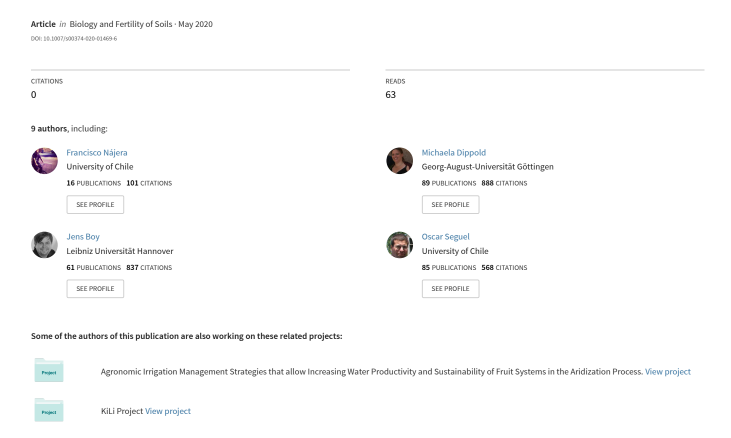
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ORIGINAL PAPER



Effects of drying/rewetting on soil aggregate dynamics and implications for organic matter turnover

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Abstract

Drying and rewetting (D/W) of soil have significant impacts on soil organic matter (SOM) turnover. We hypothesised that frequent D/W cycles would release the labile organic matter locked away in soil aggregates, increasing the priming effect (PE) (acceleration or retardation of SOM turnover after fresh substrate addition) due to preferential utilisation by microbes. ¹³Clabelled lignocellulose was added to the soil, and the effects of 0, 1, or 4 cycles of D/W were evaluated at 5 °C and 25 °C after a 27-day incubation of undisturbed soil cores from a temperate forest (Araucaria araucana). Following the incubation, macroaggregates (> 250 μm), microaggregates (250–53 μm), and silt + clay materials (< 53 μm) were separated. For each aggregate size class, three organic matter (OM) fractions (light (fPOM < 1.6 g cm⁻³), occluded (oPOM 1.6-2.0 g cm⁻³), and heavy (Hf> 2.0 g cm⁻³) were determined. D/W cycles caused macroaggregates to increase and a decrease in microaggregates (>15%) at warm temperatures, and preferential use of the novel particulate organic matter (13C labelled), formerly protected fPOM. CO₂ efflux was three times higher at 25 °C than at 5 °C. The D/W cycles at 25 °C had a strong negative impact on cumulative CO₂ efflux, which decreased by approximately -30%, induced by a negative PE of -50 mg C kg⁻¹ soil with 1 D/W cycle and -100 mg C kg⁻¹ soil with 4 D/W cycles, relative to soil under constant soil moisture receiving ¹³C-labelled lignocellulose, but no cycles. Increasing the temperature and the number of D/W cycles caused a decrease in substrate use efficiency of particulate lignocellulose. In conclusion, D/W cycles at warm temperatures accelerated OM turnover due to preferential use from fPOM, increasing macroaggregates at the expense of microaggregates. A novel pathway of OM release and PE due to the D/W cycles is discussed.

Keywords Soil priming effect · Particulate soil organic matter · Drying and rewetting cycles · Aggregate stability · Carbon turnover

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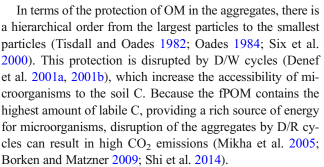


Introduction

Drying/rewetting (D/W) cycles lead to gains or losses in soil carbon (C) from soil organic matter (SOM), effects that are enhanced under extreme climatic events (Kim et al. 2012). However, soil C turnover is dependent upon other environmental conditions, e.g. temperature (Davidson and Janssens 2006). Small changes in mean annual temperature can have significant effects on soil CO₂ release (Billings and Ballantyne IV 2013). Soil organic C turnover is further modified by topography, vegetation, and soil type (Balser and Firestone 2005; Vargas et al. 2010). Other factors, such as physical protection of organic matter (OM) (von Lützow et al. 2007) or the frequency of D/W cycling during dry seasons, also play critical roles in soil C dynamics (Hibbard et al. 2005).

Cycles of D/W are assumed to affect the overall functions of soils in terrestrial ecosystems and to affect soil emission of greenhouse gases such as CO2, methane, and nitrous oxides (Corti et al. 2002; Lal 2004; Vicca et al. 2014). Increasingly frequent D/W cycles could therefore cause a breakdown of aggregates (slacking), exposing the physically protected OM to microbial decomposition (Adu and Oades 1978; Appel 1998; Oztas and Fayetorbay 2003). Greater intensities of drought could intensify negative impacts on CO₂ flow and microbial activity (Sinha and Cherkauer 2010; Meisner et al. 2015). After rewetting, an increase in gas fluxes occurs via the Birch effect (Birch 1958). The Birch effect is driven by the labile particulate organic matter (POM), determined by density fractionation as the light fraction (fPOM $< 1.6 \text{ g cm}^{-3}$), which consists mostly of pieces of plant residue and fungal hyphae. These materials can be occluded POM (oPOM 1.6-2.0 g cm⁻³), protected by the aggregates (Christensen 1992; von Lützow et al. 2007). The CO₂ efflux from the soil can decline with successive D/W events as a result of an increasingly limited pool of labile substrates (Schimel and Mikan 2005; Fernández et al. 2006; Goldberg et al. 2008).

The addition of fresh organic matter to soils results in a C cycle phenomenon known as the priming effect (PE) (Bingemann et al. 1953). The PE is a strong, short-term change in the turnover of SOM caused by an input of fresh OM (Jenkinson et al. 1985; Kuzyakov et al. 2000). It is calculated as the difference between unlabelled CO2 efflux and labelled CO₂ from ¹³C- (or ¹⁴C) added to the soil (Oades 1988; Jarvis et al. 2007). Priming can be positive (acceleration) or negative (retardation) depending on the quantity and quality of fresh input (Kuzyakov 2010; Garcia-Pausas and Paterson 2011). Although the effect is considered a short-term phenomenon, Fontaine et al. (2011) demonstrated that priming could have long-lasting effects. Hence, under frequent D/W cycles, the PE can have a significant impact on the decomposition of OM fractions, triggering CO₂ efflux from native SOM in the ecosystem (Magid and Kjærgaard 2001; Gregorich et al. 2006).



A new perspective on C release and PE due to the D/W cycle is introduced in this study. Drying and rewetting cycles are hypothesised to lead to the preferential use of new, unprotected, and labile organic matter over native C, resulting in negative PE values. Quantifying this effect under the application of ¹³C-labelled fPOM to soils will facilitate (a) differentiating the degree of physicochemical protection of the SOM in various aggregate size classes and (b) estimating the substrate use efficiency, i.e. the relative proportion of added fPOM-C that is incorporated into microbial biomass.

