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Embankment as a carbon sink: a study on carbon sequestration
pathways and mechanisms in topsoil and exposed subsoil

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“Se l’organizzazione del carbonio non si svolgesse quotidianamente attorno a noi, [...], dovunque affiori il verde di una foglia, le spetterebbe di pieno diritto il nome di miracolo.”

Primo Levi, ‘La tavola periodica’ , Einaudi, 1975, pp. 261

“If the elaboration of carbon were not a common daily occurrence, [...], wherever the green of a leaf appears, it would by full right deserve to be called a miracle.”

Primo Levi, ‘The periodic table’ , Einaudi, 1975, pp. 261

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Résumé

La séquestration du carbone (C) fait l'objet d'une attention scientifique et politique croissante dans le cadre de la réduction des gaz à effet de serre. Cependant, les sols géotechniques ont été négligés en raison de leur potentiel de séquestration du carbone, et l'attention mondiale étant concentrée sur les sols agricoles et naturels. Dans le présent projet de thèse, nous visons à évaluer le potentiel des talus géotechniques comme puits de carbone et, par l'étude des espèces végétales et des sols présentant des caractéristiques contrastées, à mettre en lumière les mécanismes de séquestration du carbone organique et les rôles des différents acteurs impliqués. Nous visons non seulement à quantifier le C gagné et perdu dans le sol, mais aussi son origine (nouveau C frais et ancien C préexistant) et comment il est réparti dans différents pools de C qui montrent une stabilité du C différente (qualité du C stocké). Tout d'abord, nous avons évalué la séquestration du carbone dans différents pools de carbone sous un sol semé de 12 espèces herbacées différentes dans une expérience de 10 mois. La caractérisation des différents traits racinaires a permis de comprendre l'influence de la stratégie d'alimentation des ressources en racines (représentée par le spectre économique de la racine) sur la séquestration du carbone. Nous avons montré que les espèces dont les caractéristiques racinaires sont associées à une production élevée de C labile entraînent une augmentation plus élevée de C dans le pool stable de SILT+CLAY (<20 μ m). Les espèces dont les traits racinaires sont associés à un faible apport de C récalcitrant favorisent plutôt l'accumulation dans la fraction POM instable. Ensuite, grâce à une expérience de marquage isotopique stable de 183 jours (CO₂ constamment enrichi en ¹³C), nous avons pu étudier la dynamique du C dans différents pools de C sous deux espèces (*Lolium perenne* and *Medicago sativa*) sur deux sols (terre végétale, profondeur 0-30 cm et sol remonté, profondeur 110-140 cm) aux caractéristiques opposées. Nous avons mis en évidence le grand intérêt de faire le pont entre l'origine du C et les pools de C lors de l'étude des destins du C du sol, ce qui permet de dévoiler des processus que les méthodes plus traditionnelles cachent. Le nouveau C et l'ancien C présentaient une covariation synergique, avec des pertes plus faibles de l'ancien C associées à de nouvelles entrées de C plus élevées. Ceci est conforme à l'hypothèse de l'utilisation préférentielle du substrat. L'hypothèse de l'Utilisation Préférentielle des Substrats a également été validée par l'étude de "priming effect" et de la respiration du sol, qui a montré que la concentration de C provenaient par le plantes dans le CO₂ inhalé par le sol était plus élevée lorsque l'apport de C par les plantes était élevé, au contraire augmentant la concentration de C provenaient par la minéralisation de l'anciennes C lorsque les input de C par le plants étaient faibles, c.-à-d. en sous sol. De plus, nous avons validé l'hypothèse de réconciliation entre 'l'hypothèse de l'Utilisation Préférentielle des Substrats' et 'l'hypothèse de la Concurrence', cette dernière déterminant le 'priming effect' dans le sous-sol à faible fertilité. Nous avons observé de nouveaux apports significatifs de C d'origine végétale dans la fraction SILT+CLAY (<20 μ m, très stable) à l'appui de la preuve de l'effet d'entombage in vivo dans l'hypothèse de la pompe à carbone microbienne du sol. L'effet de l'espèce s'est produit principalement sur les intrants de nouveaux C, mais il a été maîtrisé par l'effet du sol, avec un stockage de C plus faible dans un sol de faible qualité (faible activité et biomasse d'azote et microbienne). En général, les conditions microbiologiques ont été le principal moteur de la nouvelle accumulation de C et de l'ancienne perte de C et ont aidé à expliquer pourquoi aucun effet de la saturation en C du sol - une théorie centrale dans des études récentes sur la séquestration de C - n'a été trouvé dans le carbone protégé. Cette compréhension fondamentale des interactions plantes-sol nous aide à mieux optimiser la gestion des sols et de la végétation pour la revégétalisation des talus des routes.

