The modification of soil organic matter (SOM) decomposition by plant carbon (C) input (priming effect) represents a critical biogeochemical process that controls soil C dynamics. However, the patterns and drivers of the priming effect remain hidden, especially over broad geographic scales under various climate and soil conditions. By combining systematic field and laboratory analyses based on multiple analytical and statistical approaches, we explore the determinants of priming intensity along a 2200 km grassland transect on the Tibetan Plateau. Our results show that SOM stability characterized by chemical recalcitrance and physico-chemical protection explains more variance in the priming effect than plant, soil and microbial properties. High priming intensity (up to 137% of basal respiration) is associated with complex SOM chemical structures and low mineral-organic associations. The dependence of priming effect on SOM stabilization mechanisms should be considered in Earth System Models to accurately predict soil C dynamics under changing environments.
Soil organic matter (SOM) plays a critical role for soil fertility, food production, climate regulation and ecosystem stability. As the largest organic carbon (C) pool in terrestrial ecosystems, the soil C stock is determined by the balance between C inputs from plants and C outputs through microbial decomposition and erosion, with these processes occurring interactively in nature. Of the various plant–soil interactions, the priming effect (i.e. alteration of SOM decomposition by labile C inputs) is a key mechanism affecting the soil C cycle. Incorporating the priming effect into Earth System Models can improve the prediction of soil C stocks. However, the large variability in the priming effect among various ecosystems ranging from a 380% increase to a 50% reduction in carbon dioxide (CO2) flux from soil greatly impedes the priming representation in Earth System Models. Knowledge concerning the dominant priming driver is thus crucial for model developments to accurately predict soil C dynamics and the terrestrial feedback to climate warming.

During the past several decades, the regulation of the priming effect by plant roots (plant C input by rhizodeposition), soil (pH and clay content) and microbial properties (microbial biomass, composition and activity) has been studied intensively. Despite all the research conducted thus far, our understanding of the dominant priming driver over broad geographic scales is still inadequate since the current studies have been mainly conducted at the site level. Encouragingly, the research community has now begun to address this issue by conducting either regional-scale measurements or global-scale syntheses. Specifically, recent experimental evidence has revealed the dependence of the priming on soil nitrogen content and microbial properties at the regional scale. Likewise, the synthesis of site-level observations has also showed associations of the priming effect with the quantity and quality of plant C input, soil texture and microbial biomass over broad geographic scales. These studies provide a basic understanding of the potential priming drivers over large scales. However, the variables that have been considered only explained 7.7–22.8% of the variations in the priming intensity, with substantial variance left unexplained. The large model residuals highlight the inadequate consideration of the critical drivers in current large-scale studies.

Recently, increasing evidence has revealed the vital role of SOM stabilization mechanisms (i.e. chemical recalcitrance and physico-chemical protection) in determining soil C turnover. These mechanisms may also mediate the priming effect through the alteration of microbial C demand and the accessibility of soil C sources. However, the role of these SOM stabilization mechanisms (hereafter called SOM stability) in regulating the priming effect as well as their relative importance compared with other factors (i.e. plant, soil and microbial properties) have not been identified over broad geographic scales. Specifically, previous studies have usually characterized SOM recalcitrance indirectly by using the C : N ratio or the dissolved organic C content, and have mainly focused on the dependency of the priming effect on the clay content, or aluminum (Al) and iron (Fe) oxides. In fact, the SOM molecular composition (e.g. carbohydrate, lipid and lignin contents) can represent chemical recalcitrance more directly, and the physico-chemical protection mediated by calcium (Ca2+) and soil aggregates may outweigh the role of Al/Fe oxides in neutral and alkaline soils. Due to the interactive effects of SOM stability, plant, soil and microbial properties on SOM decomposition, a comprehensive study with systematic measurements of the priming effect together with these potential drivers over a broad geographic scale is highly needed.

In this study, we quantify the relative importance of plant (vegetation productivity and composition), soil (nutrient content, pH, texture and bulk density) and microbial properties (total biomass, composition, structure, enzyme activities and stoichiometry), and SOM stability (chemical recalcitrance and physico-chemical protection) in regulating the priming effect based on 30 sites along an ~2200 km grassland transect on the Tibetan Plateau (Supplementary Fig. 1). These sites cover a wide precipitation gradient, from 89 mm in arid climates up to 534 mm in humid climates. Across this precipitation gradient, net primary productivity (NPP) varies between 38 and 488 g m−2 yr−1. Both soil and microbial properties are also highly variable, with soil organic C (SOC) and microbial biomass C ranging from 1.1 to 118 kg C ha−1 and 23 to 1101 mg C kg−1, respectively (Supplementary Data 1). Based on the large-scale sampling along these broad environmental gradients, we determine the priming effect induced by 13C-labelled glucose during a 65-day laboratory incubation, and then combine multiple analytical approaches (acid hydrolysis, biomarker analysis, a two-pool C decomposition model, aggregate fractionation and mineral analysis) to quantify the role of SOM stability in regulating the priming intensity. We further conduct three types of statistical analysis (i.e. partial correlation, variation partitioning analysis and structural equation modelling) to identify the relative importance of the various factors in shaping the regional patterns of the priming effect.

We find that SOM stability characterized by chemical recalcitrance and physico-chemical protection exerts a more important role than other factors over a broad geographic scale.