Temperate forests in Chile have experienced increasing temperatures and more frequent extreme climatic events, such as severe droughts (Garreaud et al. 2017; Urrutia-Jalabert et al. 2018). In light of the effects of these events on soil moisture content, it is important to understand the impact of D/W on SOM turnover in these ecosystems. Three hypotheses were tested: (i) priming of native C is induced by the amendment of fresh C-input, but D/W cycles release OM, which primarily consists of the fPOM from disrupted aggregates (Fig. 1). Therefore, comparing the difference in the PE between 0 cycle and D/W cycles will allow us to quantify the effect of D/W on the actual PE. (ii) Native SOM decomposition will be retarded (negative PE) due to the preferential use of new OM by microorganisms (Fig. 1). And, (iii) a cumulatively more negative PE is expected with an increased number of D/W cycles. The aim of this study was to evaluate the effects of two frequencies of D/W events on the PE, soil aggregate size class distribution, and their OM fractions, dependent upon temperature in a temperate forest soil.

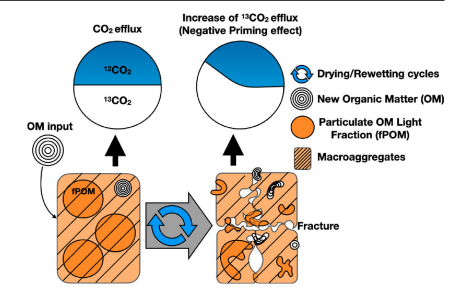
Materials and methods

Study site and sampling

Soil samples were collected from an Inceptisol (Soil Survey Staff 2014) developed under an ancient temperate forest with a dominant tree canopy of *Araucaria araucana* (Molina) K. Koch in Nahuelbuta National Park (37°47′S, 72°59′W), Chile. Important soil properties are provided in Table 1, and a more detailed description of the site can be found in Bernhard et al. (2018) and Oeser et al. (2018). Undisturbed cores (PVC 5-cm diameter × 5-cm length) were taken from the uppermost soil



Fig. 1 Schematic illustration of the impact of drying/rewetting (D/W) events on the soil C dynamics and CO₂ efflux after fresh C addition. D/W cycles breakdown soil macroaggregates and release labile particulate organic matter (fPOM) that was formerly protected. Increasing number of D/W cycles raises microbial respiration from decomposition of new organic matter of the POM fraction rather than using older, more stabilised OM, thus generating a negative PE



horizons after litter removal. Cores were stored at 4 °C and immediately transported to the laboratory of Agricultural Soil Science of Georg-August University of Göttingen, Germany.

Microcosm experiment

CO₂ effluxes were determined during the incubation period of 27 days. This timespan was selected because D/W-induced differences in mineralisation of fPOM were expected directly after the D/W cycles (Schimel and Mikan 2005; Goldberg et al. 2008). The PE and substrate use efficiency (SUE) (for details see below) were assessed to compare the microbial incorporation of the ¹³C-labelled amendment into the new

organic matter. Aggregate size distribution and density fractions from each aggregate class were determined to categorise the SOM pools via different degrees of C protection.

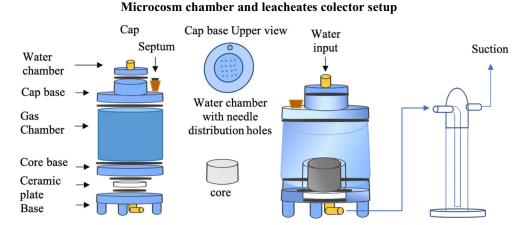
The microcosm experiment consisted of 28 undisturbed core samples (~ 78.5 g dry soil, bulk density 0.8 ± 0.1 Mg m⁻³) pre-incubated for 4 days at field capacity ($0.34 \, \mathrm{m^3 \, m^{-3}}$, $-33 \, \mathrm{kPa}$). Following incubation, the cores were placed on a ceramic pressure plate within a closed acrylic chamber, modified from Poll et al. (2010), and equipped with a septum for gas sampling (Fig. 2). Briefly, approximately 3 mg of $^{13}\mathrm{C}$ uniformly labelled lignocellulose milled residue (maize derived, isotopic purity 97 atm % $^{13}\mathrm{C}$ (IsoLife –Stable Isotope Labelled Plant Products for the Life Sciences,

Table 1 Properties and standard deviation (±) of studied soil (0–8-cm depth)

Variable	Units	Value	
pH water		4.3 ± 0.3	Acid soil (1:2.5 water)
pH CaCl ₂		3.3 ± 0.2	Acid soil (1:2.5)
Soil C	g kg ⁻¹ soil	106 ± 9.9	Total soil carbon at 0-8-cm depth
Soil N	g kg ⁻¹ soil	5.0 ± 0.5	Total soil nitrogen 0-8-cm depth
Soil C:N		21	0–8-cm depth.
Litter C:N		60	Araucaria araucana litter 0-2 cm
Al_p	g kg ⁻¹ soil	6.4 ± 2.2	Pyrophosphate extractable Al
Fe _p	g kg ⁻¹ soil	3.5 ± 1.9	Pyrophosphate extractable Fe
Al_o	g kg ⁻¹ soil	8.7 ± 2.8	Oxalate extractable Al
Fe _o	g kg ⁻¹ soil	6.7 ± 1.4	Oxalate extractable Fe
Si _o	g kg ⁻¹ soil	0.3 ± 0.2	Oxalate extractable Si
Al_d	g kg ⁻¹ soil	12.2 ± 1.6	Dithionite extractable Al
Fe_d	g kg ⁻¹ soil	4.7 ± 0.3	Dithionite extractable Fe
Al _p /Al _o		0.8	> 0.5 organo-mineral nature
$Al_o + 0.5Fe_o$	%	1.3	> 2 andic properties
Al_K	cmol ₊ kg ⁻¹	6.8 ± 2.5	Exchangeable Al
CECe	$cmol_{+} kg^{-1}$	9.0 ± 1.6	Effective cation exchange capacity



Fig. 2 Microcosm chambers (acrylic materials) setup for the drying and rewetting cycles and CO₂ collection. Note: The top of the main chamber has a small additional chamber to which several irrigation needles were connected to apply the irrigation water uniformly



Wageningen, Holland)) were suspended in 10 ml of distilled water and spread uniformly on top of each core using several injections with a syringe. Drying and rewetting cycles consisted of 3 days of drying followed by 3 days of wetting. Dry conditions were achieved using a vacuum pump (Leroy-SomerTM) from the bottom of the ceramic plate, reaching – 80 kPa for 3 h. Rewetting was conducted by watering the core soil on the top and leaving the soil to equilibrate for 30 min until the moisture content reach field capacities. This was achieved using 12 needles connected to another pump (model ISM404B, ISMATECTM). Microcosms received either 1 or 4 cycles. Control soils with labelled lignocellulose additions were subjected to zero D/W cycles and observed alongside the other treatments. In addition to determining the natural isotopic abundance of ¹³C, moist soil cores without labelled lignocellulose additions were also incubated. All treatments were replicated four times.

CO₂ sampling

 ${\rm CO_2}$ gas samples were collected during 27 days of incubation from the first day of each drying or rewetting (12 h apart) period and thereafter, with one sample collected for each day until the next drying. All samples were collected via a 10-ml syringe through the septum on top of the microcosm container (Fig. 2). The gas samples were injected into a vacutainer (Exetainer, Labco Limited, 12 ml) and stored at 5 °C until measurement. After sampling, each acrylic flask was ventilated with ${\rm CO_2}$ -free air. At the end of the 27-day incubation period, the soil was carefully extracted from each core for further analyses.

