INTRODUCTION
The formation and mineralization of soil organic carbon (SOC) are crucial for climate regulation, food provisioning, cultural heritage, habitat for organisms, and nutrient cycling (1, 2), but they remain poorly understood and quantified, particularly in different soil fractions. Separating SOC into particulate organic carbon (POC) and mineral-associated organic carbon (MAOC) is fundamental to understand the underlying processes that explain SOC formation and mineralization (3, 4). Previous studies have shown that MAOC has a relatively long persistence in the soil but requires large quantities of nitrogen for its formation and has limited storage capacity. In contrast, the POC fraction is more vulnerable to disturbance but has a lower nitrogen demand and can potentially accumulate indefinitely (3). Chemical recalcitrance of soil organic molecules and their location within large aggregates constrain POC mineralization, whereas in the MAOC fraction mineralization is constrained by physical-chemical protection, resulting from carbon (C) occlusion within microaggregates and adsorption onto mineral surfaces (5, 6). Previous research proposes that SOC formation is mainly explained by physical-chemical protection (Fig. 1A). On the other hand, “living plant experiments” consider potential protection mechanisms of SOC promoted by living roots (7). By analyzing and comparing different experimental approaches, we estimated SOC_FEs, POC_FEs, and MAOC_FEs of aboveground inputs, roots, and rhizodeposition separately using data from 35 studies and 197 observations that applied 13C as a tracer of plant inputs into the soil (Table 1). On the one hand, we compiled “litter incubation experiments” wherein the POC and aboveground inputs were added to the soil and SOC derived from each input was quantified through time (Fig. 1A). Our database includes litter incubation experiments that added litter and roots from crop and grass species, as well as forest leaves and fine roots, which performed either in controlled laboratory conditions or in the field (see Table 1 for details; 61% of the litter incubation experiments were performed in the field). Such experiments have traditionally evaluated SOC_FEs, although they do not include C-inputs from rhizodeposition and tend to ignore potential protection mechanisms of SOC promoted by living roots (Fig. 1A). On the other hand, “living plant experiments” consider the interactions of living plants with the soil matrix when estimating SOC_FEs. Our review includes living plant experiments that evaluated crop and grass species growing not only mainly in the field but also under controlled laboratory conditions (see Table 1 for details). In these experiments, rhizodeposition occurs throughout the growing season, whereas root and shoot biomass are incorporated into the soil primarily as a pulse after plant harvest (Fig. 1B). Because these different experimental approaches have important effects on SOC_FEs estimates, we combined them to estimate the SOC_FEs.
Table 1. Summary of the reviewed articles. RN, reference number; Ag, aboveground plant residues; NR, net rhizodeposition was estimated; NR*, net rhizodeposition was measured; MAP, mean annual precipitation; MAT, mean annual temperature; NA, not applicable.

