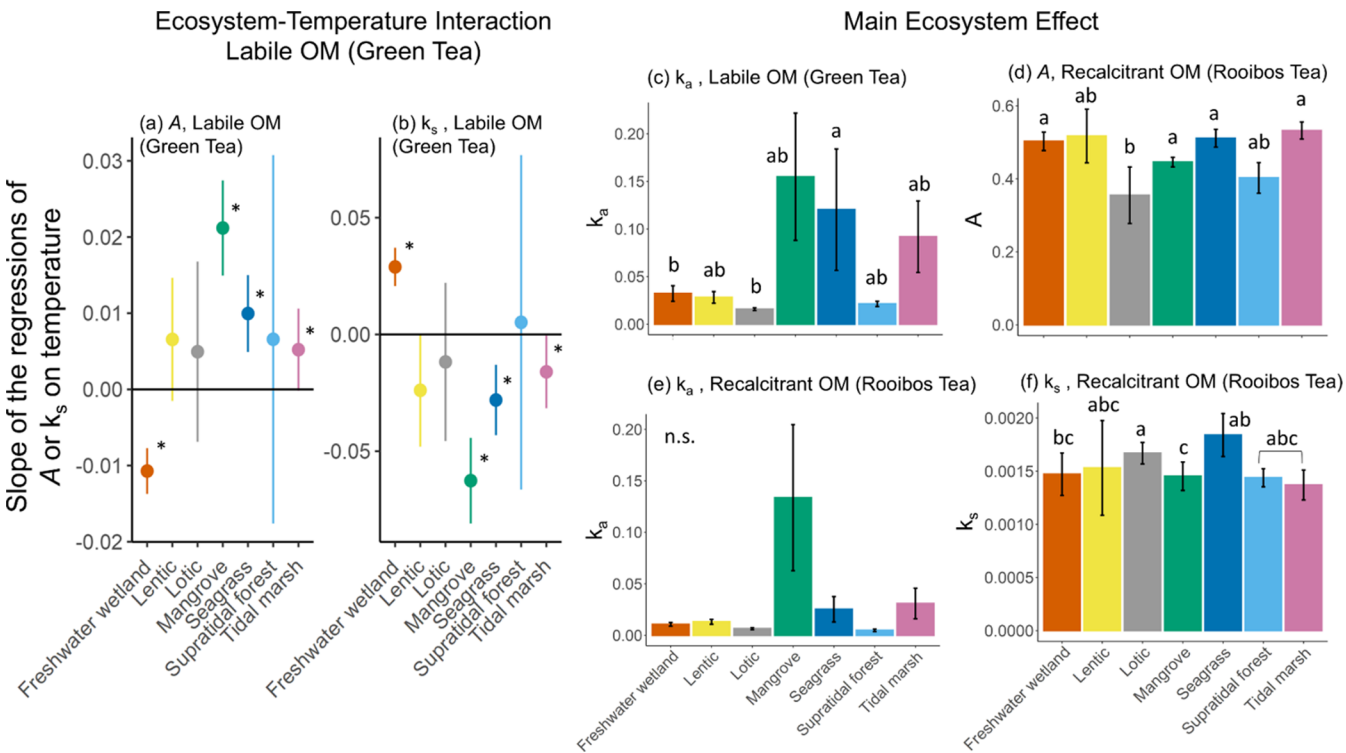


Figure 3. Ecosystem effects on the decay parameters. Each panel is the result of an independent modelrun on each OM type and decay parameter combination (Table 1). (a, b) Strength of interactions between ecosystem type and temperature for the labile organic matter (OM) type. Values represent the slope of the relationship between decomposition parameter and temperature ± the 95% confidence intervals (CIs). Asterisks indicate significant temperature effect on ecosystems (*p* < 0.05, CIs do not cross zero). (c–f) Significant main effects of the ecosystem type. Different letters indicate significant differences among the means according to Tukey's posthoc comparisons (n.s. = not significant, *p* > 0.05). Decay parameter definitions: asymptotic *A* represents the proportion of stable mass remaining (eq 1), *k_a* represents the rate constant of early decay, i.e., leaching (eq 1), and single exponential *k_s* represents the overall decay rate constant (eq 2). Data represent means ± standard errors.

assuming a Representative Concentration Pathway of 4.5 (RCP 4.5). See Supporting Methods and Table S3 for the model details. While we are mostly interested in wetland projections for inland and coastal regions of the globe, we kept

the open ocean in the scope of the projections to capture changes in decomposition parameters for wetlands of small islands. Each climate model was used to estimate yearly averages for present (January 2018 to December 2021) and