Aggregate size classes

Aggregate size distribution was determined by dry sieving. Soil was air-dried at 40 °C and sieved through 250 μm and 53 μm meshes on the Vibratory Sieve Shaker AS 200 (Retsch, Germany) for 5 min, at an amplitude of 1.5 mm. Three

aggregate size classes were obtained: macroaggregates (> $250 \mu m$), microaggregates ($250-53 \mu m$), and silt + clay size particles (< $53 \mu m$). The D/W cycles impact soil aggregate turnover, and differences in aggregate size composition between soils with 1 and 4 cycles and soils with constant moisture (0 cycle) were regarded as the proportional effects of the D/W cycles.

Organic matter density fraction

Organic matter fractions were obtained by density fractionation from each aggregate size class using sodium polytungstates (SPT) (Gunina and Kusyakov 2014). Three OM density fractions were obtained, dried at 40 °C, and weighed: light fraction (fPOM, < 1.6 g cm⁻³), occluded fraction (oPOM, 1.6–2.0 g cm⁻³), and heavy fraction (Hf> 2.0 g cm⁻³). The effect of D/W on the gain (negative values) or loss (positive values) of aggregate mass and its associated C was obtained using the difference between the 0 cycle, which received labelled residue but no D/W cycling, and the 1 cycle or 4 cycle treatments. For the aggregate calculations, the same subtraction for the proportional change in the OM density fractions was utilised.

Priming effect

The priming effect (PE) was calculated as defined by Guenet et al. (2010):

$$PE = \left(\frac{A_{\text{lignin}} - A_{\text{sample}}}{A_{\text{lignin}} - A_{\text{soil}}}\right) \times Q_{\text{sample}} - Q_{\text{soil}}$$
(1)

where $A_{\rm lignin}$, $A_{\rm sample}$, and $A_{\rm soil}$ represent the isotopic abundance of 13 C-lignocellulose residue added to the soil, the isotopic abundance of the $\rm CO_2$ from the amended soil core sample with labelled lignocellulose, and the isotopic abundance of the $\rm CO_2$ from non-amended (natural) soil core sample, respectively. $Q_{\rm sample}$ and $Q_{\rm soil}$ represent the quantity of released $\rm CO_2$ in the microcosm headspace of freshly C amended soil and the



CO₂ in the headspace of non-amended soil, respectively. Equation 1 was used to calculate the priming of SOM induced by the amendment of lignocellulose. The D/W cycles were assumed to release fPOM, which primarily consists of lignocellulose. Therefore, the difference in PE between soil with D/W cycles and soil with no D/W cycles allowed us to quantify the effect of D/W cycles on priming (PE_c).

Soil analyses

Soil C and nitrogen (N) contents were determined by dry combustion using a CN Elemental analyser (CHN NA 1500, Carlo Erba). Microbial biomass C (MB-C) was determined by the difference in extractable C in 0.5 M K₂SO₄ of fumigated (chloroform free of ethanol) and unfumigated soils and multiplied by the factor 2.64, used by Vance et al. (1987a, 1987b).

¹³C/¹²C isotope ratio

The carbon isotope ratios (13 C/ 12 C) of all fractions, CO2, MB-C, and bulk soil samples were measured at the Centre for Stable Isotope Research and Analysis (KOSI) of Georg-August-University of Göttingen, Germany. The CO2 concentration and the carbon-isotope ratio were measured in a gas chromatograph combustion isotope ratio mass spectrometer (GC-C-IRMS). Soil C contents were measured using an elemental analyser (Vario EL II, Germany), and the isotopic ratio was measured using an elemental analyser in dual-element analysis mode (Carlo Erba 1108, Milano, Italy). The C isotope ratio was expressed relative to the international Pee Dee Belemnite (PDB) limestone standard as δ^{13} C.

Substrate use efficiency

The SUE was calculated at the end of the incubation as the ratio between labelled microbial biomass ($^{13}C_B$), $^{13}CO_2$ respired, and $^{13}C_B$ (Spohn and Chodak 2015):

$$SUE = \frac{^{13}C_B}{^{13}CO_2 + ^{13}C_B}$$
 (2)

where SUE estimates the relative proportion of the labelled MB-C to respiration.

Statistical analysis

Two-way ANOVA was performed to analyse the effects of temperature and D/W cycles on CO₂ efflux, PE, aggregate size, OM-C fraction, and microbial biomass-C and its ¹³C-lignocellulose distribution. The normality of the variances was tested by the Shapiro-Wilk test, and the homogeneity of variance was tested by Levene's test. The data abnormally

distribution was log transformed until comparison data presented similar variance. Least significant difference (LSD) and post hoc Tukey tests (p < 0.05) were performed to compare mean values between variables. All analyses were conducted using SPSS statistical software v23.0.0.0 (SPSS Inc., Chicago, IL, USA). Figures were developed with DataGraph 4.3 Visual Data Tools, 2006–2018, Inc.

Results

Soil weight and C recovery (%) following dry sieving varied between 92 and 99%, respectively. The recovery of soil-labelled ¹³C fluctuated between 6 and 52% and varied between 5 and 39% in the ¹³C MB-C. Leachates were minimal and fluctuated between 0 and 7% (Table 2). The total recovered lignocellulose-derived labelled ¹³C ranged from 73 to 99% (Table 2).

Aggregate size classes

The D/W-induced change in the distribution of aggregate size classes and their C contents was obtained by subtracting D/W 0 cycle results from the aggregate size class abundance from that of the 1 or 4 D/W cycle treatment (Fig. 3). Macroaggregates (> 250 µm) were the most abundant aggregate size class in the investigated soils (609–785 g kg⁻¹), followed by microaggregates (250-53 µm) (201-308 g kg⁻¹) and silt + clay particles (< 53 μ m) (12– 23 g kg⁻¹) (Table S1, Supplementary Materials). Drying and rewetting had minimal influence on the mass of the aggregate size classes and their C content at 5 °C (Fig. 3a). At 25 °C, however, macroaggregate weight (Fig. 3b) and labelled C (Fig. 3f) increased (positive value) after 1 D/W cycle, and no significant differences were detected for 4 D/W cycles (Table S4, Supplementary Materials). The same was true for labelled C at 5 °C (Fig. 3e).

Density fractionation

Drying and rewetting cycles did influence the quantity of organic C, including labelled C (Fig. 4; Tables S2, S5, Supplementary Materials). The interaction between temperature and D/W cycles influenced the distribution of C and lignocellulose-derived ¹³C among the various OM fractions (Tables S3, S5, supplementary Materials). The quantity of D/W cycles did not have significant effects at 5 °C, but the total C and lignocellulose-derived ¹³C content always increased with 1 cycle and decreased with 4 cycles at the expense of heavy fraction, which lost the respective mass or C (Fig. 4).