Abstract

Carbon (C) sequestration is receiving increasing scientific and political attention in a framework of greenhouse gasses mitigation. However, geotechnical soils have been neglected for their C sequestration potential, with the global attention focusing on agricultural and natural soils. In the present thesis project, we aim to assess the potential of geotechnical embankments as C sink, and, through the study of plant species and soils showing contrasting features, shed light on C sequestration mechanisms and the role of the different actors involved. We aim not only to quantify the C gained and lost in soil, but even its origin (fresh new C input or old preexistent C) and how it is partitioned in different C pools characterized by different C stability (quality of stored C). First, we evaluated the C storage in different pools under soil sowed with 12 different herbaceous species in a 10 months experiment. Assessing different root traits allowed understanding the influence of root economic spectrum on C storage. We showed how traits linked to high labile C are linked to a higher C increase in the stable SILT+CLAY pool (<20µm). Root traits related to a low input of recalcitrant, instead, favor accumulation in the unstable POM fraction. Thanks to a 183 days stable isotope labelling experiment (CO₂ constantly enriched with ¹³C) we were able to study the C dynamics in different C pools under two species (*Lolium perenne* and *Medicago sativa*) sowed on two soil (topsoil, 0-30cm depth and subsoil brought to the surface, 110-140 cm depth) showing contrasting characteristics. We evidenced the great interest of bridging C origin and C pools when studying soil C fates, allowing unveiling processes those more traditional methods would hide. New C and old C showed synergetic covariation, with lower old C losses associated to higher new C inputs. This is in good accordance with the Preferential Substrate Utilization hypothesis. The Preferential Substrate Utilization hypothesis was also validated with the study of priming effect and soil respiration, that showed higher plant derived C in respired CO₂ when plant C input was high, while increasing old C mineralization when plant C input was low, i.e. in subsoil. We observed significant plant derived new C input in the SILT+CLAY fraction (<20µm, highly stable) supporting evidence of the in vivo entombing effect in the soil Microbial Carbon Pump hypothesis. The species effect mainly occurred on new C input, but it was overpowered by the soil effect, with lower C storage in low quality soil (low nitrogen and microbial biomass and activity). In general, microbiological conditions were the main driver for new C accumulation and old C loss, and helped to explain why no effect of soil C saturation – a central theory in recent studies on C sequestration - was found in the protected carbon. Such fundamental understanding of plant-soil interactions helps us to better optimize soil and vegetation management for road embankment revegetation

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CHAPTER I: General introduction

1 1.1.CONTEXT

2 Soil holds the second largest terrestrial carbon (C) pool (1500 to 2400 GtC to a depth of one meter, IPCC
3 2014; Adams et al. 1990; Anderson 1992; Eswaran et al. 1993; Batjes 1996) with possibly another 900Gt
4 at a depth of 1-2 m (Batjes 1996, Jobbagy et al. 2000), after the lithosphere but in front of vegetation (350
5 to 550 GtC, mainly in forests) and atmosphere (829 GtC, IPCC 2014). Soil shares the common interface
6 with all the other spheres and thus plays a key role in driving the global C cycle. How to prevent C loss from
7 soil and how to sequester more C into soils has become one of the most important scientific and political
8 quests in global change biology (Sauerbeck 2001; Lal 2004). The European Union is actively involved in this
9 issue, and the topsoil soil organic C content is an official indicator for the EU sustainable Development
10 Goals (EU-SDG, 2018), leading to the funding and supervision of several programs focused on soil
11 conservation and soil C increase. Some examples, among others, that involve assessment of soil organic
12 carbon and potential sequestration, showing the interest and importance of this topic, are the CIRCASA
13 project (<https://www.circasa-project.eu/>), LANDMARK project (<http://landmark2020.eu/>), iSQAPER
14 project (<http://www.isqaper-project.eu/>), and LUCAS project
15 (<https://esdac.jrc.ec.europa.eu/projects/lucas>), all funded in the framework of Horizon2020. The FAO is
16 also involved in numerous projects focusing on soil health, that among other things underline the
17 importance of soil C increase for climate change mitigation, like GSOCmap
18 (<http://54.229.242.119/GSOCmap/>) or the Intergovernmental Technical Panel on Soils (ITPS)
19 (<http://www.fao.org/global-soil-partnership/intergovernmental-technical-panel-soils/en/>). Similarly, on a
20 national basis, different projects have been developed focusing on the potential of soil C storage for
21 climate change mitigation. One of the most striking examples is the 4p1000 initiative
22 (<https://www.4p1000.org/>), launched by France on 1 December 2015 at the COP 21, stating that
23 increasing by 4 ‰ the soil C stock in agricultural soils would completely remove the excess of CO₂ in the
24 atmosphere produce by anthropic actions. However, studies on strategies of C sequestration in soils are