| RN | C-input source | Experiment type | SOC fractions* | Method for new-C estimation | Environment condition | Plant species | Soil texture | MAP (mm) | MAT (°C) | Residue management | Time (years) |
|----|----------------|-----------------|----------------|----------------------------|----------------------|---------------|--------------|----------|----------|--------------------|--------------|
| (39) | Roots | Litter incubation | SOC | Labeling | Field | *Triticum* sp. | Silt loam and silty clay loam | 697 | 10.5 | Mixed with soil | 3 |
| (40) | Ag, roots | Litter incubation | SOC, POC, and MAOC | Labeling | Field | *Pinus ponderosa* | Sandy loam | 1774 | 17 | Mixed with soil | 1.6 |
| (41) | Roots | Litter incubation | SOC | Labeling | Field | *Triticum* sp. | Sandy clay loam, loamy and silty clay | 528, 1062, 1150, and 1052 | 16.7, 5.1, 4.5, and 4.7 | Mixed with soil | 1 |
| (42) | Ag | Litter incubation | SOC, POC, and MAOC | Labeling | Field | *Andropogon gerardii* | Silty clay | 835 | 12.9 | On soil surface | 3 |
| (43) | Ag | Litter incubation | SOC, POC, and MAOC | Natural abundance | Controlled | *Eucalyptus* sp. | Sandy clay loam | NA | NA | Mixed with soil | 0.66 |
| (44) | Ag | Litter incubation | SOC | Labeling | Field | *Acer saccharum* | Clay | 1098 | 3.8 | On soil surface | 1 |
| (45) | Roots | Litter incubation | SOC, POC, and MAOC | Labeling | Field | *Triticum* sp. | Silt loam and silty clay loam | 1155 and 1062 | 4.4 and 5.8 | Mixed with soil | 1 and 3 |
| (46) | Ag, roots | Litter incubation | SOC, POC, and MAOC | Labeling | Field | *Pinus ponderosa* | Sandy loam | 1774 | 17 | Mixed with soil | 5 |
| (47) | Ag, roots | Litter incubation | SOC, POC, and MAOC | Labeling | Controlled | *Zea mays* L. | Silty loam | NA | NA | Mixed with soil | 0.23 |
| (48) | Ag, roots | Litter incubation | SOC | Labeling | Field | *Pinus ponderosa* | Silty clay loam | 1774 | 17 | Mixed with soil | |
| (49) | Ag, roots | Litter incubation | SOC | Natural abundance | Controlled | *Populus simonii* Carr. | Loamy sand | NA | NA | Mixed with soil | 0.74 |
| (50) | Ag | Litter incubation | SOC, POC, and MAOC | Labeling | Field | *Beech forest* | ND | 930 | 8.4 | On soil surface | 1 |
| (51) | Ag | Litter incubation | SOC, POC, and MAOC | Labeling | Field | *Sinapis alba* | Silt loam | 803 | 7.4 | Mixed with soil | 1.56 |
| (52) | Ag | Litter incubation | SOC | Labeling | Field | *Populus nigra* | Loam | 734 | 15 | On soil surface | 0.92 |
| (53) | Roots | Litter incubation | SOC, POC, and MAOC | Labeling | Controlled | *Oilseed rape* | Sandy loam | NA | NA | Mixed with soil | 0.37 |
| (54) | Roots | Litter incubation | SOC, POC, and MAOC | Labeling | Field | *Triticum* sp. | Silt loam and silty clay loam | 600 | 10.5 | Mixed with soil | 3 |
| (55) | Ag, roots | Litter incubation | SOC | Labeling | Controlled | *Liquidambar styraciflua* L. | Silty clay loam | NA | NA | Ag on surface and roots mixed with soil | 0.07 |

*continued to next page*
| RN | C-input source | Experiment type | SOC fractions* | Method for new-C estimation | Environment condition | Plant species | Soil texture | MAP (mm) | MAT (°C) | Residue management | Time (years) |
|----|----------------|-----------------|----------------|----------------------------|-----------------------|---------------|-------------|-----------|----------|--------------------|--------------|
| 56 | Roots          | Litter incubation | SOC            | Labeling                   | Field                 | Acer rubrum   | Sand        | 838       | 6.8      | Mixed with soil     | 1 and 2      |
| 57 | Ag, roots      | Litter incubation | SOC, POC, and MAOC | Labeling                   | Controlled            | Triticum sp.  | Silt        | NA        | NA       | Mixed with soil     | 0.18         |
| 58 | Ag, roots      | Litter incubation | SOC            | Natural abundance          | Controlled            | Canavalia ensiformis and Gliricidia sepium | Clay         | NA        | NA       | Mixed with soil     | 2            |
| 59 | Ag, roots      | Litter incubation | SOC, POC, and MAOC | Natural abundance and labeling | Controlled            | Fagus sylvatica L. and Fraxinus excelsior L. | Silt loam    | NA        | NA       | Mixed with soil     | 0.56         |
| 60 | Ag             | Litter incubation | SOC, POC, and MAOC | Natural abundance          | Controlled            | Zea mays L.  | Silt loam   | NA        | NA       | Mixed with soil     | 0.13         |
| 61 | Ag, roots, NRm | Living plant     | SOC            | Labeling                   | Field                 | Secale cereale | Fine-loamy   | 1055      | 8.2      | No-Till            | 0.42 and 1.42 |
| 62 | Ag, roots, NR  | Living plant     | SOC, POC, and MAOC | Natural abundance          | Field                 | Zea mays L.  | Silt loam   | 650       | 10.5     | Tillage            | 4            |
| 63 | Ag, roots, NR  | Living plant     | SOC            | Treatments                 | Field                 | Zea mays L.  | Silt loam   | 970       | 10       | Tillage            | 6, 10 and 12 |
| 64 | Ag, roots, NR  | Living plant     | SOC            | Natural abundance          | Field                 | Zea mays L., Hordeum vulgare L., Triticum aestivum L., Trifolium pratense L., Phleum pratense L. | Silt loam    | 1200      | 4        | Tillage            | 15           |
| 65 | Ag and roots, NR | Living plant     | SOC            | Natural abundance          | Field                 | Zea mays L., Triticum sp. | Clay loam    | 551       | 5.7      | No-Till and tillage | 13           |
| 66 | Ag, roots, NR  | Living plant     | SOC, POC, and MAOC | Labeling                   | Field                 | Vicia dasycarpa Ten. | Silt loam and silty clay loam | 450       | 16.5     | Tillage            | 0.37         |
| 67 | Ag, roots, NR  | Living plant     | SOC            | Treatments                 | Field                 | Zea mays L.  | Silty clay loam | 906       | 10       | Tillage            | 11           |