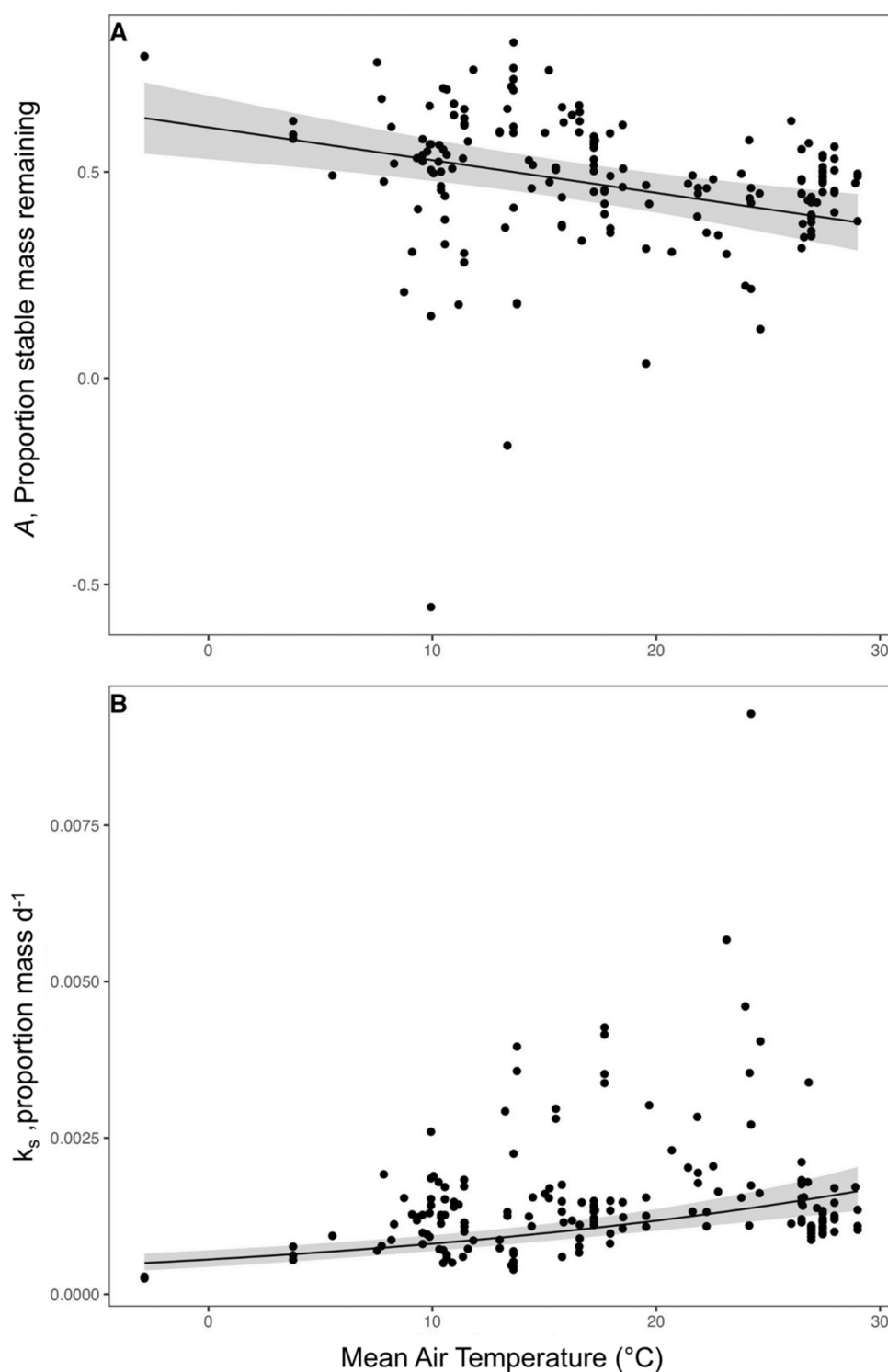


Figure 4. Temperature main effects on recalcitrant OM (rooibos tea) decay parameters. Decay parameter asymptotic A represents the proportion of stable mass remaining (eq 1) (A) and single exponential k_s represents the overall decay rate constant (eq 2) (B; d^{-1}). Raw data are plotted with a regression line with 95% confidence intervals.

future (January 2048 to December 2051) conditions, allowing us to take into consideration seasonal and interannual climate cycles. The best-fitting model was used to generate spatial predictions on decay parameters for 2020 and 2050 based on simulated conditions of rainfall and temperature extracted from each CMIP5.

3. RESULTS

3.1. Total Proportion Mass Remaining Across Ecosystem and OM Types. As expected, mass loss differed for labile and recalcitrant OM. The final remaining mass was on average $15 \pm 0.6\%$ for labile OM and $47 \pm 1\%$ for recalcitrant OM across the sites. The patterns of mass loss among ecosystem types differed between the early (3 months)

and late (≥ 24 months) stages of decay (Figure 2). Lotic and supratidal forest ecosystems had the lowest mass loss, and seagrass ecosystems had the highest mass loss for both OM types after three months of decay (Figure 2 and Table S4). However, as decay progressed, lotic and supratidal forest ecosystems experienced high mass loss for both OM types, resulting in the least mass remaining across the ecosystems (e.g., ~ 35 to 40% remaining for recalcitrant OM; Figure 2 and Table S4). By comparison, freshwater wetlands had the highest proportion of mass remaining for labile OM (19–21%), while tidal marsh ecosystems had the highest proportion of mass remaining for recalcitrant OM (~ 50 to 55%, empty circles; Figure 2 and Table S4). After 24 months, decomposition dynamics differed between OM types and between early (3 month) and late (≥ 24 months) stages of decay (Figure 2). Decomposition rates of labile and recalcitrant OM were positively correlated with each other after three months (Pearson's, p -value = 0.04, $R^2 = 0.72$). This relationship was not significant after 24+ months of decomposition (p -value = 0.4, $R^2 = 0.32$).

3.2. Effect of Ecosystem Type and Local Climate on Decomposition. Six independent models for each OM type and decay parameter combination were run for the full-factorial ecosystem-climate models, and after BIC selection, we found that ecosystem and temperature best explained OM decomposition (Table 1 and Figure 3). A statistically significant ecosystem and temperature interaction was found for stable mass remaining (asymptotic A) and overall decay rate constant (k_s) for labile OM, but not for the early decay rate constant for labile OM (k_a ; Table 1). A significant temperature effect was detected for freshwater wetlands, mangroves, seagrass, and tidal marsh ecosystems (see confidence intervals (CIs) not crossing zero in Figure 3a,b). In freshwater wetlands, increasing temperatures caused lower stable mass remaining and higher overall decay constant, while temperature had the opposite effect on these two parameters within mangrove, tidal marsh, and seagrass ecosystems (Figure 3a,b). Here, mangrove ecosystems had 2-fold and 4-fold greater effects of temperature on decomposition relative to seagrass and tidal marsh ecosystems, respectively (Figure 3a,b).

The main effects of the ecosystem type were found for early decay rate constants (k_a) of the labile OM, as well as all decay parameters for the recalcitrant OM (Figure 3c–f). For labile OM, early decay rates were significantly higher in seagrass ecosystems than in lotic and freshwater wetland ecosystems (Figure 3c). For recalcitrant OM, the remaining stable mass (A) was significantly lower in lotic ecosystems (Figure 3d). The overall decay rate of recalcitrant OM was lower in freshwater wetlands and mangroves than in seagrass and lotic ecosystems (Figure 3f). Tukey's posthoc analyses for early decay rate constants of recalcitrant OM showed only a nonsignificant effect between lotic and mangrove ecosystems ($p = 0.09$), likely due to the conservative nature of the posthoc test and the high variation in mangrove ecosystems (Figure 3e).