25 mainly limited to “green systems” (e.g. forests, grasslands, plantations, croplands, wetland etc.), where
26 the soils are considered to be or potentially to be, a C sink. We argue that, in an irrevocable era of
27 industrialization and urbanization, soil in “grey systems” connected with geotechnical infrastructure
28 industry must be taken into consideration for soil C sequestration. There are two main reasons for that: i)
29 the high environmental impact of geotechnical industry, in particular on CO₂ emissions, that needs to be
30 mitigated, and ii) the drastic increase of geotechnical infrastructures, in particular road and railroads,
31 which means that soils connected to geotechnical infrastructures can no longer be ignored for their
32 potential ecosystem services, among which is soil C sequestration.

33

34 *1.1.1. Impact of geotechnical structures on greenhouse gasses emissions and TERRE* 35 *project*

36 It is well known how construction activities and practices commonly related to geotechnical engineering
37 have a high environmental impact, negatively influencing climate change, soil sealing, erosion,
38 deforestation, desertification, ozone depletion and general air/water/soil pollution (Kibert 2008; Misra
39 and Basu, 2011). Regarding the impact of these practices on CO₂ emissions, numbers can vary according
40 to different sources, but there is general agreement that construction and infrastructure have a high
41 impact on global greenhouse gas emissions. Global CO₂ emissions from construction work are attested in
42 a range of 25-40% of the total CO₂ emissions (Dixit et al. 2010, O’Riordan et al. 2011). It is well established
43 among researchers, policy makers and practitioners how a switch towards sustainable geotechnical
44 solutions is not only desirable but absolutely vital to face the challenges of climate change mitigation, and
45 to move toward a sustainable future (Dejong et al., 2011; Misra and Basu, 2011; Gallipoli and Mendes,
46 2017). In this optic, the current thesis is financed by the Marie Skłodowska-Curie Innovative Training
47 Networks (ITN-ETN) TERRE (<http://www.terre-etn.com/>): Training Engineers and Researchers to Rethink
48 geotechnical Engineering for a low carbon future. The aim of the TERRE project is to develop new geo-

49 technologies to address the challenge of a low carbon impact European construction industry. In the TERRE
50 action, multiple interdisciplinary projects have been developed under a wide umbrella of practices:
51 optimization of energy requirement for construction, role of plants to increase soil stability via root
52 reinforcement and hydraulic suction, new low C impact materials for construction, etc. The present thesis
53 project aims to investigate the role of geotechnical embankments as C sinks.

54

55 *1.1.2. Road and railroad development*

56 In the last decades, highways and railroad systems have dramatically increased their surface. Especially in
57 developing and emerging countries, major investments have been made to increase and expand the
58 infrastructure systems, since connections among countries and cities are one of the fundamental aspects
59 of economic growth. Globally, the railroad system increased its length by 100000km in the last year
60 (<https://data.worldbank.org/topic/infrastructure>). The Chinese public roads passed from 3.5 to 4.8 million
61 km in the last 10 years (<http://statista.com>). In India, in the last 4 years, the length of the highways
62 increased by 60000km, and other 200000km of highways are expected to be finished by 2022
63 (<https://www.ibef.org/industry/roads-india.aspx>). Another striking example of the massive future
64 infrastructure development is the China's 'Belt and Road' initiative, planning to connect via a complex
65 system of roads and maritime route, Asia, Africa and Europe. Together with the development of the Trans-
66 African Highway, consisting of 60000km of highways started in 1971 by United Nations Economic
67 Commission for Africa and not yet completed, we have a picture of the dramatic increase of the global
68 infrastructure system. This overview clearly shows how the soils connected with geotechnical work, in
69 particular with the construction of road and railroad infrastructures (hereafter 'geotechnical soils'), are
70 increasingly important and any potential benefits and ecosystem services need to be explored. Road and

71 railroad embankments play a pivotal role in the interactions between environment and infrastructure, and
72 a correct design could increase the ecosystem services they can provide.