Results
Controls of SOM stability over the priming effect. Both chemical recalcitrance (i.e. the proportion of the labile and recalcitrant SOM factions) and physico-chemical protection (i.e. the proportion of C protected by aggregates, Fe and Al oxides and exchangeable calcium) were determined to characterize the SOM stability, and then used to explore their effects on priming intensity. Our results showed that the priming intensity increased with the chemical recalcitrance of SOM (Fig. 1). Specifically, both the priming intensity and the relative priming effect (as percentage of basal respiration) were negatively correlated with the proportions of labile pool I (composed of non-cellulosic polysaccharides) (all \( p < 0.01 \), Fig. 1a; Supplementary Fig. 2a), labile pool II (consisting of cellulose) (all \( p < 0.05 \), Fig. 1b; Supplementary Fig. 2b) and the labile carbohydrate content (all \( p < 0.05 \), Fig. 1d; Supplementary Fig. 2d). Conversely, increases in the recalcitrant pool (mainly composed of polymers of lipid and lignin) raised the priming intensity (all \( p < 0.01 \), Fig. 1c; Supplementary Fig. 2c). Likewise, most of the recalcitrant SOM component (e.g. cutin- and suberin-derived compounds) exerted positive effects on the priming intensity (all \( p < 0.01 \), Fig. 1e–f; Supplementary Fig. 2e–f), although no correlation was found between the priming intensity and lignin-derived phenols (Supplementary Fig. 3a). These results from acid hydrolysis and biomarker analysis were supported by the negative correlation of priming intensity with the proportion of fast C pool estimated by the two-pool C decomposition model (\( p < 0.05 \), Supplementary Fig. 3b).

Our results also revealed negative relationships between the priming effect and soil physico-chemical protection. Specifically, priming intensities were low in soils with high molar ratios of Fe/Al oxides or exchangeable Ca to SOC (all \( p < 0.01 \), Fig. 1g–i). Moreover, the priming intensity was positively correlated with the proportion of C occluded in macroaggregates (C >250 μm) (\( p < 0.01 \), Fig. 1j), but negatively related to the proportion of C occluded in microaggregates (53–250 μm) (\( p < 0.01 \), Fig. 1k) and clay + silt fractions (<53 μm) (\( p < 0.05 \), Fig. 1l). Notably, the patterns obtained for the relative priming effect were the same as those corresponding to the priming intensity (Supplementary Fig. 2g–l). Taken together, these results demonstrated that regional variations in priming intensity were jointly controlled by SOM chemical recalcitrance and physico-chemical protection.

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Importance of SOM stability relative to other properties. Three statistical methods, including partial correlation analysis, variation partitioning analysis and structural equation modelling, were used to discern the relative importance of SOM stability (i.e. chemical recalcitrance and physico-chemical protection) compared with other factors. Partial correlation analysis showed that, without controlling the role of SOM stability (zero-order in Fig. 2), the priming intensity was closely correlated with plant, soil and microbial properties. However, after controlling SOM stability, the correlation coefficients between the priming intensity and plant, soil and microbial properties decreased by 33.0%, 80.1% and 97.0%, respectively. As a consequence, except for the coverage of sedge and forb, whose correlation with priming intensity retained or increased, most of these factors were no longer associated with priming effect (Fig. 2). In contrast, the correlation coefficients only decreased by 19.6%, 19.4% and 6.0% between the priming intensity and SOM stability after controlling plant, soil and microbial properties, respectively (Supplementary Fig. 4). In particular, the abundance of cutin-derived compounds was always significantly correlated with the priming intensity, despite its correlation with the priming intensity decreased by 42.9% when the glucose input amount was controlled (Supplementary Fig. 4). Similar results were also found for the relative priming effect (Supplementary Figs. 5–6). These results highlighted that SOM stability was the dominant driver of priming intensity over a continental scale.