Increasing temperatures were found to enhance decomposition for recalcitrant OM (main temperature effect; Table 1). Higher temperatures reduced the stable mass remaining (A) and increased the overall decay constant (k_s) for recalcitrant OM (Figure 4) but did not significantly affect the early decay constant (k_a). The temperature effect was linear for A and indicated a 4% decrease in stable mass remaining for every 5 °C increase (e.g., from 52.8% at 10 °C to 44.9% at 20

°C; Figure 4A). The temperature effect on the overall decay constant (k_s) of recalcitrant OM was nonlinear. An increase from 10 to 20 °C corresponded to a 1.46-fold increase in k_s (from 0.00081 d⁻¹ at 10 °C to 0.00118 d⁻¹ at 20 °C; Figure 4B).

3.3. Effect of Macroclimate on Decomposition. Macroclimate significantly influenced decay parameters of both OM (tea) types, except for the early decay rate constant (k_a) for recalcitrant OM (Table 1). Arctic and boreal macroclimates generally had a higher proportion of stable mass remaining (A) and slower overall decay rate constant (k_s) than the other macroclimates (Figure S1a). However, arctic and boreal macroclimates were represented by only three sites, limiting our ability to make robust interpretations of the impacts of these cold macroclimates on wetland OM decay. Therefore, we focus on posthoc comparisons across the remaining six macroclimates.

The decay parameters of the two OM types generally showed that the temperate climate experienced a higher stable mass remaining (A) and slower overall decay rate constant (k_s) compared to warmer macroclimates. For labile OM, the lowest stable mass remaining values were from warm temperate and semiarid tropical climates (means 15 and 9%, respectively; Figure S1a), while the higher stable mass values (mean 21%) and slower overall decay constant (mean 0.0088 d⁻¹) were from equatorial humid climates (Figure S1a,c). For recalcitrant OM, the lowest stable mass remaining values were from warm temperate, subtropical arid, equatorial humid, and semiarid tropical climates (means 36–45%; Figure S1d). Overall decay constants of the recalcitrant OM were 2 to 3.5-fold higher ($p > 0.05$) in the semiarid tropical climate (mean 0.0038 d⁻¹) compared to temperate, warm-temperate, Mediterranean, and equatorial humid climates (means 0.0011–0.0019 d⁻¹; Figure S1e).

3.4. Effects of Freshwater Wetland IUCN Typologies on Decomposition. We used the palustrine wetland IUCN typology groupings to identify any effect of the ecosystem type on tea litter decay within the broad freshwater wetland category. The stable mass remaining of labile OM in tundras was more than 2-fold higher than that of most typologies, with the exception of boreal and temperate fens (Figure S2a). This highly stable mass remaining for the tundra coincided with a significantly slower overall decay rate constant (Figure S2e), but there was no significant difference in early decay rate constant (k_a) for the labile OM across typologies (Figure S2c). For the recalcitrant OM, permanent marshes had 4- to 24-fold and significantly higher overall decay rate constants than those of the other ecosystem types (Figure S2f). For recalcitrant OM, no significant differences across freshwater wetland typologies were found for early decay constants or the stable mass remaining (Figure S2b,d).

3.5. Projections of OM Decay Parameters. In our ecosystem-independent climate predictions, the stable mass remaining of labile OM decreased with increasing precipitation and temperature variation, while early and overall decay constants increased with temperature variation (Table S5). In contrast for recalcitrant OM, stable mass remaining decreased with increasing mean temperature and precipitation. Early decay rate constants for recalcitrant OM increased with increasing temperature variability, and overall decay rate constants were found to increase with increases in all three climatic variables (Table S5). These climatic variables were used to predict changes in each decay parameter between 2020

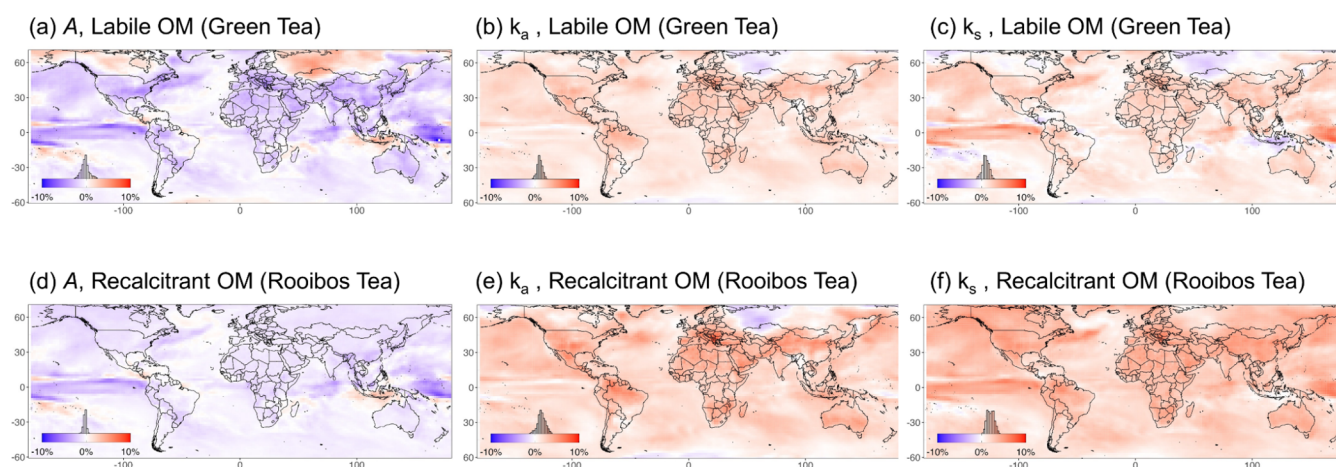


Figure 5. Percentage change of decay parameters across the globe from 2020 to 2050. Colors in the maps indicate the expected relative change in values from 2020 to 2050, with red indicating decreases, blue increases, and white no change. Model projections include open oceans to capture inland and coastal wetland habitats on the islands. The histograms show the distributions of the values in the map. The models were based on IPCC climate models (Coupled Model Intercomparison Project Phase 5, CMIP5) with an assumed representative concentration pathway (RCP) of 4.5. Decay parameter asymptotic A (a, d) represents the proportion of stable mass remaining (eq 1), k_a (b, e) represents the early decay rate constants (d^{-1} ; eq 1), and single exponential k_s (c, f) represents the overall decay rate constants (d^{-1} ; eq 2).

and 2050 by using eight CMIP5 climate models (Figures 5 and S4). The uncertainties of the predicted percent change were relatively low throughout for both OM (tea) types (Figure S4).