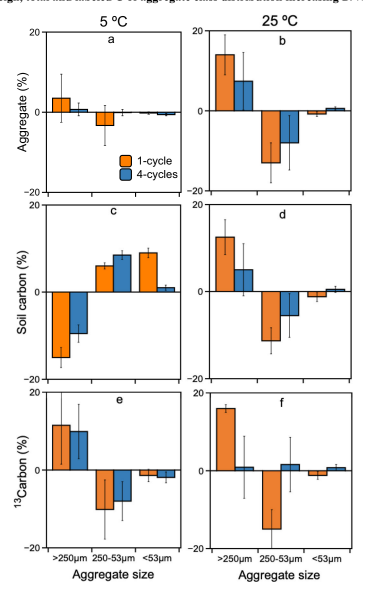
Table 2 Total recovery (%) of soil weight, soil C, and labelled C after dry sieving

	Weight	Soil C	NB- ¹³ C- ¹	$MB-^{13}C^2$	$^{13}\mathrm{CO}_2$	¹³ C- leached	Total
5 °C							
0 cycle	97	95	21	39	13	0	73
1 cycle	99	92	32	20	33	3	88
4 cycles	100	106	47	20	17	5	89
25 °C							
0 cycle	93	92	48	5	43	0	96
1 cycle	99	117	52	14	33	0.3	99
4 cycles	92	72	6.4	26	60	7	99

¹ Non-biomass

Fig. 3 Proportional change effect of aggregate size classes (macroaggregates $> 250 \mu m$, microaggregates 250-53 µm, and silt + clay size $< 53 \mu m$) obtained by subtracting zero cycles (no cycle + labelled residue) to 1 cycle D/W or 4 cycles D/W. Soils amended with lignocellulose are displayed after 27 days of incubation at 5 °C (left) and 25 °C (right), whereas relative weight (a and b), total C content (c and d), and lignocellulose-derived $^{13}\mathrm{C}$ incorporation (e and f) of the aggregate size classes is shown. Bars indicate standard errors of the means

Weigh, total and labeled C of aggregate class distribution increasing D/W cycles

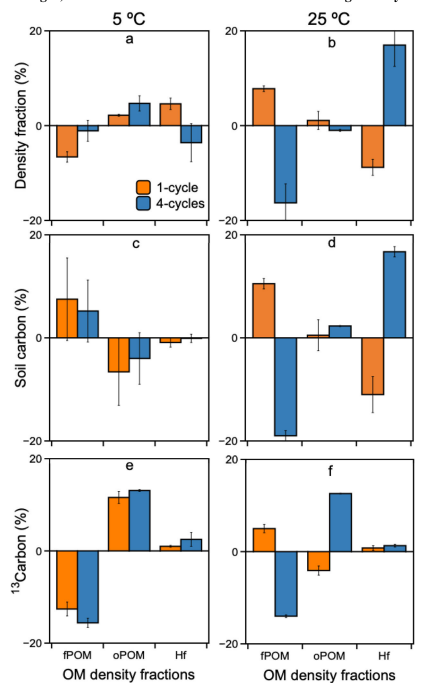




² Microbial biomass

Fig. 4 Proportional change effect of organic matter particles (POM) from the entire soil; OM fraction from the different aggregates was reunited as light fraction < 1.6 g cm^{-3} (fPOM), occluded fraction 1.6-2.0 g cm-3 (oPOM), and heavy fraction $> 2.0 \text{ g cm}^{-3}$ (Hf) obtained by subtracting the zero cycles (no cycle + labelled residue) to 1 cycle D/W or 4 cycle D/W. Soils amended with lignocellulose after 27 days of incubation at 5 °C and 25 °C are presented regarding the relative weight of the OM fraction (a and **b**), their total C content (**c** and **d**), and their lignocellulose-derived ¹³C incorporation (**e** and **f**). Bars indicate standard errors of the means

Weight, total and labeled C of OM fractions increasing D/W cycles



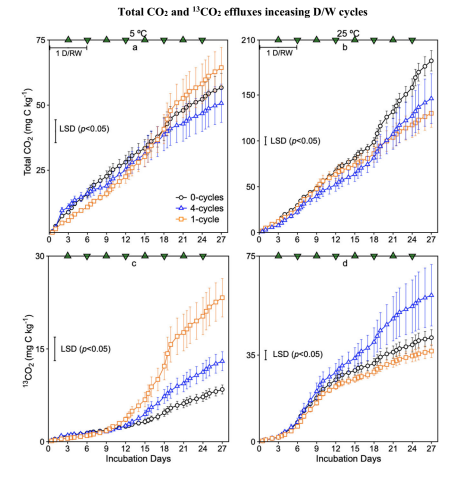
CO₂ effluxes

Soil respiration was responsive to temperature, as demonstrated by the accumulated total $\rm CO_2$ and labelled $^{13}\rm CO_2$ efflux in the undisturbed cores (Table S6, Supplementary Materials). On average, the total amount of $\rm CO_2$ released at 25 °C was approximately three times that released at 5 °C (Fig. 5). The mineralisation of lignocellulose-derived $^{13}\rm C$ was roughly 2.5 times higher at 25 °C than that at 5 °C.

D/W cycle effects were isolated at each temperature by one-way ANOVA. After day 18, the total CO_2 efflux was significantly higher for soils exposed to 1 D/W cycle than those exposed to 4 cycles or provided with constant moisture content (0 cycle) at 5 °C (p < 0.05) (Fig. 5a). Lignocellulose mineralisation displayed the same pattern as the total CO_2 ; it was higher in soils exposed to only a single D/W cycle, compared with those experiencing 0 or 4 cycles (Fig. 5c). Soil Incubated at 25 °C with no D/W cycles had a higher total



Fig. 5 Total CO₂ evolved during 27 days of incubation at 5 °C (a) and 25 °C (b) from soil with lignocellulose addition and D/W (0 cycle, 1 cycle, and 4 cycles). Dry period (triangle) started on day 3 and continued for another 3 days of incubation. The wet period (inverted triangle) started on day 6 until the next drying. ¹³CO₂ efflux through 27 days of incubation at 5 °C (c) and 25 °C (d). Small bars on the data point indicate standard errors of the mean. Large bars indicate the least significant differences (LSD) (p < 0.05)



 ${\rm CO_2}$ efflux than those of soils with 1 or 4 D/W cycles after 18 days of incubation (p < 0.05) (Fig. 5b). In the warmer soil, the release of lignocellulose $^{13}{\rm C}$ was the highest when exposed to 4 D/W cycles (cf. Fig. 5 b and d).

Priming effect

The PE response varied with the temperature, and differences between D/W treatments began to become evident after 18 days of incubation (Fig. 6). Only the 0 D/W cycle soil at 5 °C showed a positive PE, although it was not significantly different from zero PE; soils with D/W cycles showed a negative PE, and the soil with 4 D/W cycles was the only treatment significantly different from zero at the end of the incubation time (Fig. 6a). At 25 °C; however, the PE was always negative and soils exposed to D/W, regardless of the number of cycles, showed the most negative values (p < 0.05) (Fig. 6b). At 5 °C PEc, the differences between soil with 1 or 4 D/W cycles and 0 cycle were significant between 0 and 18 days and at the end of the incubation (Fig. 6c). However, at 25 °C the differences were expressed from day 9 and were not perceptible at the end of the incubation (Fig. 6d).

Microbial biomass and substrate use efficiency

Temperature and D/W cycles had significant impacts on microbial biomass 13 C incorporation (cf., Tables 2 and 3) and SUE (Table 3; Fig. 7; and Table S7, Supplementary Materials). High SUE occurred preferentially under lower temperatures and was on average two times higher (p < 0.05) at 5 °C than at 25 °C (Fig. 7). Drying and rewetting had a decreased effect on SUE values at 5 °C (Fig. 7a), but at 25 °C, D/W cycles did not induce any significant effects on SUE (Fig. 7b).