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| RN | C-input source | Experiment type | SOC fractions* | Method for new-C estimation | Environment condition | Plant species | Soil texture | MAP (mm) | MAT (°C) | Residue management | Time (years) |
|----|----------------|-----------------|----------------|-----------------------------|-----------------------|---------------|--------------|-----------|-----------|-------------------|-------------|
| (67) | Ag, roots, NR | Living plant | POC | Natural abundance | Field | Zea mays L. and Glycine max L. | Silt loam | 1200 | 17 | No-Till | 2 |
| (68) | Ag, roots, NRm | Living plant | SOC | Labeling | Field | Vicia villosa Roth subsp. Villosa | Clay loam | 1053 | 13 | Tillage | 0.41 |
| (69) | Ag, roots, NRm | Living plant | SOC | Labeling | Field | Triticum aestivum L., Pisum sativum L., and Vicia sativa L. | Sandy loam | 769 | 19.3 | No-Till | 0.49 |

*POC: includes either the fractions with particle size higher than 53 μm or the light fraction separated by density (3). MAOC: includes either the fractions with particle size lower than 53 μm or the heavy fraction separated by density. It was assumed that SOC is equal to the sum of POC plus MAOC; therefore, when the studies reported only one fraction and total SOC, the second fraction was calculated by difference (70).
of roots, aboveground inputs, and rhizodeposition separately, specifically selecting papers that measured POC and MAOC contents to estimate POC<sub>FE</sub> and MAOC<sub>FE</sub> of all C-input sources.

**RESULTS AND DISCUSSION**

**Formation efficiencies of belowground and aboveground inputs into total SOC**

Within the reviewed experiments, SOC formation efficiencies (SOC<sub>FE</sub>) were relatively stable at different C-input levels (Fig. 2), as shown by the linear relationships between C-inputs and C in new SOC in both experimental methods (litter incubations and living plant experiments). Average aboveground-input-SOC<sub>FE</sub> and root-input-SOC<sub>FE</sub> were similar in litter incubation experiments (0.31 and 0.36, respectively) (Fig. 2A) but differed markedly in living plant experiments where average aboveground-input-SOC<sub>FE</sub> was 0.11, one-third the value of average belowground-input-SOC<sub>FE</sub> of 0.31 (belowground inputs included roots and rhizodeposition in this type of experiments) (Fig. 2B). These results suggest that SOC<sub>FE</sub> of traditional litter incubation experiments may overestimate real SOC<sub>FE</sub> occurring under normal field conditions, particularly for aboveground inputs. In addition, the higher belowground-input-SOC<sub>FE</sub> as compared to aboveground-inputs-SOC<sub>FE</sub> observed only in experiments with living roots suggests that reasons other than biochemical composition likely explain the higher retention of belowground inputs into bulk SOC. Our estimates agree with recently reported values for aboveground-input-SOC<sub>FE</sub> but are lower than reported values for belowground-input-SOC<sub>FE</sub> (12) because our calculations include rhizodeposition amounts plus roots (as belowground inputs) in the SOC<sub>FE</sub> estimation. Because belowground-input-SOC<sub>FE</sub> is typically calculated as the ratio between root biomass and belowground-derived-SOC, ignoring rhizodeposition as an input, reported values of belowground-input-SOC<sub>FE</sub> in living plant experiments may overestimate real formation efficiencies by almost 50% because net rhizodeposition is not included in the denominator (assuming a net rhizodeposition:root ratio of 0.5; see Table 2). Overall, our results suggest that SOC<sub>FE</sub> values measured in litter incubation experiments may not represent decomposition dynamics under field conditions with living plants and highlight the importance of estimating SOC<sub>FE</sub> in ways that consider all plant inputs including rhizodeposition.