Overall for labile OM, the stable mass remaining (A) was predicted to decrease by 1.7% from 2020 to 2050, while early (k_a) and overall (k_s) decay rate constants were predicted to increase by 1.7 and 1.8%, respectively (Figure S4). Across the globe, the range of change was highest for labile OM stable mass remaining A , varying from approximately 4% decreases to 3% increases with extremes to $\pm 10\%$ (Figure 5a). Similarly, the overall decay constant, k_s , for labile OM was also predicted to vary, with overall changes ranging from 2% decrease to 4% increase (Figure 5c). In contrast, early decay for labile OM had generally predicted increases of $<5\%$ globally (Figure 5b). Specifically, for labile OM, hotspots for enhanced decomposition (i.e., decreased A and increased k_a and k_s) included parts of North America (particularly south/southeast), eastern Europe, and northern South America (Figure 5a–c). In contrast, parts of SE Asia and arctic regions were predicted to have an increased stable mass remaining and a decreased overall decay rate constant.

For recalcitrant OM, future climatic conditions were predicted to generally increase decomposition globally. The proportion of stable mass remaining (A) was predicted to decrease by 1%, with 2.5 and 3.1% increases in early (k_a) and overall (k_s) decay rate constants, respectively (Figure S4). For A , most of the change was predicted to occur in the range of ~ 0 to -2% (Figure 5d), while early and overall decay rate constants were predicted to have greater relative increases than labile OM, in some cases $>4\%$ (Figure 5e,f). Recalcitrant OM had hotspots of enhanced decomposition similar to those of labile OM, although with greater predicted increases in both decay constant parameters (Figure 5d–f). Small pockets of reduced decomposition were predicted for SE Asia and Central America (Figure 5d–f).

4. DISCUSSION

4.1. Impacts of Climate and Ecosystem on Long-Term OM Decomposition. Our global long-term study comprising over 180 wetland sites showed that temperature was a key

factor in belowground litter decomposition in wetlands. The long-term decomposition of recalcitrant OM (rooibos tea) was enhanced by higher mean annual temperatures and within warmer macroclimates independent of ecosystem type, similar to previous tea litter studies (~ 12 -month incubation).^{43,46} This temperature effect was found for late-stage stable mass remaining and overall decay parameters (A , k_s), but not for early decay rate constant k_a that are often associated with the passive leaching of OM (Table 1 and Figure 4). A similar persistent temperature effect has been shown for aboveground litter and belowground rhizome/root decay in coastal wetlands across similar temperature ranges as in this study.^{10,25} Increased aboveground decay rates in coastal wetlands were found in warmer climates (over $\sim 25^\circ\text{C}$),¹⁰ although the positive effect of warmer temperature on recalcitrant OM decay found here (Figure 4) was not as pronounced in comparison. A possible explanation is that the belowground environment considered in this study comprises a different set of external factors that influence decomposition compared to above-soil conditions (e.g., slower oxygen diffusion, presence of microbial activity and root exudates, porewater salinity, porewater pumping),^{10,13,25} thereby causing a slightly different realized temperature response.

Ecosystem-dependent effects of temperature were found for labile OM; here, decay was enhanced with increasing temperatures in freshwater wetlands, while tidal wetlands showed the opposite trend. Some ecosystems (e.g., seagrass meadows) had as low as 5% mass remaining (Table S4), losing much of its nonhydrolyzable fraction (fresh green tea litter has $\sim 16\%$ lignin and ash).³⁹ Perhaps, these labile OM residues are less susceptible to temperature effects on further decomposition compared to ecosystems that have relatively more OM remaining (e.g., freshwater wetlands), due to lack of accessible OM, insensitivity of microbes to temperature differences, or physicochemical conditions limiting microbial attack within tidal wetland soils.^{3,16,56,57} As such, the remaining highly decomposed OM from green tea litter in this study may be less impacted by parameters that typically enhance microbial metabolism (e.g., temperature, moisture, solubility of nutrients). Labile OM, such as that represented by green

tea, may not make substantial contributions to long-term carbon stocks in wetland soils, but certain hydrologic conditions may promote the preservation or stabilization of the remaining OM. We could not resolve if longer incubations beyond 3 years of the recalcitrant rooibos tea litter would also show an inundation-inhibited temperature effect observed for labile OM.

Ecosystem-type differences across decomposition parameters indicate that the driving forces of ecosystem characteristics on decay, such as moisture and inundation, change throughout the decomposition process. For both OM types, early decay rate constant (k_a) and mass loss at three months were generally higher for tidally influenced ecosystem types (Figure 3). This is consistent with soil moisture and inundation enhancing abiotic, leaching-driven mass loss of tea litter in short-term cross- and within-ecosystem studies for aquatic and wetland ecosystems.^{42,43,58} In contrast, the ecosystem differences were variable across the parameters that represented longer-term decay (asymptotic A , overall decay constant k_s , and mass loss at 24+ months; Figures 2 and 3). Freshwater wetlands had double the proportion of mass remaining for labile OM compared to lentic, lotic, and seagrass ecosystems. Such ecosystem differences were less pronounced for recalcitrant OM, but tidal marshes and freshwater wetlands had 3–10% more mass remaining compared to the other ecosystem types at the end of the incubation (Figure 3 and Table S4). Additionally, within the freshwater wetland typologies, permanently inundated marshes had enhanced decomposition compared to that in other freshwater wetland typologies (Figure S2). We suspect that inundation or flow rates are influencing a range of conditions that moderate biotic microbial processes at multiple scales that can increase or decrease recalcitrant OM breakdown, including soil temperature, nutrient availability, salinity, and oxygen.^{59–63}