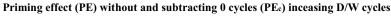
Discussion

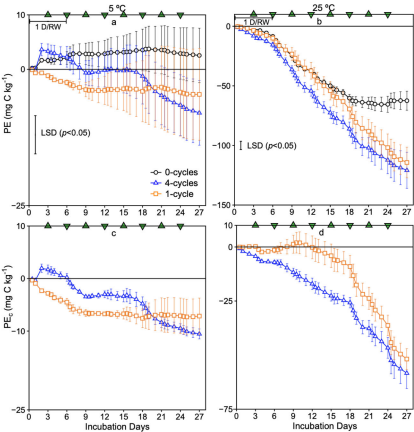
Aggregates and particulate organic matter

Physical protection of SOM by aggregates is an important mechanism for C stabilisation; differentiating the degree of physicochemical protection afforded to the SOM by various aggregate size classes remains challenging. Drying and rewetting cycles were investigated in terms of their disruptive effects on the degradation and formation of macroaggregates (> 250 µm) and microaggregates at (250—53 µm) (i.e. the



Fig. 6 Priming effect (PE) through 27 days of incubation at 5 °C (a) and 25 °C (b) for soil with lignocellulose addition. Dry period (triangle) started on day 3 and continued for another 3 days of incubation. The wet period (inverted triangle) started on day 6 until the next drying. Relative priming effect as affected by drying and rewetting (PEc), calculated by substratcing the zero cycles (no cycle + labelled residue) to 1 cycle D/W or 4 cycles D/W, is shown for 27 days of incubation at 5 °C (c) and 25 °C (d) for soil with lignocellulose addition. Small bars on the data point indicate standard errors of the mean. Large bars indicate the least significant differences (LSD) (p < 0.05)





macroaggregate turnover) and the associated C dynamics in the respective aggregate size classes found in a temperate forest. Soil with D/W cycles experienced accelerated macroaggregate turnover at the expense of the microaggregates size class, particularly at warmer temperatures of 25 °C (Fig. 3b).

Table 3 Total microbial biomass C (MB-C), MB- 13 C, and standard error of the mean (\pm) of four replicates. Different lowercase letters in each column and within each temperature indicate significant differences (p < 0.05). Different capital letters in each column and between temperatures indicate significant differences (p < 0.05)

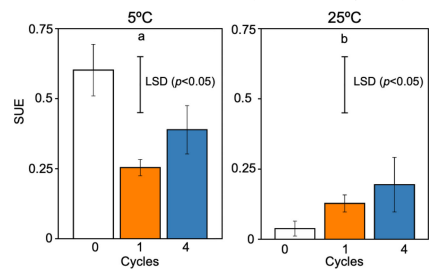
Drying and rewetting	Microbial biomass C			
	Total MB-C (mg C kg ⁻¹)	MB- ¹³ C		
5 °C				
0 cycle	$456\pm61aA$	$15 \pm 4.0 \text{ aA}$		
1 cycle	$372\pm28~aB$	$7.5\pm1.0~b\mathrm{B}$		
4 cycles	$346 \pm 51 \text{ aB}$	$7.6 \pm 3.0 \text{ bB}$		
25 °C				
0 cycle	$317 \pm 48 \text{ aC}$	$2.0\pm1.5~bD$		
1 cycle	$313 \pm 25 \text{ aC}$	$5.4 \pm 1.1 \ aC$		
4 cycles	$345 \pm 57 \text{ aBC}$	9.8 ± 4.0 aB		

Generally speaking, we found strongly contrasting effects between the two temperature treatments, indicating a systematic process underlying influence of temperature. Although we did not determine the microbial community composition, fungal growth could be significantly reduced at low temperatures and fungal hyphae development could be stunted at higher temperatures. Fungi can function as binding agents in soil (Denef et al. 2001a), and at high temperatures, fungi could represent a significant portion of the microbial biomass and could have key function in building and stabilising macroaggregates (Denef et al. 2001a). After 27 days of incubation at 25 °C, the proportion of microaggregates' weight (Fig. 3b), their C content (Fig. 3d), and lignocellulose-derived ¹³C (Fig. 3f) decreased in soils exposed after 4 D/W cycles, but the effect was not observed after 1 D/W cycle (Fig. 3b-f). Increasing the number of D/W events could therefore have a negative impact on microaggregate formation, while novel protected OM within macroaggregates could contribute to its formation. Density fractionation indicated a depletion of the fPOM and an increase in Hf, supporting the acceleration of macroaggregate turnover (Fig. 4b, d) and increasing the ¹³C fraction in the oPOM (Fig. 4f). Microaggregates were depleted by accelerated turnover and the formation of new macroaggregates, which only occurred in the short term under enhanced D/W cycles



Fig. 7 Substrate use efficiency (SUE) of 13 C-lignocellulose at 5 °C (a) and 25 °C (b) estimated after 27 days of incubation of the D/W treatments, 0 cycle, 1 cycle, and 4 cycles. Small bars indicate standard errors of the mean. Large bars indicate the least significant differences (LSD) (p < 0.05)

Microbial substrate use efficiency (SUE) inceasing D/W cycles



(Denef et al. 2001a; Dorodnikov et al. 2011; Gunina and Kusyakov 2014).

CO₂ efflux

Drying and rewetting cycles affected the cumulative C mineralisation at 25 °C. The total CO₂ emitted throughout the 27 days of incubation was strongly reduced compared to the CO₂ produced by soil at optimum moisture content (0 cycle). Thus, rewetting does not compensate for the lower mineralisation during drought periods but rather has a medium-term impact (Fig. 5). Cumulative mineralisation is linked to the intensity and duration of drying and the accessible pool of organic C during drying and rewetting (Borken and Matzner 2009). Therefore, under field conditions, increasing droughts could result in reduced C mineralisation, whereas increasing spring/summer precipitation could accelerate C losses from physically protected organic matter. In our laboratory study, drying and rewetting cycles caused higher microbial respiration of new organic matter added to the soil (Fig. 5). This effect for ¹³CO₂ became evident after 15 days of incubation at both temperatures (Fig. 5c and d). Araucaria araucana litter fall displayed a C:N ratio of 60 (Table 1) with a high lignin: N ratio (70–90), which is generally associated with a low mineralisation rate (Diehl et al. 2003; Bertiller et al. 2006). The lignocellulose-rich OM decomposed at a rate between 50 and 150 mg C kg soil⁻¹ day⁻¹, comparable to other rates determined through similar incubations of Chilean forest soils (Matus et al. 2008; Muñoz et al. 2016) or Mediterranean forest soils (Almagro et al. 2009; Guntiñas et al. 2013). Despite the recalcitrance of the fresh substrate, there was significant C mineralisation from this material, supported by the negative PE value (Fig. 6d).