73

74 *1.1.3. Development of sustainable geotechnical practices*
75 Efforts to increase the sustainability of geotechnical structures have already been made. Practices included
76 the use of alternative ecofriendly materials, use of bio-engineering on slopes, reuse and restoration of
77 older structures, underground energy storage, and use of geothermal energy (Misra and Basu, 2011).
78 However, all these practices are based on the reduction of CO₂ emissions. Such a framework relies on a
79 passive role of geotechnical structures (new technologies to reduce CO₂ emission), but ignoring the
80 potential active role that geotechnical soil can have in reducing atmospheric CO₂ concentration via soil
81 organic C sequestration.

82

83 *1.1.4. Geotechnical embankments: a new hotspot for soil carbon sequestration?*
84 Geotechnical soils present some unique features that could potentially make them achieve efficient soil
85 carbon sequestration. The main general feature is that geotechnical soils do not present specific
86 constraints regarding their use. The objective of agricultural soils is the production of food and goods for
87 direct consumption or to be placed on the market. Therefore, agricultural soils have an “economical
88 constraint”, and the objective of stakeholders, even in a framework of sustainability, is to increase or
89 maintain production without depleting soils. In a natural ecosystem, it is possible to talk about an
90 ‘ecological constraint’, in the sense that it is not possible to modify the environment to increase soil C
91 storage without disturbing the ecological balance and networks of the systems, affecting the health of the
92 system itself and, ultimately, the ecosystem services that it provides to the community.

93 Geotechnical soils, instead, are heavily anthropized soils, where the ecological balance has already been
94 disturbed. Moreover, soils are frequently moved from other areas or dug and brought to the surface,
95 changing the soil composition, microbiology, fertility and, ultimately, their ecological value. Vegetation
96 planted on geotechnical soils, especially on embankments, is not used for agricultural production. Re-
97 vegetation is therefore artificially implanted, and there are few ecological or economic constraints. These
98 soils and the plants used for revegetation can be chosen and planned to promote regulating, supporting
99 and cultural ecosystem services, including embankment stability maintenance, erosion control, noise
100 dissipation, traffic air pollution isolation, biodiversity conservation and aesthetical effect against driver
101 fatigue. Among these ecosystem services, in particular, we argue that, contrary to agricultural and natural
102 systems, geotechnical soils can be actively designed for CO₂ sequestration. Dejong et al. (2011) advocated
103 the possibility of using geotechnical soils to efficiently store C by i) selecting plants that efficiently fix and
104 move C into soil, ii) study different microbial communities that influence soil C cycle and the potential of
105 inoculation, iii) selection of different soils with higher potential for organomineral interaction and C
106 protection, and iv) using soil improvers (like recycled concrete and furnace slag) to increase C
107 sequestration. However, no specific studies have been implemented to really investigate the C
108 sequestration potential of geotechnical soils and how to maximize it. Therefore, in this thesis I aim to start
109 investigating the potential for designing efficient C sequestering embankments, starting with the main
110 issue of soil and plant selection.

111

112 *1.1.5. Embankment design*

113 When designing an embankment, the structure is based on a core of clay soil compacted according to a
114 Proctor compaction test, to achieve maximum dry density (Standard Australia, 2003). This compacted soil
115 core is usually covered with a 30-50 cm layer of uncompacted soil for revegetation (Fig. 1). The