4.2. OM Decomposition under Future Climate Scenarios. Our model based on RCP 4.5 predicts that decomposition will increase under future climate scenarios for both labile and recalcitrant OM. Labile OM (e.g., green tea litter) will have a higher percent change for loss of stable OM (parameter A) under future conditions compared to recalcitrant OM (e.g., rooibos tea litter). However, since the labile OM type is relatively low in mass and largely represented by OM residues at this stage of decay, the net change in labile OM will likely be small. By comparison, decomposition of recalcitrant OM occurs over a longer period and will likely be more susceptible to increasing temperatures in global wetlands, which is important for future predictions of soil carbon stocks and flux (i.e., blue and teal carbon). Overall, our results for recalcitrant OM are consistent with the trends of climate change-induced warming increasing plant decomposition in terrestrial (14–27% by 2070s)⁶⁴ and coastal ecosystems (2–3-fold increase in carbon loss between 2050 and 2100).¹⁰ Ouyang and colleagues¹⁰ also found that the potential for plant biomass production of carbon in the future to out-pace carbon remineralization (i.e., net carbon gains), in addition to ecosystem-specific differences for predicted enhancement of decomposition (mangrove > tidal marsh > seagrass). Warming can also impact wetland plant diversity and composition,⁶⁵ which could in turn influence the production of recalcitrant OM, e.g., lignocellulose, in a wetland.⁶⁶ While ecosystem type was not included in the model in our study, studies on natural litter and the ecosystem-climate model responses of the different OM types from this study using standardized litter

highlight the importance of resolving ecosystem-specific climate responses of standardized OM, to better predict the effects of climate change on wetlands at regional or global scales.

There are uncertainties and gaps in our approach and data set that can be addressed in future studies to increase the breadth and scope of long-term tea decay datasets for improved prediction and interpretation, including a greater sampling of poorly studied ecosystems (macroalgal, brackish ecosystem) and macroclimate types (Arctic, boreal, and arid temperate). There are uncertainties associated with modeling based on future air temperatures instead of future soil temperatures, which may impact the variability of temperature change the litter would experience.⁶⁷ Additionally, our study was designed to explore the global-scale drivers of decay. Measuring additional environmental metrics at the site level, such as salinity, inundation/flooding period and groundwater metrics, vegetation, nutrient concentrations, and soil mineral composition, will provide further insight into local-scale drivers of belowground litter decay in wetlands in the context of regional- and global-scale drivers (e.g., micro- vs macroclimate controls),^{25,68,69} as well as vulnerability to droughts and sea-level rise.^{70–72}

4.3. Environmental Implications. This study provides the first empirical comparison of multiyear wetland decomposition using standardized OM, providing new insight on the benefits and limitations of using tea litter to resolve longer-term controls on belowground litter turnover. By deriving multiple decay parameters that describe different stages of decomposition, we show that few climatic- or ecosystem-type variables influenced early leaching-dominated decay (k_a) for either OM (tea) type. We also observed relative changes in mass loss among ecosystem types between 3 and ≥ 24 months time points (Figure 2). These results highlight that short-term decomposition incubations capture less of the processes occurring during OM turnover and are likely not suitable for making inferences about soil carbon preservation and long-term storage (blue carbon and teal carbon). Rather, the stable portion of mass (A) and overall decay rate constant (k_s) derived from multiyear time series datasets reveal that increased temperatures and ecosystem traits, aggregated at a global scale, can significantly increase belowground litter remineralization.

Additionally, our study reveals a significant positive relationship between modeled stable mass remaining (A , eq 1) and raw final proportion mass remaining values at the end of the incubation for both OM types, as well as a consistent inverse relationship between A and k_s across all our statistical models (Figure S5). Only in a few sites was the asymptote not reached, resulting in negative A values (Figure S5). While model-fitting of time-series datasets is recommended for short-term 3 months incubations,⁷³ these correlative relationships in this study suggest that the final mass remaining data from longer-term incubations (e.g., >1 year) are representative of wetland long-term decomposition processes. As longer-term tea litter studies become more common,^{44,74} these findings highlight how the final mass remaining data of standardized OM may be useful as a simple metric for long-term ecosystem decomposition modeling without the need for gathering time- and cost-intensive time-series datasets.

We propose that standardized tea litter decay data could contribute to addressing the limited long-term decay datasets for wetlands, which represents a current barrier in predicting

wetland carbon-cycling at the scale and biogeographic levels available for terrestrial ecosystems. Multiyear and cross-ecosystem decay time series data (e.g., LIDET) and modeling frameworks, such as WARMER-2, have an immense value for evaluating the drivers of decomposition from local and regional to global scales now and in the future.^{28,29,37,75} Since we have shown that the final mass remaining can be used as a predictor of longer-term decay rates, standardized litter datasets can contribute to existing efforts to fill this gap in wetland ecosystems, including their value as translatable datasets across existing networks and databases.⁷⁶ A key limitation of tea litter studies is their limited chemical diversity compared to natural litter, and this may limit our interpretation of OM turnover.⁷⁷ Transplant and long-term studies that directly compare local wetland OM and tea litters⁷⁸ would help reveal if any and what uncertainties exist and the ecosystems where tea litters are less representative in characterizing natural OM turnover (e.g., homefield advantage),^{77,79} as well as allowing for the quantification of such uncertainties in predictive analyses. Future studies measuring carbon fluxes (e.g., respiration, loss of carbon with decay) to estimate potential changes in greenhouse gas emissions, as well as microbial community and function, can complement standardized decay metrics to predict net change in carbon and parameterize system models under future climate change.^{10,32,34,80} Together, addressing these unknowns concerning OM preservation in wetlands, using multiple approaches including both standardized and natural litters, will advance our knowledge on potential threats to soil carbon sequestration capacity of wetlands now and in the future.

■ ASSOCIATED CONTENT

SI Supporting Information

The Supporting Information is available free of charge at <https://pubs.acs.org/doi/10.1021/acs.est.4c02116>.