Fig. 1 Relationships of the priming effect with soil organic matter chemical recalcitrance and physico-chemical protection. Soil organic matter chemical recalcitrance is characterized by the proportion of labile pool I (a), labile pool II (b), the recalcitrant pool (c), relative abundances of carbohydrates (d), cutin-derived compounds (e) and suberin-derived compounds (f). Physico-chemical protection is reflected by the molar ratios of free Fe/Al oxides (g) and amorphous Fe/Al oxides (h) to SOC, the ratio of exchangeable Ca to SOC (i), and the SOC proportions in macroaggregates (j), microaggregates (k), and clay + silt fractions (l). The solid lines represent the fitted ordinary least-squares model, and the grey areas correspond to 95% confidence intervals. A base-10 log scale is used for the x-axis of b, d and g–i. The orange dots and green squares represent data in alpine steppe and alpine meadow, respectively. **Significant correlation between the priming effect and the corresponding variables at the 0.01 level.
priming intensity than plant (6.7%), soil (8.1%) and microbial properties (4.0%) (Fig. 3a). Similarly, this predominant contribution of SOM stability was observed for the relative priming effect (Fig. 3b). Structural equation modelling analysis further revealed that SOM stability and microbial properties were two direct factors that regulated the priming effect. The combination of plant, soil and microbial properties as well as SOM stability explained 54% and 43% of the variance in the priming intensity and relative priming effect, respectively (Fig. 4). Of the factors tested in this study, SOM stability had the largest direct effect among which the correlations with the priming effect are examined. The size and colour of the circles indicate the strength and sign of the correlation. Differences in circle size and colour between the zero-order and controlled factors indicate the level of dependency of the correlation between the priming effect and the examined factor on the controlled variable (no change in circle size and colour between the controlled factor and zero-order = no dependency; a decrease/increase in circle size and colour intensity = loss/gain of correlation). ANPP, aboveground net primary productivity; EVI, enhanced vegetation index; CInp, C input amount; Gra, relative coverage of grass; Sed, relative coverage of sedge; Forb, relative coverage of forb; SOC, soil organic carbon content; BD, bulk density; PLFA: total PLFAs; Fun, fungal PLFAs; Bat, bacterial PLFAs; F/B, fungi/bacteria ratio; Cenz, C-acquiring enzyme activity; Nenz, N-acquiring enzyme activity; Penz, P-acquiring enzyme activity; CNenz, C:N ratio of enzyme activity; NPenz, N:P ratio of enzyme activity; CPenz, C:P ratio of enzyme activity; LPI, labile pool I; LPII, labile pool II; RP, recalcitrant pool; Carb, carbohydrate; Cut, cutin-derived compound; Sub, suberin-derived compound; FeAlₓ, molar ratio of free Fe/Al oxides to SOC; FeAlₓ, molar ratio of amorphous Fe/Al oxides to SOC; Ca, ratio of exchangeable Ca to C; Mac, proportion of C protected by macroaggregates; Mic, proportion of C protected by microaggregates; C + S, proportion of C associated with clay + silt fractions. *p < 0.05; **p < 0.01

Discussion

Based on multiple analytical and statistical approaches, this study provided empirical evidence that SOM stability, characterized by chemical recalcitrance and physico-chemical protection, had higher predictive power than other factors in explaining regional-scale patterns of the priming effect. The priming intensity increased with the proportion of recalcitrant pool and the content of recalcitrant components (e.g. cutin and suberin), but decreased with physico-chemical protection by minerals and aggregates (Fig. 5). The regulation of SOM chemical recalcitrance on the priming effect could be attributed to its potential impacts on microbial C and N requirements. On the one hand, compared with soils rich in labile C fractions (e.g. carbohydrates), soils with more recalcitrant polymers usually provide less bioenergetically favourable substrates to microorganisms, leading to the relatively higher microbial C limitation and a larger proportion of microbial dormancy. Consequently, more microorganisms could be activated by labile C inputs to these soils, further stimulating SOM decomposition. On the other hand, high SOM recalcitrance can also trigger great microbial N demand, since the complex molecular structure can hinder the breakdown of N-containing polymers (e.g. chitin) to access available N. The high microbial N requirement could then contribute to CO₂ release from these soils by either stimulating SOM decomposition to acquire N (microbial N mining theory) or accelerating microbial N recycling at the expense of C use efficiency.

The decrease in the priming intensity with physico-chemical protection by minerals and aggregates could be due to its constraint on SOM availability. Specifically, Fe/Al (hydro)oxides have negatively charged functional groups that enable them to participate in ligand exchange reactions and effectively adsorb...
SOM protecting them against biodegradation\(^{30}\). In addition to Fe/Al oxides, Ca\(^{2+}\) can also constrain SOM availability by forming both inner- and outer-sphere cation bridging\(^{22}\), with its protective effect outweighing Fe/Al oxides in neutral to alkaline soils\(^{22}\). Additionally, these oxides and cations can accelerate the formation of stable aggregates by their cementation of other minerals\(^{29,30}\), which results in the compartmentalization of substrates from enzymes and restricts the diffusion of oxygen, thus decreasing SOM decomposition. Consistent with this deduction, the proportion of C associated with clay + silt fractions increased with the molar ratios of metal to SOC across our study area (Supplementary Fig. 8). Taken together, mineral and aggregate protection could inhibit the priming effect through their constraints on organic C availability.

Although both chemical recalcitrance and physico-chemical protection of SOM could regulate the priming effect over a broad geographic scale, the effect of chemical composition depends largely on the degree of physico-chemical protection\(^{16,31–33}\). When a relatively larger proportion of SOM is protected by mineral–organic associations and aggregate occlusion, SOM stability could be primarily governed by the degree of protection rather than chemical recalcitrance\(^{32}\). Nevertheless, chemical recalcitrance still regulates microbial decomposition when SOM is weakly stabilized by physico-chemical interactions\(^{34}\). This phenomenon can be observed with dissolved organic C\(^{35}\), free light-fraction SOM\(^{36}\) and, to a certain extent, organic components with fewer associations with minerals\(^{36,34}\). Given that only 15.8% of the soil C pool was protected by minerals across our study area\(^{37}\), both SOM decomposition and the associated priming effect were jointly affected by the chemical composition and mineral protection.