Priming effect

At 5 °C, there was a small but significant PE induced by the lignocellulose amendment, while at 25 °C a stronger negative PE was produced. The PE was negatively correlated with respired 13 C (r = -0.59, p < 0.04, n = 12). This result indicates that when less organic C is consumed from native SOM by microorganisms (negative PE), more fresh ¹³C is mineralised. This correlation clearly demonstrates the preferential C use of the substrate, which is, in this study, a representative compound of the soil's fPOM fraction. These results were also supported by the PE_c, the difference in PE between 1 or 4 cycles and 0 cycle (Fig. 6). In other studies, where complex OM was added in the form of leaf and stem residues (¹³C-labelled wheat residues), the results also showed intensive mineralisation of the added OM, yielding a negative PE for extended periods (Shahbaz et al. 2017); with the addition of other materials, a temporary negative PE (up to 40 days) was observed (Wang et al. 2015), likely due to preferential utilisation of the added substrate and thus a pool substitution (Shahbaz et al. 2017). This mechanism was effectively revealed by the SUE, particularly at 5 °C (Fig. 7), because the lignocellulose was incorporated into the microbial biomass by a well-adapted microbial community (Borken and Matzner 2009). Soil microbial communities are resilient and able to quickly recover after wetting in soils with high SOC stock (Canarini et al. 2017). At 25 °C, however, lignocellulosederived C could be invested for highly energy demanding processes, e.g. for enzyme synthesis, and less C was converted into structural cell components (Ågren and Bosatta 1987; Blagodatsky et al. 2000; Davidson and Janssens 2006; Manzoni et al. 2012). This finding is in accordance with other studies, which obtained similar results, i.e. a decreasing C use efficiency with increasing temperature between 2 and 28 °C



(Manzoni et al. 2012; Qiao et al. 2019). This phenomenon can likely be associated with a change in metabolism rather than a change in the microbial community structure (Manzoni et al. 2012; Bölscher et al. 2017; Di Lonardo et al. 2017). When the number of D/W cycles was increased on undisturbed soil, fPOM was negatively affected by warming and the acceleration of aggregate turnover. Warming and aggregate turnover could have significant implications for temperate forests facing global change. Faster macroaggregate turnover and depletion of fPOM may prevent large stabilised SOM from decomposing, leading to a lack of C stabilisation after repeated D/W cycles.

Conclusions

One drying-rewetting (D/W) cycle disrupted approximately 15% of the aggregates after 27 days. The fine particulate fraction of organic matter (fPOM, < 1.6 g cm⁻³) released from aggregates disrupted by D/W contained the most physically exposed C (microbially available) as compared to the other C pools. This fPOM, which we traced by added ¹³C-labelled lignocellulose, was decomposed preferentially, covering microbial energy demand but not converted to microbial biomass and therefore not contributing to long-term stabilisation in the form of necromass residues. Increasing the number of D/W cycles caused a negative priming effect (PE), i.e. preferential utilisation of the added ¹³C-labelled lignocellulose released by D/W cycles rather over other OM fractions. A new pathway is proposed for C release by aggregate disruption, resulting in negative PE due to D/W cycling. This scenario was reflected by low substrate use efficiency (SUE), i.e. microbes preferentially respiring the accessible fPOM, particularly at a high temperature. Consequently, the D/W cycles at 25 °C significantly increased microbial activity, which primarily regulated fPOM decomposition.

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References

- Adu JK, Oades JM (1978) Physical factors influencing decomposition of organic materials in soil aggregates. Soil Biol Biochem 10:109–115. https://doi.org/10.1016/0038-0717(78)90080-9
- Ågren GI, Bosatta E (1987) Theoretical analysis of the long-term dynamics of carbon and nitrogen in soils. Ecology 68:1181–1189. https://doi.org/10.2307/1939202
- Almagro M, López J, Querejeta JI, Martínez-Mena M (2009) Temperature dependence of soil CO₂ efflux is strongly modulated by seasonal patterns of moisture availability in a Mediterranean ecosystem. Soil Biol Biochem 41:594–605. https://doi.org/10.1016/j.soilbio.2008.12.021
- Appel T (1998) Non-biomass soil organic N—the substrate for N mineralization flushes following soil drying–rewetting and for organic N rendered CaCl₂-extractable upon soil drying. Soil Biol Biochem 30: 1445–1456. https://doi.org/10.1016/S0038-0717(97)00230-7
- Balser T, Firestone M (2005) Linking microbial community composition and soil processes in a California annual grassland and mixed-conifer forest. Biogeochemistry 73:395–415. https://doi.org/10.1007/s10533-004-0372-y
- Bernhard N, Moskwa L-M, Schmid K, Oeser RA, Aburto F, Bader M, Baumann K, von Blanckenburg F, Boy J, van den Brink L, Brucker E, Büdel B, Canessa R, Dippold M, Ehlers T, Fuentes JP, Godoy R, Jung P, Karsten U, Köster M, Kuzyakov J, Leinweber P, Neidhardt H, Matus F, Carsten Mueller CM, Oelmann Y, Oses R, Osses P, Paulino L, Samolov E, Schaller M, Schmid M, Spielvogel S, Spohn M, Stock S, Stroncik N, Tielbörger K, Übernickel K, Scholten T, Seguel O, Wagner D, Kühn P (2018) Pedogenic and microbial interrelations to regional climate and local topography: new insights from a climate gradient (arid to humid) along the Coastal Cordillera of Chile. Catena 170:335–355. https://doi.org/10.1016/j.catena.2018.06.018
- Bertiller MB, Mazzarino MJ, Carrera AL, Diehl P, Satti P, Gobbi M, Sain CL (2006) Leaf strategies and soil N across a regional humidity gradient in Patagonia. Oecologia 148:612–624. https://doi.org/10.1007/s00442-006-0401-8
- Billings SA, Ballantyne F IV (2013) How interactions between microbial resource demands, soil organic matter stoichiometry, and substrate reactivity determine the direction and magnitude of soil respiratory responses to warming. Glob Chang Biol 19:90–102. https://doi.org/10.1111/gcb.12029
- Bingemann CW, Varner JE, Martin WP (1953) The effect of the addition of organic materials on the decomposition of an organic soil. Soil Sci Soc Am J 17:34–38. https://doi.org/10.2136/sssaj1953.03615995001700010008x
- Birch H (1958) The effect of soil drying on humus decomposition and nitrogen availability. Plant Soil 10:9–31. https://doi.org/10.1007/BF01343734
- Blagodatsky S, Heinemeyer O, Richter J (2000) Estimation the active and total soil microbial biomass by kinetic respiration analysis. Biol Fertil Soils 32:73–81. https://doi.org/10.1007/s003740000219
- Bölscher T, Paterson E, Freitag T, Thornton B, Herrmann AM (2017) Temperature sensitivity of substrate-use efficiency can result from altered microbial physiology without change to community composition. Soil Biol Biochem 109:59–69. https://doi.org/10.1016/j.soilbio.2017.02.005