Detailed methodology; descriptions of sites and models used in the study; decay parameters from different incubation periods, local climate conditions, macro-climates, and freshwater wetland IUCN typologies; predicted change in litter decay and coefficient of variables for predicted change between 2020 and 2025; and regressions between decay parameters (PDF)

■ AUTHOR INFORMATION

Corresponding Author

Stacey M. Trevathan-Tackett — Deakin Marine Research and Innovation Centre, School of Life and Environmental Sciences, Deakin University, Melbourne, Victoria 3125, Australia; Centre for Nature Positive Solutions, Biosciences and Food Technology Discipline, RMIT University, Melbourne, Victoria 3001, Australia; orcid.org/0000-0002-4977-0757; Phone: +61 3 9925 3888; Email: stacey.trevathan-tackett@rmit.edu.au

Authors

Sebastian Kepfer-Rojas — Department of Geosciences and Natural Resource Management, University of Copenhagen, Frederiksberg 1165, Denmark
Martino Malerba — Deakin Marine Research and Innovation Centre, School of Life and Environmental Sciences, Deakin University, Melbourne, Victoria 3125, Australia; Centre for Nature Positive Solutions, Biosciences and Food Technology

Discipline, RMIT University, Melbourne, Victoria 3001, Australia; orcid.org/0000-0002-7480-4779

Peter I. Macreadie — Deakin Marine Research and Innovation Centre, School of Life and Environmental Sciences, Deakin University, Melbourne, Victoria 3125, Australia; Centre for Nature Positive Solutions, Biosciences and Food Technology Discipline, RMIT University, Melbourne, Victoria 3001, Australia

Ika Djukic — Swiss Federal Institute for Forest, Snow and Landscape Research WSL, Birmensdorf 8903, Switzerland

Junbin Zhao — Norwegian Institute of Bioeconomy Research, 1430 As, Norway; Institute of Environment and Department of Biological Sciences, Florida International University, Miami, Florida 33199, United States; orcid.org/0000-0001-5142-4901

Erica B. Young — Department of Biological Sciences and School of Freshwater Sciences, University of Wisconsin-Milwaukee, Milwaukee, Wisconsin 53211, United States

Paul H. York — Centre for Tropical Water and Aquatic Ecosystem Research, James Cook University, Cairns, Queensland 4870, Australia

Shin-Cheng Yeh — Graduate Institute of Sustainability Management and Environmental Education, National Taiwan Normal University, Taipei 11677, Taiwan

Yanmei Xiong — Research Institute of Tropical Forestry, Chinese Academy of Forestry, Guangzhou 510520, China

Gidon Winters — The Dead Sea and Arava Science Center, Masada National Park, Mount Masada 8698000, Israel; Eilat Campus, Ben-Gurion University of the Negev, Hatmarim Boulevard, Eilat 8855630, Israel

Danielle Whitlock — School of Applied Sciences, Edinburgh Napier University, Edinburgh EH11 4BN, U.K.

Carolyn A. Weaver — Department of Biology, Millersville University, Millersville, Pennsylvania 17551, United States

Anne Watson — School of Natural Sciences, University of Tasmania, Hobart, TAS 7005, Australia

Inger Visby — Derwent Estuary Program, Hobart, TAS 7000, Australia

Jacek Tytkowski — Institute of Geocology and Geoinformation, Adam Mickiewicz University, 61-680 Poznan, Poland

Allison Trethowan — Education Officer, RiverConnect - Greater Shepparton, Shepparton, Victoria 3630, Australia

Scott Tiegs — Department of Biological Sciences, Oakland University, Rochester, Michigan 48309, United States

Ben Taylor — Nature Glenelg Trust, Mount Gambier, South Australia 5290, Australia

Jozef Szpikowski — Institute of Geocology and Geoinformation, Adam Mickiewicz University, 61-680 Poznan, Poland

Grażyna Szpikowska — Geoecological Station in Storkowo, Adam Mickiewicz University, 78-450 Poznan, Poland

Victoria L. Strickland — Long Beach High School, Lido Beach, New York 11561, United States

Normunds Stivrins — Department of Geography, University of Latvia, Riga LV-1004, Latvia; Department of Geology, Tallinn University of Technology, Tallinn 19086, Estonia; Faculty of Forest and Environmental Sciences, Latvia University of Life Sciences and Technologies, Jelgava LV-3001, Latvia

Ana I. Sousa — ECOMARE - Laboratory for Innovation and Sustainability of Marine Biological Resources, CESAM - Centre for Environmental and Marine Studies, Department of