In addition to SOM stability, plant properties (i.e. plant productivity and community composition) were also important in regulating the priming effect, and exerted strong indirect impacts through their associations with SOM stability (Fig. 4). Particularly, all the variables of SOM stability were significantly correlated with 75% of the plant properties (e.g. aboveground net primary production (ANPP), enhanced vegetation index (EVI), C input rate and the coverage of grass) (Supplementary Fig. 9). The strong dependence of SOM stability on plant properties could be ascribed to the following aspects. On the one hand, as the main SOM source, the rate of plant C input directly determines the amount of plant-derived C, which could be newly stabilized in soil\(^{32}\) and affects the decomposition of stable SOM pool via the priming effect\(^{3}\). Plant C input can also mediate geochemical processes (e.g. soil acidification), which affect soil pH and the amount of cations\(^{38}\), thus indirectly regulating the SOM stability. On the other hand, plant community composition controls the chemistry of plant litter and root exudates\(^{39}\), thereby directly affecting the chemical recalcitrance of SOM. The organic acids (e.g. oxalic acid) from root exudates can also destabilize SOM by liberating C, which are associated with clay minerals, and thus increase C accessibility\(^{18}\). Overall, plant properties can regulate the priming effect through their modification on SOM stability.

Although our study demonstrated the important role of chemical recalcitrance and physico-chemical protection in regulating the priming effect over a broad geographic scale, there are still some limitations that need to be addressed in future studies. Particularly, despite being a frequently used approach in priming experiments, a single pulse of glucose addition with an amount equal to the microbial biomass cannot realistically characterize plant C input in terms of quantity, quality\(^{3,40}\) and input frequency in natural ecosystems. This single addition may also induce changes in microbial biomass\(^{9}\), community structure\(^{41}\) and microbial C use efficiency\(^{42}\). Furthermore, due to the decreased microbial diversity\(^{43}\) and enhanced microbial C starvation during the late stage of incubation\(^{44}\), this single addition may result in a lower priming effect or even negative priming compared with repeated substrate additions\(^{45}\). In situ experiments are thus encouraged to better capture realistic plant C input and elucidate the role of plant properties in regulating the priming effect over a large scale. In addition, the limited ecosystem types and climate zones involved in this study may have also induced uncertainties when generalizing patterns and drivers observed in this study. More empirical studies with diverse ecosystem types (forests, shrubs, etc.) and climate zones (tropical, temperate zone, etc.) are thus needed to further advance our understanding on this issue.

In summary, based on systematic measurements of the priming effect together with plant, soil and microbial properties and SOM stability, this study provides empirical evidence that SOM stability, as regulated by plant productivity and community composition, determines the glucose-induced priming effect over a large scale. The stronger predictability of SOM stability than other factors implies that the priming intensity is also governed by SOM stabilization mechanisms, which is similar to what has been observed for soil C pools\(^{46}\) and their turnover\(^{47}\). The higher predictive power of SOM stability also infers potential uncertainties in priming simulations among current models, in which priming intensity is solely assumed to depend on the amount of plant C input\(^{36}\). Future modelling studies should thus consider both SOM chemical recalcitrance and physico-chemical
The Tibetan Plateau is the largest and highest plateau in the world and has broad environmental gradients, as such, this area serves as an ideal platform for exploring the dominant drivers of the priming effect over a broad geographic scale. Specifically, the climate is characterized as cold and dry across the plateau, with a southeast to northwest precipitation gradient ranging from 84 to 593 mm per year. The mean annual temperature in this area varies from −4.9 to 6.9 °C. The dominant ecosystem on the plateau is alpine grassland, which shifts from alpine steppe in the northwestern area to alpine meadow in the southeastern area. Of the two major grassland types, the alpine steppe is mainly dominated by *Stipa purpurea* and *Carex moorcroftii* and is characterized by low precipitation amounts, low plant productivity and a low soil C content. In contrast, the alpine meadow, dominated by *Kobresia pygmaea*, *K. humilis* and *K. tibetica*, receives relatively high amounts of precipitation and corresponds to high plant productivity and higher soil C content.

**Methods**

**Study area and field sampling.** The Tibetan Plateau is the largest and highest plateau in the world and has broad environmental gradients, as such, this area serves as an ideal platform for exploring the dominant drivers of the priming effect over a broad geographic scale. Specifically, the climate is characterized as cold and dry across the plateau, with a southeast to northwest precipitation gradient ranging from 84 to 593 mm per year. The mean annual temperature in this area varies from −4.9 to 6.9 °C. The dominant ecosystem on the plateau is alpine grassland, which shifts from alpine steppe in the northwestern area to alpine meadow in the southeastern area. Of the two major grassland types, the alpine steppe is mainly dominated by *Stipa purpurea* and *Carex moorcroftii* and is characterized by low precipitation amounts, low plant productivity and a low soil C content. In contrast, the alpine meadow, dominated by *Kobresia pygmaea*, *K. humilis* and *K. tibetica*, receives relatively high amounts of precipitation and corresponds to high plant productivity and higher soil C content.
Fig. 5  Soil organic matter stability, as regulated by plant C input and community composition, determines the priming effect. With the increase of plant productivity and forb cover, more plant-derived recalcitrant SOM fractions (e.g. cutins and suberins) accumulated in the soil. The proportion of C occluded in macroaggregates increased at the expense of C associated with microaggregates and clay + silt fractions, and the physico-chemical protection by aggregations and minerals became weaker. Consequently, either (a) the increased chemical recalcitrance or (b) the reduced physico-chemical protection (increased C accessibility) could lead to the stronger priming intensity. Notably, this conceptual diagram mainly focused on the most important direct (i.e. soil organic matter stability) and indirect factors (i.e. plant properties) derived from the structural equation modelling analysis.
Particularly, the 13C-labelled CaCO3 experiment revealed that CaCO3-derived CO2 only accounted for 1.0% of the total CO2 release. These analyses demonstrated a minor contribution of carbonate to the CO2 production and also its δ13C. Therefore, as done in most priming studies on soil with pH > 7.0 [3,5], the effects of soil carbonates on CO2 release were considered limited in this study. The fractions of CO2-C derived from the added 13C-glucose (fglucose) and from the SOM pool (fSOM) were then determined by Eqs. (2) and (3).