**Formation efficiency of aboveground, root, and rhizodeposition inputs into POC and MAOC**

We found a significant effect of time on POC<sub>FE</sub> (P < 0.001; Table 4). POC<sub>FE</sub> decreased during the first year of experimentation, after which time it appears to stabilize, but MAOC<sub>FE</sub> estimates were not significantly related to time (evaluated using linear and nonlinear models) (Fig. 3). The different formation mechanisms of POC and
MAOC could explain these results. POC is largely made up of relatively undecomposed fragments of plant residues, whereas MAOC consists of microscopic fragments of organic material associated with soil minerals (7). Therefore, it is expected that new POC derived from plant inputs follows a decomposition pattern similar to that from litter inputs, and therefore, POCFE decreases across time. However, it is likely that new MAOC has a different decomposition pattern because it is rapidly formed during early stages of litter decomposition and then remains relatively stable because of soil protection (3). Thus, we expected a smaller or lack of correlation of MAOCFE with experimental time. We found these same results when analyzing two particular experiments in our dataset that resampled soils through experimental time and estimated POCFE and MAOCFE variations (see fig. S2).

![Figure 2](https://www.science.org)

**Fig. 2.** Relationships between C-inputs and new SOC formation in litter incubation experiments and living plant experiments. Panel A shows litter incubation experiments and panel B living plant experiments. Average SOC formation efficiencies (SOCFE) are shown for each type of inputs and experimental method. Regression lines are shown in red, and CI_{intercept} and CI_{slope} are the 95% confidence intervals (for the intercept and slope, respectively). Belowground C-inputs in living plant experiments include net rhizodeposition (see Materials and Methods).

### Table 2. Mean, SD, and sample size (n) net rhizodeposition-to-root ratios observed in this paper and in Pausch and Kuzyakov review (10).

| Ratio                      | Land use      | Mean | SD  | n   | Reference                  |
|----------------------------|---------------|------|-----|-----|----------------------------|
| Net rhizodeposition/root   | Crops         | 0.54 | 0.07| 99  |                            |
| biomass                   | Grasses       | 0.50 | 0.06| 128 | Pausch and Kuzyakov (10)   |
|                           | Trees         | 0.49 | 0.11| 9   |                            |
|                           | Crops and cover crops | 0.44 | 0.12| 7   | This review                |
|                           | All           | 0.51 | 0.07| 243 | Weighted average           |
After correcting for time effects, we found that belowground inputs have substantially higher formation efficiencies than aboveground inputs, not only in total SOC but also in both POC and MAOC fractions, with differential effects of roots, rhizodeposition, and aboveground inputs in the formation efficiencies of each soil fraction. In living plant experiments that reflect real plant growing conditions, belowground-POC<sub>FE</sub> (roots and rhizodeposition) was ~180% higher than aboveground-input-POC<sub>FE</sub>, with similar results for MAOC (~170% increase) (Fig. 4). However, different mechanisms seem to explain the observed higher POC<sub>FE</sub> and MAOC<sub>FE</sub> of roots and rhizodeposition, as discussed below.

Our results for the MAOC fraction suggest that plant rhizodeposition is critical for building SOC in stabilized pools because average net rhizodeposition-MAOC<sub>FE</sub> in living plant experiments was high (0.46 ± 0.21; Fig. 4B) compared to the MAOC<sub>FE</sub> of roots or aboveground inputs (~0.07; Fig. 4B). Belowground-MAOC<sub>FE</sub> in living plant experiments (0.19 ± 0.07) was nearly three times higher than MAOC<sub>FE</sub> of aboveground inputs (0.07 ± 0.03) in the same type of experiment, whereas root-input-MAOC<sub>FE</sub> did not differ from aboveground-input-MAOC<sub>FE</sub> in litter experiments (Fig. 4B). These results suggest that the presence of living roots is a more efficient way to increase MAOC. Our calculations show that the majority (more than 75%) of total belowground derived MAOC was contributed by rhizodeposition (Fig. 5). These results assume a net rhizodeposition:root biomass ratio of 0.51 (± 0.07, n = 242) extracted from a literature synthesis (Table 2), but a sensitivity analyses revealed that our estimates remain unchanged for a wide range of net rhizodeposition:root biomass ratio, varying only with net rhizodeposition:root biomass ratios lower than ~0.3, which are uncommon given that 0.3 is two SDs away from the mean (see the Supplementary Materials and fig. S1 for details).