116 construction and design of the embankment is outside the scope of this research, where only the soil layer
117 used for revegetation is considered for potential soil C storage. This soil layer is usually stripped topsoil (\cong
118 30cm) collected in the area and conserved, while the clay core is usually subsoil excavated, mixed and, if
119 needed, adjusted with additional soils or soil improvers to achieve the optimal density level to support the
120 structure. However, often the layer of topsoil is collected and transported to the construction site from
121 other fertile areas, with a high impact on CO₂ emissions (for the transport) and on environment (for the
122 ecological value of fertile topsoil). We argue that, to improve the sustainability of the embankments,
123 instead of using valuable topsoil for revegetation, mineral subsoil (> 1m depth) collected in the area can
124 be prepared and used for revegetation. Compared to stripped topsoil, subsoil embankments are more
125 economically interesting, but usually demand higher constraints in plant selection due to their less
126 favorable growing conditions (although recent soil inoculation techniques can improve this). Herbaceous
127 plants are essential materials for embankment revegetation. Herbaceous plants usually demand low
128 maintenance cost and intensity, with one or two cuts per year to maintain vegetation vigor. The choice of
129 soil (organic topsoil *versus* mineral subsoil) and vegetation will deeply influence the potential for C
130 sequestration.

131 However, to effectively enhance C sequestration in geotechnical soils, a better understanding of the
132 mechanisms behind the plant-soil C-cycle is necessary. There is a need to understand the influence of
133 different plants on soil C sequestration and their relationships with soil and microbiological communities,
134 to allow the design of the best practices for soil C sequestration, in geotechnical and non-geotechnical
135 soils.

136

137 *1.1.6. Plants: the primary source of carbon input in soil*

138 Plants act as conduits to transport C from atmosphere to soil (Fig. 2). Plants regulate the uptake and
139 fixation of CO₂ in different organic forms via photosynthesis, using water and atmospheric CO₂ as 'raw
140 materials' and light as an energy source (Chan, 2008). Plants also regulate the input of C in soil via two
141 main processes: 1) plant biomass from roots and shoots in the form of litter, forming soil particulate
142 organic matter (POM) and 2) root exudates and other labile C compounds released by roots during plant
143 growth (Hungate et al. 1997; Lal, 2004) (Fig. 2).

144 With regard to C input, the first process strongly influencing the C-cycle is C input in forms of residues
145 derived from vegetation. The selection of plants can considerably influence the C input in soil in terms of
146 quantity (biomass production) and quality. Biomass production and related amount of C input is not the
147 only driver for soil organic C accumulation. It has now been observed that the litter quality, especially
148 regarding the C:N ratio of plant tissues, will strongly influence the decomposability of POM and its
149 residence time in the soil (Castellano et al. 2015). C from exudates also represents a major amount of C
150 that plants transfer from the atmosphere to soil (Balesdent and Balabane, 1996). Estimates vary from more
151 conservative values, such as 5 - 33% of daily photoassimilates (De Deyn et al. 2008), to 40-60% (Högberg
152 et al. 2001; Clemmensen et al. 2013; Keiluweit et al. 2015;) to up to 80% of photosynthetically assimilated
153 C moved in 10 days via exudates in soil (Reid and Mexal, 1977). The input from exudates has traditionally
154 been seen as the 'labile C input' that is consumed and respired quickly in the soil system. However, recent
155 studies showed how C protected via organomineral complexation on minerals and in aggregates mainly
156 derive from plant exudates or microbiological exudates and exopolysaccharides, that in turn originate from
157 plant labile C input consumption and complexation (Lorenz and Lal 2005; Six et al. 2006; Cotrufo et al.,
158 2013; Vidal et al., 2018).

159 In a recent review, Poirier *et al.*, (2018) has argued that the root traits that most influence C stabilization
160 are those related to chemical composition, root exudation and the presence of symbionts (mycorrhizas
161 and dinitrogen (N₂) -fixing *Rhizobium*), whereas the role of morphological traits is not yet clear. More
162 specifically, root traits increasing recalcitrance promote short-term C stabilization by slowing
163 decomposition rates, but traits that reduce recalcitrance contribute to long-term C stabilization via the
164 reaction of microbial products with mineral surfaces. Although several studies have analyzed the link
165 between plant functional traits, microbial activity and C accumulation (Chapin 2003; Lavorel *et al.*, 2007;
166 De Deyn *et al.*, 2008; Poirier *et al.* 2018), as yet, no study has focused on how root growth and specific
167 traits can alter the accumulation and potential persistence of different soil C pools, that are linked to the
168 physical structure of soil itself (see Cardinael *et al.*, 2015; Fujisaki *et al.* 2018).