- Biology, University of Aveiro, 3810-193 Aveiro, Portugal; orcid.org/0000-0003-0783-5177
- Sutinee Sinutok** – Faculty of Environmental Management, Prince of Songkla University, Hat Yai 90110, Thailand
- Whitney A. Scheffel** – Pensacola and Perdido Bays Estuary Program, Pensacola, Florida 32514, United States
- Rui Santos** – Center of Marine Sciences, University of Algarve, 8005-139 Faro, Portugal
- Jonathan Sanderman** – Woodwell Climate Research Center, Falmouth, Massachusetts 02540, United States
- Salvador Sánchez-Carrillo** – Department of Biogeochemistry and Microbial Ecology, National Museum of Natural Sciences (MNCN-CSIC), E-28006 Madrid, Spain
- Joan-Albert Sanchez-Cabeza** – Unidad Académica Mazatlán, Instituto de Ciencias del Mar y Limnología, Universidad Nacional Autónoma de México, Mazatlán 82040, Mexico; orcid.org/0000-0002-3540-1168
- Krzysztof G. Rymer** – Adam Mickiewicz University Polar Station “Petuniabukta”, Adam Mickiewicz University, 61-680 Poznan, Poland
- Ana Carolina Ruiz-Fernandez** – Unidad Académica Mazatlán, Instituto de Ciencias del Mar y Limnología, Universidad Nacional Autónoma de México, Mazatlán 82040, Mexico
- Bjorn J. M. Robroek** – Department of Ecology, Radboud Institute for Biological and Environmental Sciences, Radboud University, Nijmegen 6500 HC, The Netherlands
- Tessa Roberts** – Nature Glenelg Trust, Mount Gambier, South Australia 5290, Australia
- Aurora M. Ricart** – Institut de Ciències del Mar (ICM-CSIC), 08003 Barcelona, Spain; Bigelow Laboratory for Ocean Sciences, East Boothbay, Maine 04544, United States
- Laura K. Reynolds** – Soil Water and Ecosystem Sciences Department, University of Florida, Gainesville, Florida 32603, United States
- Grzegorz Rachlewicz** – Institute of Geoecology and Geoinformation, Adam Mickiewicz University, 61-680 Poznan, Poland
- Anchana Prathep** – Division of Biological Science, Faculty of Science, Prince of Songkla University, Hat Yai 90110, Thailand
- Andrew J. Pinsonneault** – Smithsonian Environmental Research Center, Edgewater, Maryland 21037, United States
- Elise Pendall** – Hawkesbury Institute for the Environment, Western Sydney University, Penrith, NSW 2751, Australia
- Richard Payne** – Department of Environment and Geography, University of York, York YO10 5NG, U.K.
- Ilze Ozola** – Lake and Peatland Research Centre, Puikule, Purvisi LV-4063, Latvia
- Cody Onufrock** – Long Beach High School, Lido Beach, New York 11561, United States
- Anne Ola** – School of the Environment, The University of Queensland, St. Lucia, Queensland 4072, Australia
- Steven F. Oberbauer** – Institute of Environment and Department of Biological Sciences, Florida International University, Miami, Florida 33199, United States
- Aroloye O. Numbere** – Department of Animal and Environmental Biology, University of Port Harcourt, P.M.B 5323 Choba, Nigeria
- Alyssa B. Novak** – Department of Earth and Environment, Boston University, Boston, Massachusetts 02215, United States
- Joanna Norkko** – Tvärminne Zoological Station, University of Helsinki, 10900 Hanko, Finland; orcid.org/0000-0001-9885-8408
- Alf Norkko** – Tvärminne Zoological Station, University of Helsinki, 10900 Hanko, Finland
- Thomas J. Mozdzer** – Bryn Mawr College, Department of Biology, Bryn Mawr, Pennsylvania 19010, United States
- Pam Morgan** – University of New England, Biddeford, Maine 04005, United States
- Diana I. Montemayor** – Instituto de Investigaciones Marinas y Costeras (IIMYC), Facultad de Ciencias Exactas y Naturales, Universidad Nacional de Mar del Plata, CONICET, 7600 Mar del Plata, Argentina
- Charles W. Martin** – Stokes School of Marine and Environmental Sciences, University of South Alabama and Dauphin Island Sea Lab, Dauphin Island, Alabama 36528, United States
- Sparkle L. Malone** – Yale School of the Environment, New Haven, Connecticut 06511, United States
- Maciej Major** – Institute of Geoecology and Geoinformation, Adam Mickiewicz University, 61-680 Poznan, Poland
- Mikolaj Majewski** – Institute of Geoecology and Geoinformation, Adam Mickiewicz University, 61-680 Poznan, Poland
- Carolyn J. Lundquist** – National Institute of Water & Atmospheric Research, Hamilton 3251, New Zealand; School of Environment, The University of Auckland, Auckland 1142, New Zealand
- Catherine E. Lovelock** – School of the Environment, The University of Queensland, St. Lucia, Queensland 4072, Australia
- Songlin Liu** – Key Laboratory of Tropical Marine Bio-resources and Ecology, South China Sea Institute of Oceanology, Chinese Academy of Sciences, Guangzhou 510301, China; orcid.org/0000-0002-6478-4938
- Hsing-Juh Lin** – Department of Life Sciences and Innovation and Development Center of Sustainable Agriculture, National Chung Hsing University, Taichung 402202, Taiwan
- Ana Lillebo** – ECOMARE - Laboratory for Innovation and Sustainability of Marine Biological Resources, CESAM - Centre for Environmental and Marine Studies, Department of Biology, University of Aveiro, 3810-193 Aveiro, Portugal
- Jinqian Li** – Ministry of Education Key Laboratory for Biodiversity Science and Ecological Engineering, National Observations and Research Station for Wetland Ecosystems of the Yangtze Estuary, Institute of Biodiversity Science and Institute of Eco-Chongming, School of Life Sciences, Fudan University, Shanghai 200438, China; orcid.org/0000-0002-2219-2571
- John S. Kominoski** – Institute of Environment and Department of Biological Sciences, Florida International University, Miami, Florida 33199, United States; orcid.org/0000-0002-0978-3326
- Anzar Ahmad Khuroo** – Centre for Biodiversity & Taxonomy, Department of Botany, University of Kashmir, Srinagar 190006 Jammu & Kashmir, India
- Jeffrey J. Kelleway** – School of Science and Environmental Futures Research Centre, University of Wollongong, Wollongong, New South Wales 2522, Australia
- Kristin I. Jinks** – Coastal and Marine Research Centre, Australian Rivers Institute, School of Environment and Science, Griffith University, Gold Coast, Queensland 4222, Australia