Although several studies show that rhizodeposition promotes SOC decomposition (i.e., priming) (14, 15), our results suggest that in the MAOC fraction it may have the opposite result—MAOC formation. This result may occur because rhizodeposition is rich in simple carbohydrates (16) that are easily consumed by microbial activity that produce microbial necromass and microbial-derived compounds. These compounds are recovered in the dissolved organic C fraction, which is the dominant and more efficient pathway of MAOC formation (7, 13). In addition, sorption of other low–molecular weight compounds (e.g., organic acids) from rhizodeposition could be another pathway of MAOC formation. When fine minerals of soil (clay + silt) are nonsaturated with organic compounds (17), these simple molecules have a strong capacity to interact with minerals and contribute to MAOC formation (13, 18). Furthermore, MAOC<sub>FE</sub> of roots and aboveground inputs was not different in litter incubation experiments (~0.11), therefore suggesting that biochemical composition is less important for retaining C in this stable SOC fraction (Fig. 4B).

Mycorrhizal contribution to rhizodeposition is unclear but could partly explain the high net rhizodeposition-MAOC<sub>FE</sub> found in our study. Substantial quantities of rhizosphere exudates come from extraradical hyphae secretions, and, in contrast, death and turnover of mycorrhizal tissues may also contribute substantially as C-inputs to soil decomposers, therefore increasing microbial necromass and microbial-derived compounds (19). As discussed above, these two pathways (increasing the microbial necromass or producing low–molecular weight compounds) are key C-inputs for MAOC formation. Most of the data analyzed in our study came from croplands (Table 1), where fertilization practices or tillage could have reduced mycorrhizal abundance. Therefore, MAOC formation rates from rhizodeposition could be potentially higher in nonmanaged and more natural ecosystems with abundant mycorrhizal colonization.

In contrast to MAOC formation, root biomass was the more efficient C-source for forming POC because root-input-POC<sub>FE</sub> was highest in litter incubation experiments (0.19 ± 0.07), even higher than belowground-input-POC<sub>FE</sub> in living plant experiments (0.12 ± 0.06) where both root + rhizodeposition inputs are occurring jointly (Fig. 4A). This differential stabilization into POC is likely attributable to the chemical recalcitrance of roots because root-input-POC<sub>FE</sub> was higher than aboveground-input-POC<sub>FE</sub> (0.10 ± 0.07) in litter experiments (Fig. 4A). In such experiments, root and aboveground inputs are incubated under similar soil conditions without rhizodeposition, the main difference being their chemical composition. Our estimates also showed that the majority of belowground-derived-POC was contributed by roots (Fig. 5). However, because POC is usually...
a smaller fraction of total SOC in croplands (20), the chemical recalcitrance mechanism would usually play a minor role in the overall SOC formation (21, 22), with POC formation becoming relatively more important in forest and coarser textured soils (23). It was unclear from our data how root stimulation of soil aggregation promotes POC accumulation, as suggested previously (24, 25). The larger differences between aboveground-input-POC$_{FE}$ and belowground-input-POC$_{FE}$ in living plant experiments as compared to the difference between aboveground-input-POC$_{FE}$ and root-input-MAOC$_{FE}$ in litter incubation experiments may also suggest that root stimulation of soil aggregation could increase root conservation in POC in addition to its chemical recalcitrance.

Our results show that rhizodeposition increased MAOC formation (as explained above) but reduced POC formation, likely because it increased decomposition rates of new POC. Rhizodeposition seems to decrease POC$_{FE}$ based on observations comparing average root-input-POC$_{FE}$ in litter experiments (0.19 ± 0.07) with belowground-input-POC$_{FE}$ (root + rhizodeposition) in living plant experiments (0.12 ± 0.07) (Fig. 4A). Even assuming that rhizodeposition does not form POC because this soil fraction contains coarse plant materials and not simpler molecules of low molecular weight through rhizodeposition (9), and considering an average net rhizodeposition:root ratio of 0.5 ± 0.1 (Table 2), root-input-POC$_{FE}$ in living plant experiments would be 0.18, still lower than the observed in litter incubation experiments (0.19 ± 0.07). This lower root-input-POC$_{FE}$ in living plant experiments may be explained if root decomposition is increased by rhizodeposition inputs (26). This increased decomposition rate could explain the negative values of net rhizodeposition-POC$_{FE}$ (Fig. 4A) and the negative contribution of net rhizodeposition to belowground-derived-POC (Fig. 5). Other research suggests that rhizodeposition induces greater microbial activity and increases SOC mineralization (i.e., priming effect) (14), either by contributing compounds that serve as cometabolites or by facilitating the release of mineral protected C into more accessible pools (26). These
mechanisms not only could explain the increased decomposition rates of new POC but also could be operating on preexisting POC (i.e., “old POC”) producing a priming effect (not accounted for in our study). Therefore, in the presence of rhizodeposition, a higher proportion of roots and aboveground inputs may be respired due to an increase in POC decomposition rates (both new and old), as observed for new POC only in living plant experiments. Our results strongly suggest that rhizodeposition increases decomposition rates only for roots entering the POC fraction, where recalcitrance limits its decomposition (7), reconciling previous evidence showing that rhizodeposition may either induce SOC decomposition (in the POC fraction) or increase its formation (in the MAOC fraction).