\[ f_{\text{glucose}} = \frac{(a_{\text{treat}} - a_{\text{control}}) / a_{\text{glucose}}}{a_{\text{SOM}}} \]

\[ f_{\text{SOM}} = 1 - f_{\text{glucose}} \]

where \(a_{\text{SOM}}\) is the SD of the SOM-derived CO2 flux in the glucose treatment, \(a_{\text{glucose}}\) and \(a_{\text{SOM}}\) are the SDs of the total CO2 fluxes in the glucose-treated soil and the SOM pool fraction, respectively, and \(f_{\text{glucose}}\) and \(f_{\text{SOM}}\) are their respective mean values.

After obtaining the means and SDs of SOM-derived CO2 release under both the glucose treatment and control conditions, we performed 1000 Monte Carlo simulations for the priming effect during each time’s measurement. First, the SOM-derived CO2 release in both the glucose treatment and control conditions were randomly generated based on normal distributions using the above-mentioned means and SDs. The priming effect was then calculated as the difference in the cumulative priming effect over the incubation time. Finally, we calculated the 95% confidence interval of the cumulative priming effect based on a total of 1000 estimates derived from four groups to explore their roles in regulating the priming effect: plant, soil and microbial properties.

Identification of SOM stability. SOM stability was quantified by two types of stabilization mechanisms: SOM chemical recalcitrance and physico-chemical protection. For SOM chemical recalcitrance, we used the relative abundances of the labile and recalcitrant SOM pools measured by the acid hydrolysis method [5], the relative abundances of particular SOM components determined by biomarker analysis and the proportion of fast C pool inversed from the two-pool C decomposition model [6]. Specifically, the proportions of labile pool I, labile pool II and the fast C pool, and the content of carbohydrates were used to represent the labile SOM fractions, while the proportion of recalcitrant C pool and the relative abundances of lignin-, cutin- and suberin-derived compounds were used to reflect the recalcitrant SOM fractions. For physico-chemical protection, the proportion of C in macro-aggregates, microaggregates and clay + silt fractions was determined by soil C fractionation [7], and the ratios of metal to SOC (i.e. free and amorphous Fe and Al oxides: molar Fe2+Al/SOC, molar Fe2+Al/SOC; and exchangeable calcium: Caex/SOC) were applied [8].

Acid hydrolysis method. The soil labile C pool (composed of polysaccharides and cellulose) and the recalcitrant C pool (composed of wax-derived long-chain aliphatics and aromatic components) were determined by the acid hydrolysis method [9]. Briefly, soil samples were first hydrolysed with 2.5 M H2SO4 at 105 °C for 30 min. The hydrolysates were centrifuged and decanted. The residue was washed with distilled water and the supernatant was added to the hydrolysate, which was regarded as labile pool I. The remaining residue was further hydrolysed with 13 M H2SO4 and shaken overnight at room temperature. Subsequently, the distilled water was added to dilute the acid concentration to 1 M, and the sample was heated at 105 °C for 30 min, and then cooled to room temperature. The remaining soil residue was rinsed twice with distilled water and dried at 60 °C. This fraction was considered the recalcitrant SOM pool. The relative abundances of labile pool I and II and the recalcitrant pool were calculated as the ratio of SOC in each pool to the total SOC [10].

SOM biomarker analysis. The biomarker extraction and analysis followed the standard procedures, in which sequential chemical extractions (solvent extraction, base hydrolysis and CuO oxidation) were conducted to separate the solvent-extractable compounds, cutin- and suberin-derived compounds and lignin-derived phenols, respectively [11]. Of the solvent-extractable compounds, we focused on the most labile C fractions (carbohydrates). The detailed chemical extraction and analysis are provided in the Supplementary Methods.

Plant C decomposition model. The proportion of fast and slow C pool, another proxy for SOM chemical recalcitrance, were estimated by a two-pool model [12,13]. Specifically, we applied the two-pool model to each of the 30 sites in the control group as follows:

\[ R(t) = k_1f_1 e^{-k_1t} + k_2 (1-f_1) e^{-k_2t} \]

where \(R(t)\) is the CO2 emission rate (mg C g−1 SOC day−1) at time t, \(k_1\) is the maximum C loss percentage (i.e. 100% = 1000 mg C g−1 180C, \(f_1\) is the proportion of the fast pool, and \(k_1\) and \(k_2\) are the decay rates of the fast and slow pools (day−1), respectively. The three parameters \((f_1, k_1, k_2)\) were estimated by the Markov chain Monte Carlo (MCMC) approach [14]. Before applying the MCMC approach, the prior parameter range in the initial model (Supplementary Table 2) was set as wide as possible to cover two possibilities for all sites. After the MCMC simulations, maximum likelihood estimates were used to quantify the well-constrained parameters, while mean values were calculated for the poorly

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constrained parameters (Supplementary Table 3). The data-model comparison revealed good model performance for the 30 study sites ($R^2 = 0.95; \text{RMSE} = 0.22$). Notably, a two-pool model rather than a three-pool model was chosen because the relatively short-term (65 days) incubation made it less possible to estimate the true turnover time of the passive C pool in the three-pool model.