169 C entering the soil can face two main fates: be consumed by microorganisms and leave the soil pool via
170 microbiological respiration, or be stored in the soil for different periods of time, often after being
171 processed by microorganisms.

172

173 **1.1.7. Microbiological communities: the carbon pump in different soil fractions**

174 Microbiological communities can be identified as a further main actor for the C storage in soil (Fig. 2). Soil
175 organic C consumption by microorganisms will mainly depend by their substrate use efficiency, meaning
176 the proportion of the C used by microorganisms for biomass growth or enzyme production (part of the C
177 stock in soil) and the C respired or mineralized (Lekkerkerk *et al.*, 1990). The balance between these two
178 fluxes, accumulation in biomass and/or via microbial exudation and loss via respiration, will depend on
179 different factors related to substrate quality, (C:N ratio, molecular complexity, molecular weight and
180 solubility) and the efficiency of different microbiological communities to degrade organic C in soil
181 (Lekkerkerk *et al.*, 1990, Cotrufo *et al.*, 2013) which can vary by microbiological abundance, composition

182 and partition between bacterial and fungal communities (Six et al., 2006). Microorganisms are also mainly
183 responsible for C transformation in soil, consuming C input of plants in form of exudates or root debris,
184 and 'pumping' it into the soil structures and in contact with mineral surfaces, in the form of microbial
185 exudates and exopolysaccharides (Cotrufo et al., 2013; Vidal et al.; 2018). This active role of microbes have
186 been formalized by the conceptual framework of 'microbial C pump' by Liang et al. (2017). However, this
187 framework does not consider the destination of microbial derived C in different soil C pools. When in
188 symbiosis with plants, the C substrates that are assimilated by microorganisms at the root apex are utilized
189 rapidly for respiration and growth, or lost as microbial exudates or exopolysaccharides that are used as a
190 substrate for subsequent microbial communities. Certain microbial species, such as *Rhizobium*, present in
191 nodules of N₂-fixing species, produce large amounts of exopolysaccharides (Downie, 2010) that can also
192 be adsorbed onto fine silt and clay particles (Fehrmann and Weaver, 1978).

193

194 ***1.1.8. Soil: responsible for carbon protection***

195 Finally, the last main actor to consider in the determination of the fate of soil organic C is the soil itself
196 (Fig. 2). The residence time of C is controlled by the protection mechanisms that contribute to stabilize it
197 (Luo et al. 2004; Jastrow et al. 2005). C in soil can be divided into three main pools: an unprotected C pool,
198 referring to the labile particulate organic matter (POM) in the soil, a biochemically protected pool (BPC)
199 (Fig. 3), when C is moved in soil in recalcitrant forms and is difficult for microorganisms to consume it, and
200 a physically protected pool (PPC), when C is protected inside aggregates or absorbed on clay/silt particles
201 and cannot easily be consumed by microorganisms (Fig. 3). The POM and BPC pool fate depends on the
202 nature of the organic matter and the microbiological communities, as discussed previously. The PPC is
203 considered to be the most stable C pool, and therefore the most important for soil C storage (Rumpel et
204 al. 2012). Regarding the PPC, it is particularly worthwhile to explore the linkage between PPC and labile C

205 from plants, i.e. C from exudates. Exudates were usually considered to be immediately consumed by
206 microorganisms and to play a marginal role in C sequestration (van Veen et al. 1991; Van Geijn et al. 1993;
207 Hungate et al. 1997). This assumption is now questioned by more recent studies that state that labile soil
208 C compounds are just partially consumed, and dissolved labile organic C can be protected by soil
209 absorption inside aggregates or on clay and silt (see review by Kalbitz and Kaiser, 2008). Moreover, it has
210 been demonstrated that microbiological exudates and exopolysaccharides are the main precursor of
211 organomineral protected C (Cotrufo et al., 2013; Vidal et al., 2018). To understand the fate of C and
212 increase the PPC, the two main mechanisms of C protection need to be investigated.