**Shoot:root effects on C-input contributions to POC and MAOC formation**

When considering a shoot:root ratio of 1, 81% of the new SOC formed annually was derived from belowground inputs (root + rhizodeposition), but belowground contributions were 43% when the shoot:root ratio was 6 (Fig. 6A). SOC formation depends on both input formation efficiencies and total input amounts from vegetation. High aboveground inputs explain why even with shoot:root ratios of 6, our estimates show that SOC is mainly aboveground-derived, despite aboveground inputs having a relatively low POCFE and MAOCFE (Fig. 4). The relative contributions of aboveground and belowground
from 1 to 6, SOC FE decreased from 25 to 16%, representing a 36% shoot:root ratios (Fig. 6B). Therefore, when shoot:root ratios increased 327, SOC fractions is needed (34), which varies between 1 and 6 in our dataset (Fig. 6A and Table 3). To estimate C-input contributions to POC and MAOC formation, we combined three hypothetical shoot:root ratios (1, 3, and 6) with POC and MAOC formation efficiencies of aboveground input, roots, and rhizodeposition in living plant experiments (Fig. 4). For shoot:root ratios of 1 and 3, POC and MAOC were mainly formed from root and rhizodeposition inputs, respectively. However, when the shoot:root ratio increased to 6, both fractions were formed more from aboveground inputs (Fig. 6A). Overall, when considering all the experiments reviewed, SOCFE decreased with increasing shoot:root ratios (Fig. 6B). Therefore, when shoot:root ratios increased from 1 to 6, SOCFE decreased from 25 to 16%, representing a 36% decrease in total SOC formation efficiencies (Fig. 6A). These results confirm earlier studies, suggesting that increasing C allocation to roots (and rhizodeposition) may be an important tool for increasing SOC storage (13). Our estimates of root:shoot ratios do not consider crop harvesting, which would remove substantial amounts of aboveground biomass (grains) decreasing the proportion of POC and MAOC formed from aboveground inputs.

Predicting SOC dynamics is important for developing sustainable land use strategies in the context of global change. The latest discoveries in SOC dynamics are not typically included in traditional simulation models, and a new generation of models based on measurable SOC fractions is needed (3, 27, 28). Our findings on formation efficiencies of different C sources and shoot:root ratios could directly inform these models and improve management of the C cycle and soil C sequestration. We showed that roots and rhizodeposition are highly efficient C sources for POC and MAOC formation, respectively. Thus, the inclusion of plants with higher C allocation to belowground biomass may increase SOC stocks, especially in croplands where SOC stock depletion has occurred worldwide (29, 30). However, plant breeding has traditionally selected crops for aboveground C allocation because of its relation to harvestable products (31). Therefore, potential trade-offs between production and SOC formation (12), and multiple-objective breeding programs (32, 33), could emerge. A good strategy that would contribute to reducing these trade-offs could be the inclusion of service crops (cover crops) in agricultural rotations, with high root production and elevated rhizodeposition (34, 35). Our work shows that root production and rhizodeposition, often overlooked in agronomy and plant breeding programs, should be evaluated for novel C cycle management options.

**MATERIALS AND METHODS**

**Literature search and database building**

We searched for peer-reviewed studies that evaluated SOC formation from aboveground and/or belowground inputs in Scopus (www.scopus.com). The following combination of terms was used: (“decomposition” OR “humification” OR “mineralization” OR “stabilization”) AND (“soil organic carbon” OR “soil organic matter” OR “SOM” OR “SOC”) AND (“root” OR “litter” OR “belowground” OR “aboveground”) AND (“isotopes” OR “label” OR “labelled” OR “labeling” OR “labeled”). The search resulted in 248 articles. To build a database with the papers, a two-step selection process was carried out. First, titles and abstracts of all articles were examined to exclude those that clearly do not focus on SOC formation. Second, only full papers that met the above criteria were reviewed and the papers where SOC, POC, or MAOC formation efficiency was reported, or could be calculated from data, were kept. When a study...
had measurements through time, we emphasize the final (i.e., longest) sampling time. The isotopic techniques used as tracers included the isotopic labeling of plants or the use of plant species with a different natural abundance of $^{13}\text{C}$ ($C_3$ versus $C_4$ species). In addition to the selected papers, two older articles were also included in our review, because they were pioneer works in the topic, and although they did not use isotope tracers, their experimental setup allowed us to estimate separately belowground from aboveground SOC formation efficiencies (36, 37). The final database included 35 articles with 197 observations (Table 1).