- Borken W, Matzner E (2009) Reappraisal of drying and wetting effects on C and N mineralization and fluxes in soils. Glob Chang Biol 15: 808–824. https://doi.org/10.1111/j.1365-2486.2008.01681.x
- Canarini A, Pødenphant Kiær L, Dijkstra FA (2017) Soil carbon loss regulated by drought intensity and available substrate: a meta-analysis. Soil Biol Biochem 112:90–99. https://doi.org/10.1016/j. soilbio.2017.04.020
- Christensen BT (1992) Physical fractionation of soil and organic matter in primary particle size and density separates. In: Stewart B.A. (eds) Advances in soil science. Advances in soil science, vol 20. Springer, New York, pp 1–90. https://doi.org/10.1007/978-1-4612-2930-8 1
- Corti G, Ugolini FC, Agnelli A, Certini G, Cuniglio R, Berna F, Fernández Sanjurjo MJ (2002) The soil skeleton, a forgotten pool of carbon and nitrogen in soil. Eur J Soil Sci 53:283–298. https://doi. org/10.1046/j.1365-2389.2002.00442.x
- Davidson EA, Janssens IA (2006) Temperature sensitivity of soil carbon decomposition and feedbacks to climate change. Nature 440:165– 173. https://doi.org/10.1038/nature04514
- Denef K, Six J, Bossuyt H, Frey SD, Elliott ET, Merckx R, Paustian K (2001a) Influence of dry- wet cycles on the interrelationship between aggregate, particulate organic matter, and microbial community dynamics. Soil Biol Biochem 33:1599–1611. https://doi.org/10.1016/S0038-0717(01)00076-1
- Denef K, Six J, Paustian K, Merckx R (2001b) Importance of macroaggregate dynamics in controlling soil carbon stabilization: short-term effects of physical disturbance induced by dry-wet cycles. Soil Biol Biochem 33:2145–2153. https://doi.org/10.1016/S0038-0717(01) 00153-5
- Di Lonardo DP, De Boer W, Klein Gunnewiek PJA, Hannula SE, Van der Wal A (2017) Priming of soil organic matter: chemical structure of added compounds is more important than the energy content. Soil Biol Biochem 108:41–54. https://doi.org/10.1016/j.soilbio.2017.01. 017
- Diehl P, Mazzarino MJ, Funes F, Fontenla S, Gobbi M, Ferrari J (2003) Nutrient conservation strategies in native Andean-Patagonian forests. J Veg Sci 14:63–70. https://doi.org/10.1658/1100-9233(2003) 014[0063:NCSINA]2.0.CO:2
- Dorodnikov M, Kuzyakov Y, Fangmeier A, Wiesenberg G (2011) C and N in soil organic matter density fractions under elevated atmospheric CO₂: turnover vs. stabilization. Soil Biol Biochem 43:579–589. https://doi.org/10.1016/j.soilbio.2010.11.026
- Fernández DP, Neff JC, Belnap J, Reynolds RL (2006) Soil respiration in the cold desert environment of the Colorado Plateau (USA): abiotic regulators and thresholds. Biogeochemistry 78:247–265. https://doi. org/10.1007/s10533-005-4278-0
- Fontaine S, Henault C, Aamor A, Bdioui N, Bloor JMG, Maire V, Nary B, RevaillotS MP (2011) Fungi mediate long term sequestration of carbon and nitrogen in soil through their priming effect. Soil Biol Biochem 43:86–96. https://doi.org/10.1016/j.soilbio.2010.09.017
- Garcia-Pausas J, Paterson E (2011) Microbial community abundance and structure are determinants of soil organic matter mineralization in the presence of labile carbon. Soil Biol Biochem 43:1705–1713. https://doi.org/10.1016/j.soilbio.2011.04.016
- Garreaud R, Alvarez-Garreton C, Barichivich J, Boisier JP, Christie DA, Galleguillos M, Le Quesne C, McPhee J, Zambrano-Bigiarini M (2017) The 2010–2015 mega drought in Central Chile: impacts on regional hydroclimate and vegetation. Hydrol Earth Syst Sci 21: 6307–6327. https://doi.org/10.5194/hess-2017-191
- Goldberg SD, Muhr J, Borken W, Gebauer G (2008) Fluxes of climaterelevant trace gases between a Norway spruce forest soil and atmosphere during repeated freeze-thaw cycles in mesocosms. J Plant Nutr Soil Sc 171:729–739. https://doi.org/10.1007/s10533-009-9294-z
- Gregorich EG, BeareMH MKUF, Skjemstad JO (2006) Chemical and biological characteristics of physically uncomplexed organic matter.

- Soil Sci Soc Am J 70:975–985. https://doi.org/10.2136/sssaj2005.
- Guenet B, Danger M, Abbadie L, Lacroix G (2010) Priming effect: bridging the gap between terrestrial and aquatic ecology. Ecology 91: 2850–2861. https://doi.org/10.1890/09-1968.1
- Gunina A, Kusyakov Y (2014) Pathways of litter C by formation of aggregates and SOM density fractions: implications from ¹³C natural abundance. Soil Biol Biochem 71:95–104. https://doi.org/10.1016/j.soilbio.2014.01.011
- Guntiñas ME, Gil-Sotres F, Leirós MC, Trasar-Cepeda C (2013) Sensitivity of soil respiration to moisture and temperature. J Soil Sci Plant Nut 13:445–461. https://doi.org/10.4067/S0718-95162013005000035
- Hibbard KA, Law BE, Reichstein M, Sulzman J (2005) An analysis of soil respiration across northern hemisphere temperate ecosystems. Biogeochemistry 73:29–70. https://doi.org/10.1007/s10533-004-2946-0
- Jarvis P, Rey A, Petsikos C, Wingate L, Rayment M, Pereira J, Banza J, David J, Miglietta F, Borghetti M, Manca G, Valentini R (2007) Drying and wetting of Mediterranean soils stimulates decomposition and carbon dioxide emission: the "Birch effect". Tree Physiol 27: 929–940. https://doi.org/10.1093/treephys/27.7.929
- Jenkinson DS, Fox RH, Rayner JH (1985) Interactions between fertilizer nitrogen and soil nitrogen-the so-called 'priming' effect. J Soil Sci 36:425–444. https://doi.org/10.1111/j.1365-2389.1985.tb00348.x
- Kim DG, Vargas R, Bond-Lamberty B, Turetsky MR (2012) Effects of soil rewetting and thawing on soil gas fluxes: a review of current literature and suggestions for future research. Biogeosciences 9: 2459–2483. https://doi.org/10.5194/bg-9-2459-2012
- Kuzyakov Y (2010) Priming effects: interactions between living and dead organic matter. Soil Biol Biochem 42:1363–1371. https://doi.org/10.1016/j.soilbio.2010.04.003
- Kuzyakov Y, Friedel JK, Stahr K (2000) Review of mechanisms and quantification of priming effects. Soil Biol Biochem 32:1485– 1498. https://doi.org/10.1016/S0038-0717(00)00084-5
- Lal R (2004) Soil carbon sequestration impacts on global climate change and food security. Science 304:1623–1627. https://doi.org/10.1126/ science.1097396
- Magid J, Kjærgaard C (2001) Recovering decomposing plant residues from the particulate soil organic matter fraction: size versus density separation. Biol Fertil Soils 33:252–257. https://doi.org/10.1007/s003740000316
- Manzoni S, Schimel JP, Porporato A (2012) Responses of soil microbial communities to water stress: results from a meta-analysis. Ecology 93:930–938. https://doi.org/10.1890/11-0026.1
- Matus F, Lusk H, Maire C (2008) Effects of soil texture, carbon input rates, and litter quality on free organic matter and nitrogen mineralization in Chilean rain forest and agricultural soils. Comm in Soil Sci and Plant Anal 39:187–201. https://doi.org/10.1080/00103624. 2017.1395447
- Meisner A, Rousk J, Bååth E (2015) Prolonged drought changes the bacterial growth response to. Re-wetting. Soil Biol Biochem 88: 314–322. https://doi.org/10.1016/j.soilbio.2015.06.002
- Mikha MM, Rice CW, Milliken GA (2005) Carbon and nitrogen mineralization as affected by drying and wetting cycles. Soil Biol Biochem 37:339–347. https://doi.org/10.1016/j.soilbio.2004.08.003
- Muñoz C, Cruz B, Rojo F, Campos J, Casanova M, Doetterl S, Boeckx P, Zagal E (2016) Temperature sensitivity of carbon decomposition in soil aggregates along a climatic gradient. J Soil Sci Plant Nut 16: 461–476. https://doi.org/10.4067/S0718-95162016005000039
- Oades JM (1984) Soil organic matter and structural stability: mechanisms and implications for management. Plant Soil 76:319–337. https:// doi.org/10.1007/BF02205590
- Oades JM (1988) The retention of organic matter in soils. Biogeochemistry 5(1):35–70. https://doi.org/10.1007/bf02180317