**Physical protection parameters.** The proportion of C in the different functional SOM fractions was used to characterize the role of physical protection by agglomerates. The functional C fractions were obtained following the method based on the conceptual C fraction model. Briefly, soil C was fractionated into macroaggregate-associated C (>250 μm), microaggregate-associated C (250–53 μm) and non-aggregated silt + clay C (<53 μm). The isolated fractions were analysed for their SOC contents using an elemental analyser (mstil EA 4000) after removing the inorganic C with 1 M HCl. Based on these C fractions and the SOC contents in the bulk soil, we calculated the proportion of C associated with the different fractions.

**Mineral protection parameters.** The contents of Fe and Al oxides and exchangeable Ca and exchangeable Al were used to calculate the mineral protection parameters. The reactive Fe and Al contents were determined by the citrate-bicarbonate-dithionite method ($Fe_C + Al_C$) and acid oxalate extraction ($Fe_N + Al_N$). The $Fe_C + Al_C$ represents the amount of pedogenic Fe and Al within oxides, silicates and organic complexes, whereas $Fe_N + Al_N$ originates from poorly crystalline mineral complexes. The exchangeable C was extracted by NH₄Cl-ethanol. Based on these measurements and the SOC content in bulk soil, the molar ratio of metals to SOC ($molar_\text{Fe}/\text{SOC}$ and $molar_\text{Al}/\text{SOC}$) and the ratio of exchangeable Ca to SOC ($Ca_{\text{ex}}/\text{SOC}$) were calculated to represent the potential of minerals to adsorb SOC.

**Statistical analyses.** We first conducted ordinary least squares regression to evaluate the relationships between the priming effect and SOM stability. We then used three types of statistical models (partial correlation, variation partitioning analysis and structural equation modelling) to test whether the importance of SOM stability was maintained when accounting for other three types of factors. Given the strong connections and inter-correlations among the various factors (Supplementary Fig. 9), partial correlation was first conducted to evaluate the relationships between the priming effect and the various factors. For example, after controlling every single variable of SOM stability, we examined the relationships of the priming effect with plant, soil and microbial properties. Similarly, plant, soil and microbial properties were separately set as the controlling factors to explore the relationship between the priming effect and SOM stability. The stronger the difference in the partial correlation coefficient between the zero-order and controlling correlation, the stronger the effect of the factor being controlled. Moreover, the relatively greater difference also indicated the stronger interaction effects among these predictors. These analyses were conducted using the packages ggm and psych of the R statistical software v.3.2.4 (R Development Core Team, 2016). Variation partitioning analysis that partitioned the variance shared by all factors was then used to quantify the unique contribution of each group of factors. A negative value in the variance explained for a group of factors was interpreted as zero, which indicated that the explanatory variables explained less variation than random normal variables. The variation partitioning analyses were conducted with the R package vegan v.3.2.4 (R Development Core Team, 2016).

Structural equation modelling was further used to evaluate the direct and indirect relationships between the priming effect and SOM stability, plant, soil and microbial properties. This approach can partition the direct and indirect effects that one variable may have on another and is therefore useful for exploring complex relationships in natural ecosystems. Owing to strong correlations among the factors within each group, we conducted principal component (PC) analysis to create a multivariate functional index before structural equation modelling construction. The first component (PC1), which explained 49.7–66.0% of the total variance for these four groups, was then introduced as a new variable to represent the combined group properties into the subsequent analysis (Supplementary Table 4). The fit of the final model was evaluated using the model $R^2$ test and the root mean-squared error of approximation. The structural equation modelling analyses were conducted using AMOS 21.0 (Amos Development Corporation, Chicago, IL, USA).

**Data availability**

All plant, soil and microbial predictors used in this study are available as a supplementary file (Supplementary Data 1). Additional data that support the findings of this study are available from Y.Y. upon reasonable request.