Experiments were divided into litter incubation experiments and living plant experiments (Table 1), as explained in Introduction (Fig. 1). In litter incubation experiments, roots and aboveground plant tissues were added to the soil and, after a time period, the amount of new SOC formed from these tissues was measured (Fig. 1A). These experiments were carried out under controlled conditions in the laboratory or in the field (in situ) and assessed crops and forest litter. We included forest and grass treatments because no statistical differences were found between vegetation types (fig. S3). The living plant experiments have two phases. The first phase included the crop’s growing period, from sowing to crop termination. During this period of crop growth, C accumulates in aboveground and roots tissues, while gross rhizodeposition occurs and net rhizodeposition can be measured at crop termination in the soil (Fig. 1B). C accumulated at crop termination in roots and net rhizodeposition is considered as the total belowground inputs, and C accumulated in aboveground litter as the total aboveground inputs (Fig. 1B). In the second phase of living plant experiments, aboveground, roots, and net rhizodeposition begin to decompose and form new SOC (similar to the litter incubation experiments; Fig. 1A). All the living plant experiments were carried out under field conditions, excepting Comeau et al. (38), where plants were grown under greenhouse conditions.

Data analysis
The data collected from the reviewed papers included experimental method (litter incubation or living plant experiment), soil texture, soil depth, experiment time, initial SOC, total new SOC, new SOC in each soil fraction (POC or MAOC), and C-input sources [aboveground, root, or net rhizodeposition (when available)]. When net rhizodeposition was neither measured nor estimated, it was assumed as 50% of root biomass (Table 2). Because data distribution was not normal [tested with Shapiro test (71)], differences between groups (Fig. 4) were assessed with Wilcoxon test (72). All statistical analyses were performed with R software version 4.0.2 with stats package (73).

Estimates of formation efficiencies of aboveground, roots, and rhizodeposition into POC and MAOC
Formation efficiencies of a given C-input (aboveground, root, and belowground) were estimated by dividing the amount of input by the amount of new SOC (POC or MAOC) formed from that C-input (Eq. 1)

$$
\text{C-input-SOC}_{\text{FE}} = \frac{\text{New C derived from C-input/input amount}}{(1 - f_{\text{nr}}) * \text{root-SOC}_{\text{FE}}}
$$

To differentiate POC and MAOC formation efficiencies originated from rhizodeposition or roots, two steps were made. First, we considered that belowground-input-POC$_{\text{FE}}$ and belowground-input-MAOC$_{\text{FE}}$ are a weighted average of net rhizodeposition and root formation efficiencies (Eqs. 2 and 3)

$$
\text{Belowground-input-POC}_{\text{FE}} = f_{\text{nr}} * \text{net rhizodeposition-POC}_{\text{FE}} + (1 - f_{\text{nr}}) * \text{root-POC}_{\text{FE}}
$$

$$
\text{Belowground-input-MAOC}_{\text{FE}} = f_{\text{nr}} * \text{net rhizodeposition-MAOC}_{\text{FE}} + (1 - f_{\text{nr}}) * \text{root-MAOC}_{\text{FE}}
$$

where Belowground-input-POC$_{\text{FE}}$ is the formation efficiency of belowground input to POC, $f_{\text{nr}}$ is the fraction of belowground input that is net rhizodeposition, net rhizodeposition-POC$_{\text{FE}}$ is the formation efficiency of net rhizodeposition to POC, root-POC$_{\text{FE}}$ is the formation efficiency of roots to POC, Belowground-input-MAOC$_{\text{FE}}$ is the formation efficiency of belowground inputs to MAOC, net rhizodeposition-MAOC$_{\text{FE}}$ is the formation efficiency of net rhizodeposition to MAOC, and root-MAOC$_{\text{FE}}$ is the formation efficiency of roots to MAOC. It is important to note that in previous works that estimate SOC formation efficiencies [see review by Jackson et al. (12)], estimates are based only on root biomass, ignoring rhizodeposition inputs, and therefore, SOC formation efficiencies of roots are overestimated and will be lower if rhizodeposition is considered as an input.