- Daniel Jerónimo** — ECOMARE - Laboratory for Innovation and Sustainability of Marine Biological Resources, CESAM - Centre for Environmental and Marine Studies, Department of Biology, University of Aveiro, 3810-193 Aveiro, Portugal
- Christopher Janousek** — Department of Fisheries, Wildlife and Conservation Sciences, Oregon State University, Corvallis, Oregon 97331, United States; orcid.org/0000-0003-2124-6715
- Emma L. Jackson** — Coastal Marine Ecosystems Research Centre, CQUniversity, Gladstone, Queensland 4680, Australia
- Oscar Iribarne** — Instituto de Investigaciones Marinas y Costeras (IIMYC), Facultad de Ciencias Exactas y Naturales, Universidad Nacional de Mar del Plata, CONICET, 7600 Mar del Plata, Argentina
- Torrance Hanley** — Department of Marine and Environmental Science, Marine Science Center, Northeastern University, Nahant, Massachusetts 01908, United States
- Maroof Hamid** — Centre for Biodiversity & Taxonomy, Department of Botany, University of Kashmir, Srinagar 190006 Jammu & Kashmir, India
- Arjun Gupta** — Long Beach High School, Lido Beach, New York 11561, United States
- Rafael D. Guariento** — Laboratory of Ecology, Federal University of Mato Grosso do Sul, Campo Grande 79070-900, Brazil
- Ieva Grudzinska** — Lake and Peatland Research Centre, Puikule, Purvisi LV-4063, Latvia
- Anderson da Rocha Gripp** — Institute of Biodiversity and Sustainability, Federal University of Rio de Janeiro, Rio de Janeiro 27965045, Brazil; orcid.org/0000-0002-0099-6990
- María A. González Sagrario** — Instituto de Investigaciones Marinas y Costeras (IIMYC), Facultad de Ciencias Exactas y Naturales, Universidad Nacional de Mar del Plata, CONICET, 7600 Mar del Plata, Argentina
- Laura M. Garrison** — Santa Clara Valley Water District, San Jose, California 95118, United States
- Karine Gagnon** — Environmental and Marine Biology, Åbo Akademi University, 20500 Turku, Finland; Institute of Marine Research, Flødevigen Research Station, 4817 His, Norway; orcid.org/0000-0002-0971-7740
- Esperança Gacia** — Integrative Freshwater Ecology Group, Centre for Advanced Studies of Blanes (CEAB-CSIC), 17300 Girona, Spain
- Marco Fusi** — Dove Marine Laboratory, School of Natural and Environmental Sciences, Newcastle University, Newcastle-upon-Tyne NE1 7RU, U.K.
- Lachlan Farrington** — Nature Glenelg Trust, Warrnambool, Victoria 3280, Australia
- Jenny Farmer** — School of Biological Sciences, University of Aberdeen, Aberdeen AB243UL, U.K.
- Francisco de Assis Esteves** — Institute of Biodiversity and Sustainability, Federal University of Rio de Janeiro, Rio de Janeiro 27965045, Brazil
- Mauricio Escapa** — Instituto de Investigaciones Marinas y Costeras (IIMYC), Facultad de Ciencias Exactas y Naturales, Universidad Nacional de Mar del Plata, CONICET, 7600 Mar del Plata, Argentina
- Monika Domańska** — Geoeological Station in Storkowo, Adam Mickiewicz University, 78-450 Poznan, Poland
- André T. C. Dias** — Departamento de Ecologia, Universidade Federal do Rio de Janeiro, Rio de Janeiro 21941-902, Brazil
- Carmen B. de los Santos** — Center of Marine Sciences, University of Algarve, 8005-139 Faro, Portugal; orcid.org/0000-0002-7013-494X
- Daniele Daffonchio** — Biological and Environmental Sciences and Engineering Division, King Abdullah University of Science and Technology, Thuwal 23955-6900, Kingdom of Saudi Arabia
- Paweł M. Czyryca** — Natural Environment Monitoring Station, Adam Mickiewicz University, 72-510 Międzyzdroje, Poland
- Rod M. Connolly** — Coastal and Marine Research Centre, Australian Rivers Institute, School of Environment and Science, Griffith University, Gold Coast, Queensland 4222, Australia
- Alexander Cobb** — Center for Environmental Sensing and Monitoring, Singapore-MIT Alliance for Research and Technology, Singapore 138602, Singapore; orcid.org/0000-0002-3128-3002
- Maria Chudzińska** — Integrated Monitoring of Natural Environment Station Poznań-Morasko, Adam Mickiewicz University, 61-680 Poznan, Poland
- Bart Christiaen** — Washington State Department of Natural Resources, Seattle, Washington 98504, United States
- Peter Chiffard** — Department for Geography, Philipps-University Marburg, D-35032 Marburg, Germany
- Sara Castelar** — Integrative Freshwater Ecology Group, Centre for Advanced Studies of Blanes (CEAB-CSIC), 17300 Girona, Spain
- Luciana S. Carneiro** — Department of Ecology, Federal University of Rio Grande do Norte, Natal 59072 970, Brazil
- José Gilberto Cardoso-Mohedano** — Estación del Carmen, Instituto de Ciencias del Mar y Limnología, Universidad Nacional Autónoma de México, Ciudad del Carmen 24157, Mexico
- Megan Camden** — Department of Biological Sciences, Oakland University, Rochester, Michigan 48309, United States
- Adriano Caliman** — Department of Ecology, Federal University of Rio Grande do Norte, Natal 59072 970, Brazil
- Richard H. Bulmer** — Tidal Research, Ltd., Auckland 0626, New Zealand
- Jennifer Bowen** — Department of Marine and Environmental Science, Marine Science Center, Northeastern University, Nahant, Massachusetts 01908, United States
- Christoffer Boström** — Environmental and Marine Biology, Åbo Akademi University, 20500 Turku, Finland
- Susana Bernal** — Integrative Freshwater Ecology Group, Centre for Advanced Studies of Blanes (CEAB-CSIC), 17300 Girona, Spain; orcid.org/0000-0002-6726-8840
- John A. Berges** — Department of Biological Sciences and School of Freshwater Sciences, University of Wisconsin-Milwaukee, Milwaukee, Wisconsin 53211, United States
- Juan C. Benavides** — Faculty of Environmental and Rural Studies, Pontificia Universidad Javeriana, 110321 Bogota, Colombia; orcid.org/0000-0002-9694-2195
- Savanna C. Barry** — Nature Coast Biological Station, University of Florida, Cedar Key, Florida 35625, United States; orcid.org/0000-0002-8743-4383
- Juha M. Alatalo** — Environmental Science Center, Qatar University, 2713 Doha, Qatar; orcid.org/0000-0001-5084-850X
- Alia N. Al-Haj** — Smithsonian Environmental Research Center, Edgewater, Maryland 21037, United States; Department of

Earth and Environment, Boston University, Boston,
Massachusetts 02215, United States

Maria Fernanda Adame – Australian Rivers Institute, Centre
for Marine and Coastal Research, Griffith University,
Nathan, QLD 4111, Australia

Complete contact information is available at:
<https://pubs.acs.org/10.1021/acs.est.4c02116>

Author Contributions

S.M.T.-T.: Conceptualization, data curation, formal analysis, funding acquisition, investigation, methodology, project administration, visualization, and writing—original draft preparation; S.K.-R.: Conceptualization, formal analysis, methodology, visualization, and writing—review and editing; M.M.: Formal analysis, methodology, visualization, and writing—review and editing; P.I.M.: Conceptualization, funding acquisition, and writing—review and editing; I.D.: Conceptualization, methodology, resources, and writing—review and editing; and co-authors of TeaComposition H₂O (J.Z. to F.A.): Investigation and writing—review and editing.

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Notes

The authors declare no competing financial interest.

[†]R. Payne co-authorship is posthumous (June 2019).

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