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**References**

1. Jackson, R. B. et al. The ecology of soil carbon: pools, vulnerabilities, and biotic and abiotic controls. *Annu. Rev. Ecol. Evol. Syst.* 48, 419–445 (2017).
2. Battin, T. J. et al. The boundless carbon cycle. *Nat. Geosci.* 2, 598–600 (2009).
3. Kuzyakov, Y. Priming effects: interactions between living and dead organic matter. *Soil Biol. Biochem.* 42, 1363–1371 (2010).
4. Fontaine, S. et al. Stability of organic carbon in deep soil layers controlled by fresh carbon supply. *Nature* 531, 279–282 (2016).
5. Guenet, B. et al. Impact of priming on global soil carbon stocks. *Glob. Change Biol.* 24, 1873–1883 (2018).
6. Sulman, B. N., Phillips, R. P., Oishi, A. C., Shevlakova, E. & Pacala, S. W. Microbe-driven turnover offsets mineral-mediated storage of soil carbon under elevated CO₂. *Nat. Clim. Change* 4, 1099–1102 (2014).
7. Xiao, C., Luo, Y. & Chen, X. Rhizosphere priming effect: a meta-analysis. *Soil Biol. Biochem.* 111, 78–84 (2017).
8. Qiao, N. et al. Carbon and nitrogen additions induce distinct priming effects along an organic-matter decay continuum. *Sci. Rep.* 6, https://doi.org/10.1038/srep19865 (2016).
9. Chen, R. et al. Soil C and N availability determine the priming effect: microbial N mining and stoichiometric decomposition theories. *Glob. Change Biol.* 20, 2356–2367 (2014).
10. Xiao, C., Guenet, B., Zhou, Y., Su, J. & Janssens, I. A. Priming of soil organic matter decomposition scales linearly with microbial biomass response to litter input in steppe vegetation. *Oikos* 124, 649–657 (2015).
11. Atkinson, V. N. et al. Understanding of priming effect: an analysis using thirty five soils with contrasted properties sampled from five continents. *Soil Biol. Biochem.* 134, 162–171 (2019).
12. Razanampalala, K. et al. Soil microbial diversity drives the priming effect along climate gradients: a case study in Madagascar. *ISME J.* 12, 451–462 (2017).
13. Luo, Z., Wang, E. & Smith, C. Fresh carbon input differentially impacts soil carbon decomposition across natural and managed systems. *Ecology* 96, 2806–2813 (2015).
14. Luo, Z., Wang, E. & Sun, O. J. A meta-analysis of the temporal dynamics of priming soil carbon decomposition by fresh carbon inputs across ecosystems. *Soil Biol. Biochem.* 101, 96–103 (2016).
15. Lehmann, J. & Kleber, M. The contentious nature of soil organic matter. *Nature* 528, 60–68 (2015).
16. Schmidt, M. W. et al. Persistence of soil organic matter as an ecosystem property. *Nature* 478, 49–56 (2011).
17. Hamer, U. & Marschner, B. Priming effects in different soil types induced by fructose, alanine, oxalic acid and catechol additions. *Soil Biol. Biochem.* 37, 445–454 (2005).
18. Keiluweit, M. et al. Mineral protection of soil carbon counteracted by root exudates. *Nat. Clim. Change* 5, 588–595 (2015).
19. Chowdhury, S., Farrell, M. & Bolan, N. Priming of soil organic carbon by malic acid addition is differentially affected by nutrient availability. *Soil Biol. Biochem.* 77, 158–169 (2014).
20. Finley, B. K. et al. Soil mineral assemblage and substrate quality effects on microbial priming. *Geoderma* 322, 38–47 (2018).
21. Chen, L. Y. et al. Determinants of carbon release from the active layer and permafrost deposits on the Tibetan Plateau. *Nat. Commun.* 7, https://doi.org/10.1038/ncomms13046 (2016).
22. Rowley, M. C., Grand, S. & Verrecchia, É. P. Calcium-mediated stabilisation of soil organic carbon. *Biogeochemistry* 137, 27–49 (2018).
23. Rovira, P. & Valloge, V. R. Examination of thermal and acid hydrolysis procedures in characterization of soil organic matter. *Commun. Soil Sci. Plant Anal.* 31, 81–100 (2000).
24. Rovira, P. & Valloge, V. R. Labile and recalcitrant pools of carbon and nitrogen in organic matter decomposing at different depths in soil: an acid hydrolysis approach. *Geoderma* 107, 109–141 (2002).
25. Blagodatskaya, E. & Kuzyakov, Y. Active microorganisms in soil: critical review of estimation criteria and approaches. *Soil Biol. Biochem.* 67, 192–211 (2013).
26. Treat, C. C. et al. Temperature and peat type control CO₂ and CH₄ production in Alaskan permafrost peats. *Glob. Change Biol.* 20, 2674–2686 (2014).
27. Bagley, J. A., Chapman, S. K. & Hungate, B. A. Ectomycorrhizal colonization slows root decomposition: the post-mortem fungal legacy. *Ecol. Lett.* 9, 955–959 (2006).
28. Chen, L. et al. Nitrogen availability regulates topsoil carbon dynamics after permafrost thaw by altering microbial metabolic efficiency. *Nat. Commun.* 9, https://doi.org/10.1038/s41467-018-06232-y (2018).
29. Singh, M. et al. Stabilization of soil organic carbon is influenced by clay mineralogy. *Adv. Agron.* 148, 33–84 (2018).
30. Keiluweit, M. et al. Nano-scale investigation of the association of microbial nitrogen residues with iron (hydr)oxides in a forest soil O-horizon. *Geochim. Cosmochim. Acta* 95, 213–226 (2012).
31. Buttera, R., Kleber, M., Torn, M. S. & Jahn, R. Stabilization of soil organic matter: association with minerals or chemical recalcitrance? *Biogeochemistry* 77, 25–56 (2006).
32. Dungait, J. A. J., Hopkins, D. W., Gregory, A. S. & Whitmore, A. P. Soil organic matter turnover is governed by accessibility not recalcitrance. *Glob. Change Biol.* **18**, 1781–1796 (2012).

33. Kleber, M. What is recalcitrant soil organic matter? *Environ. Chem.* **7**, 320–332 (2010).

34. Sjögersten, S. et al. Organic matter chemistry controls greenhouse gas emissions from permafrost peatlands. *Soil Biol. Biochem.* **98**, 42–53 (2016).

35. Marschner, H. & Kalbitz, K. Controls of bioavailability and biodegradability of dissolved organic matter in soils. *Geoderma* **113**, 211–235 (2003).