- Oeser R, Stroncik N, Moskwa L, Bernhard N, Schaller M, Canessa R, van den Brink L, Köster M, Brucker E, Stock S, Fuentes JP, Godoy R, Matus F, Oses Pedraza R, Osses McIntyre P, Paulino L, Seguel O, Bader MY, Boy J, Dippold M, Ehlers T, Kühn P, Kuzyakov Y, Leinweber P, Scholten T, Spielvogel S, Spohn M, Übernickel K, Tielbörger K, Wagner D, von Blanckenburg F Von (2018) Chemistry and microbiology of the critical zone along a steep climate and vegetation gradient in the Chilean Coastal Cordillera. Catena 170:183–203. https://doi.org/10.1016/j.catena.2018.06.002
- Oztas T, Fayetorbay F (2003) Effect of freezing and thawing processes on soil aggregate stability. Catena 52:1–8. https://doi.org/10.1016/S0341-8162(02)00177-7
- Poll C, Pagel H, Devers-Lamrani M, Martin-Laurent F, Ingwersen J, Streck T, Kandeler E (2010) Regulation of bacterial and fungal MCPA degradation at the soil-litter interface. Soil Biol Biochem 42:1879–1887. https://doi.org/10.1016/j.soilbio.2010.07.013
- Qiao Y, Wang J, Liang G, Du Z, Zhou J, Zhu C, Huang K, Zhou X, Luo Y, Yan L, Xia J (2019) Global variation of soil microbial carbon-use efficiency in relation to growth temperature and substrate supply. Sci Rep 9:5621. https://doi.org/10.1038/s41598-019-42145-6
- Schimel JP, Mikan C (2005) Changing microbial substrate use in Arctic tundra soils through a freeze-thaw cycle. Soil Biol Bioch 37:1411– 1418. https://doi.org/10.1016/j.soilbio.2004.12.011
- Shahbaz M, Kuzyakov Y, Sanaullah M, Heitkamp F, Zelenev V, Kumar A, Blagodatskaya E (2017) Microbial decomposition of soil organic matter is mediated by quality and quantity of crop residues: mechanisms and thresholds. Biol Fertil Soils 53:287–301. https://doi.org/10.1007/s00374-016-1174-9
- Shi Z, Thomey ML, Mowll W, Litvak M, Brunsell NA, Collins SL, Luo Y (2014) Differential effects of extreme drought on production and respiration: synthesis and modeling analysis. Biogeosciences 11: 621–633. https://doi.org/10.5194/bg-11-621-2014
- Sinha T, Cherkauer KA (2010) impacts of future climate change on soil frost in the midwestern United States. J Geophys Res-Atmos 115:1– 16. https://doi.org/10.1029/2009JD012188
- Six J, Elliott ET, Paustian K (2000) Soil macroaggregate turnover and microaggregate formation: a mechanism for C sequestration under no-tillage agriculture. Soil Biol Biochem 32:2099–2103. https://doi. org/10.1016/S0038-0717(00)00179-6
- Soil Survey Staff (2014) Keys to soil taxonomy, Twelfth Edition NRCS USDA
- Spohn M, Chodak M (2015) Microbial respiration per unit biomass increases with carbon-to-nutrient ratios in forest soils. Soil Biol Biochem 81:128–133. https://doi.org/10.1016/j.soilbio.2014.11.008

- Tisdall JM, Oades JM (1982) Organic matter and water-stable aggregates in soils. J Soil Sci 33:141–163. https://doi.org/10.1111/j.1365-2389. 1982.tb01755.x
- Urrutia-Jalabert R, Gonzalez ME, Gonzalez-Reyes A, Lara A, Garreaud R (2018) Climate variability and forest fires in central and south-central Chile. Ecosphere 9(4):e02171. https://doi.org/10.1002/ecs2.2171
- Vance ED, Brookes PC, Jenkinson DS (1987a) Microbial biomass measurements in forest soils: the use of the chloroform fumigation-incubation method in strongly acid soils. Soil Biol Biochem 19: 697–702. https://doi.org/10.1016/0038-0717(87)90050-2
- Vance ED, Brookes PC, Jenkinson DS (1987b) An extraction method for measuring soil microbial biomass C. Soil Biol Biochem 19:703– 707. https://doi.org/10.1016/0038-0717(87)90052-6
- Vargas R, Detto M, Baldocchi DD, Allen MF (2010) Multiscale analysis of temporal variability of soil CO₂ production as influenced by weather and vegetation. Glob Chang Biol 16:1589–1605. https://doi.org/10.1111/j.1365-2486.2009.02111.x
- Vicca S, Bahn M, Estiarte M, van Loon EE, Vargas R, Alberti G, Ambus P, Arain MA, Beier C, Bentley LP, Borken W, Buchmann N, Collins SL, de Dato G, Dukes JC, Escolar C, Fay P, Guidolotti G, Hanson PJ, Kahmen A, Kröel-Dulay G, Ladreiter-Knauss T, Larsen KS, Lellei-Kovacs E, Lebrija-Trejos E, Maestre FT, Marhan S, Marshall M, Meir P, Miao Y, Muhr J, Niklaus PA, Ogaya R, Peñuelas J, Poll C, Rustad LE, Savage K, Schindlbacher A, Schmidt IK, Smith AR, Sotta ED, Suseela V, Tietema A, van Gestel N, van Straaten O, Wan S, Weber U, Janssens IA (2014) Can current moisture responses predict soil CO₂ efflux under altered precipitation regimes? A synthesis of manipulation experiments. Biogeosciences 11:2991–3013. https://doi.org/10.5194/bg-11-2991-2014
- Von Lützow M, Kögel-Knabner I, Ekschmitt K, Flessa H, Guggenberger G, Matzner E, Marschner B (2007) Soil fractionation methods: relevance to functional pools and stabilization mechanisms. Soil Biol Biochem 39:2183–2207. https://doi.org/10.1016/j.soilbio.2007.03.007
- Wang H, Boutton TW, Xu W, Hu G, Jiang P, Bai E (2015) Quality of fresh organic matter affects priming of soil organic matter and substrate utilization patterns of microbes. Sci Rep 5:10102. https://doi.org/10.1038/srep10102

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