Second, we solved Eqs. 2 and 3 to estimate net rhizodeposition-POC$_{\text{FE}}$ and net rhizodeposition-MAOC$_{\text{FE}}$ (Eqs. 4 and 5) for each living plant experiment

$$
\text{Net rhizodeposition-POC}_{\text{FE}} = \text{Belowground-input-POC}_{\text{FE}} - \frac{f_{\text{nr}} * \text{root-POC}_{\text{FE}}}{f_{\text{nr}}}
$$

$$
\text{Net rhizodeposition-MAOC}_{\text{FE}} = \text{Belowground-input-MAOC}_{\text{FE}} - \frac{f_{\text{nr}} * \text{root-MAOC}_{\text{FE}}}{f_{\text{nr}}}
$$

where

1. Belowground-POC$_{\text{FE}}$ and belowground-MAOC$_{\text{FE}}$ were taken from the observed values in living plant experiments.
2. Root-POC$_{\text{FE}}$ and root-MAOC$_{\text{FE}}$ were assumed to be the median of observed values in the litter incubation experiments (0.18 and 0.06, respectively) because in these experiments roots are added to the soil manually and therefore no rhizodeposition exists. We used the median instead of the average for this assumption because median is a measure of central tendency not distorted by skewed data (74).
3. The net rhizodeposition:root ratio was assumed to be 0.5, and therefore, the fraction of net rhizodeposition ($f_{\text{nr}}$) used was 0.33 of total belowground inputs (see Table 2).

Estimates of the relative contributions of root and net rhizodeposition to the total belowground-derived SOC pool
After solving Eqs. 4 and 5, we estimated the contributions of roots and net rhizodeposition to new POC and new MAOC as a percentage of total belowground inputs (Fig. 5), with Eqs. 6 to 9 as follows

$$
\text{Root contribution to POC} (%) = \left( \frac{(1 - f_{\text{nr}}) * \text{root-POC}_{\text{FE}}}{f_{\text{nr}} * \text{net rhizodeposition-POC}_{\text{FE}} + (1 - f_{\text{nr}}) * \text{root-POC}_{\text{FE}}} \right) * 100
$$

$$
\text{Net rhizodeposition contribution to POC} (%) = \left( \frac{f_{\text{nr}} * \text{net rhizodeposition-POC}_{\text{FE}}}{f_{\text{nr}} * \text{net rhizodeposition-POC}_{\text{FE}} + (1 - f_{\text{nr}}) * \text{root-POC}_{\text{FE}}} \right) * 100
$$

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Root contribution to MAOC (%) = \((1-f_{nt})\times f_r\times MAOC_{FE}\) / \((f_{nt}\times f_r\times MAOC_{FE} + (1-f_{nt})\times f_r\times MAOC_{FE})\times 100\) (8)

Net rhizodeposition contribution to MAOC (%) = \((f_{nt}\times f_r\times MAOC_{FE}) / (f_{nt}\times f_r\times MAOC_{FE} + (1-f_{nt})\times f_r\times MAOC_{FE})\times 100\) (9)

**Estimates of new POC and MAOC formation under varying shoot:root ratios**

The contribution of different C-inputs to SOC pools depends not only on formation efficiencies but also on the relative amount of inputs added to the soil, which depends on plant allocation patterns (i.e., shoot:root ratio). To evaluate the impact of different shoot:root ratios on SOC formation, we performed an additional analysis (Fig. 6). We varied the relative contributions of C-inputs assuming shoot:root ratios of 1, 3, and 6 (which are within the reported range in the reviewed experiments; see Table 4) and multiplied them by the aboveground-input-POC, root-POC, net rhizodeposition-POC, aboveground-input-MAOC, root-MAOC, or net rhizodeposition-MAOC estimated for each experiment to calculate the relative contributions (in percentage) of each C-input to the different pools of new SOC formed, as follows

New C derived from aboveground (%) = 100 \times f_a \times \text{aboveground-POC or MAOC}_{FE} (10)

New C derived from root (%) = 100 \times f_r \times \text{root-POC or MAOC}_{FE} (11)

New C derived from net rhizodeposition (%) = 100 \times f_{nt} \times \text{net rhizodeposition-POC or MAOC}_{FE} (12)

where \(f_a\), \(f_r\), and \(f_{nt}\) are the fractions of total input (\(f_a + f_r + f_{nt} = 1\)) that is aboveground, root, or net rhizodeposition, respectively. These fractions vary with the different shoot:root ratios considered.

**SUPPLEMENTARY MATERIALS**

Supplemental material for this article is available at http://advances.sciencemag.org/cgi/content/full/7/16/eabd3176/DC1

View/request a protocol for this paper from Bio-protocol.

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Plant rhizodeposition: A key factor for soil organic matter formation in stable fractions
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