36. Wagai, R. et al. Linking temperature sensitivity of soil organic matter decomposition to its molecular structure, accessibility, and microbial physiology. *Glob. Change Biol.* **19**, 1114–1125 (2013).

37. Fang, K., Qin, S., Chen, L., Zhang, Q. & Yang, Y. Al/Fe mineral controls on soil organic carbon stock across Tibetan alpine grasslands. *J. Geophys. Res. Biogeosci.* **124**, 247–259 (2019).

38. Angst, G. et al. Soil organic carbon stability in forests: distinct effects of tree species identity and traits. *Glob. Biol. Change* **25**, 1529–1546 (2018).

39. Hodgkin, S. B. et al. Changes in peat chemistry associated with permafrost thaw increase greenhouse gas production. *Proc. Natl Acad. Sci. USA* **111**, 5819–5824 (2014).

40. Cheng, W. X. et al. Synthesis and modeling perspectives of rhizosphere priming. *New Phytol.* **201**, 31–44 (2014).

41. Fontaine, S., Mariotti, A. & Abbadi, L. The priming effect of organic matter: a question of microbial competition? *Soil Biol. Biochem.* **35**, 837–843 (2003).

42. Schuur, E. A. G. et al. Ecosystem carbon vulnerability to climate change and soil warming. *Proc. Natl Acad. Sci. USA* **105**, 1237–1243 (2008).

43. Grassi, R. et al. Microbial priming decomposes easily available organic C and N stimulates microbial decomposition of soil organic matter in arctic permafrost soil. *Soil Biol. Biochem.* **75**, 143–151 (2014).

44. Mau, R. L. et al. Linking soil bacterial biodiversity and soil carbon stability. *ISME J.* **9**, 1477–1480 (2015).

45. Qiao, N. et al. Labile carbon retention compensates for CO$_2$ released by priming in forest soils. *Glob. Change Biol.* **20**, 1943–1954 (2014).

46. Hamer, U. & Marschner, B. Priming effects in soils after combined and repeated substrate additions. *Geoderma* **128**, 38–51 (2005).

47. Doetterl, S. et al. Soil carbon storage controlled by interactions between geochemistry and climate. *Nat. Geosci.* **8**, 780–783 (2015).

48. Herold, N., Schöning, I., Michalzik, B., Trumbore, S. & Scharpf, M. Controls on soil carbon storage and turnover in German landscapes. *Biogeochemistry* **119**, 435–451 (2014).

49. Yang, Y. H., Fang, J. Y., Pan, Y. D. & Ji, C. J. Aboveground biomass in Tibetan grasslands. *J. Arid Environ.* **73**, 91–95 (2009).

50. Ding, J. Z. et al. The permafrost carbon inventory on the Tibetan Plateau: a new evaluation using deep sediment cores. *Glob. Change Biol.* **22**, 2688–2701 (2016).

51. Shi, X. Z. et al. Soil database of 1,000,000,000 soil survey and reference system of the Chinese genetic soil classification system. *Soil Survey Horizons* **45**, 129–136 (2004).

52. Yang, Y. et al. Storage, patterns and controls of soil organic carbon in the Tibetan alpine grassland. *Glob. Change Biol.* **14**, 1592–1599 (2008).

53. Chen, Y. L. et al. Patterns and drivers of soil microbial communities in Tibetan alpine and global terrestrial ecosystems. *J. Biogeogr.* **43**, 2027–2039 (2016).

54. Fierer, N., Colman, B. P., Schimel, J. P. & Jackson R. B. Predicting the temperature dependence of microbial respiration in soil: a continental-scale analysis. *Glob. Biogeochem. Cycle* **20**, https://doi.org/10.1029/2005GB002644 (2006).

55. Chapin, F. S., Matson, P. A. & Vitousek, P. M. Principles of Terrestrial Ecosystem Ecology 2nd edn (Springer, 2012).

56. Bastida, F. et al. Global ecological predictors of the soil priming effect. *Nat. Commun.* **10**, 3481 (2019).

57. Zhu, B. & Cheng, W. Constant and diurnally-varying temperature regimes lead to different temperature sensitivities of soil organic carbon decomposition. *Soil Biol. Biochem.* **43**, 866–869 (2011).

58. Yang, Y., Fang, J., J. C. & Han, W. Above- and belowground biomass allocation in Tibetan grasslands. *J. Veg. Sci.* **20**, 177–184 (2009).

59. Nelson, D. W. & Sommers, L. E. Total carbon, organic carbon, and organic matter. in *Methods of Soil Analysis Part II* (American Society of Agronomy, Madison, 1982).

60. Bastida, F. et al. Can the labile carbon contribute to carbon immobilization in semiarid soils? Priming effects and microbial community dynamics. *Soil Biol. Biochem.* **57**, 892–902 (2013).

61. Bossio, D. A. & Sow, K. M. Impacts of carbon and flooding on soil microbial communities: phospholipid fatty acid profiles and substrate utilization patterns. *Micro. Ecol.* **35**, 265–278 (1998).

62. German, D. P. et al. Optimization of hydrolytic and oxidative enzyme methods for ecosystem studies. *Soil Biol. Biochem.* **43**, 1387–1397 (2011).

63. Liang, J. et al. Methods for estimating temperature sensitivity of soil organic matter based on incubation data: a comparative evaluation. *Soil Biol. Biochem.* **80**, 127–135 (2015).