213

214 *1.1.9. Soil structure and carbon physical protection in aggregates*
215 Aggregate protection of C is due to the C physical protection from microorganisms by occlusion of C in the
216 smaller pores, limiting the gas and nutrient diffusion and, therefore, microbiological activity, and
217 separating enzymes from substrates on mineral and humic surfaces (O'Brien and Jastrow, 2013).
218 Aggregates are formed by binding of soil particles by fine roots and fungal hyphae (Tisdall and Oades 1982)
219 and cementation by microbiological and plants exudates, like glycoproteins, polysaccharides, and
220 mucilage, directly influencing the stability of aggregates (Tisdall and Oades 1982; Caesar-Tonthat 2002;
221 Nichols and Wright 2005). The formation of aggregates is different in regards to their size. Abiotic factor,
222 such as ligand exchange and polyvalent cation bridging promoted by drying-rewetting cycles (Bronick and
223 Lal, 2005; Keil and Mayer, 2014), are known to form stable microaggregates. Microaggregates are then
224 complexed in small macroaggregates thanks to the biotic action and cementation from microbes that
225 produce extracellular polymeric substances acting as glues to connect soil aggregates (Blankinship et al.,
226 2016). Fine roots and hyphae of fungi further complex aggregates in bigger structures thanks to their
227 enmeshing action (Tisdall and Oades, 1982; Blankinship et al., 2016). The silt and clay particles are

228 connected with the formation of microaggregates, while sand particles are mostly associated with macro-
229 aggregates (Blagodatskaya and Kuzyakov, 2008). The stability of aggregates is an important factor
230 influencing C protection since it will directly influence the aggregation and disaggregation processes in
231 soils. However, aggregation is a dynamic process, with aggregates continuously forming and being
232 destroyed by natural cycles and animal or anthropogenic disturbance (Eyles et al. 2015). Aggregate size is
233 another important characteristic influencing C protection: Jastrow (2006) states that C turnover is higher
234 in macroaggregates (>212 μm) compared to microaggregates (53-212 μm), implying that microaggregates
235 have a higher C protection potential.

236

237 *1.1.10. Organomineral interactions with fine silt and clay minerals and soil carbon* 238 *saturation*

239 The other main mechanism for C protection is organomineral interactions with cations in soil that decrease
240 the soil C lability (Eyles et al., 2015; O'Brien and Jastrow, 2013). This process can happen inside aggregates
241 or in loose soil and relies on chemical sorption on mineral surfaces, polyvalent cation bridging and layered
242 chemical binding on mineral surface, of microbiological products primarily adsorbed on minerals and
243 covered by exudates (Cotrufo et al., 2015; Kleber et al. 2007; O'Brien and Jastrow, 2013). The soil potential
244 for organomineral interactions relies on the amount of fine elements in the soil (especially clay particles),
245 cations of different reactive elements, in particular Fe and Al (Swanston et al. 2009), and, particularly
246 interesting for this study, C saturation level of the soil. The concept of soil C saturation has been highlighted
247 after some studies reported no increase of soil organic C in soils even after further increase of C input (e.g.
248 Campbell et al. 1991, Solberg et al. 1997, Gill et al. 2002). To explain this behavior of soil, Six et al. (2002)
249 introduced the concept of 'soil C saturation', where it was suggested that the different C pools have
250 different saturation points after which they cannot effectively store C anymore. The capacity of these pools
251 to store C depends on their nature. For example, the physically protected C pool relies on the surface area

252 of particles, meaning that after the available areas are occupied by adsorbed C and further C input will not
253 be associated anymore and therefore not protected (Six et al. 2002). This concept was further elaborated
254 by Stewart et al. (2007), who stated that the soil C pool can be saturated with respect to the C inputs and
255 that a linear model cannot efficiently describe the input-storage behavior of a soil (Fig. 4). In this respect,
256 Stewart et al (2007) conclude that a soil poor in C, can store C more efficiently than a soil rich in organic C
257 and, therefore, closer to its C saturation threshold (Fig. 4), depending on the content of clay/silt of the
258 soils, the aggregation capacity, and their adsorption capacity. Several studies suggested that subsoil might
259 protect C more efficiently in fine soil fractions due to lower C saturation that increase the possibility for
260 organomineral interactions (Rasse et al., 2005; Lorenz and Lal 2005; Thomas et al. 2007; Horrocks et al.
261 2010;Rumpel et al., 2012). However, to our knowledge, no studies focused on the potential of subsoil
262 revegetation and the influence on the C-cycle and organomineral interactions.

263

264 *1.1.11. Soil carbon pools associated to different soil fractions*

265 When analyzing C content in soil, it is difficult to assess the different pools of C present in the soil and their
266 protection (biological protection determined by recalcitrance or physical protection from aggregate
267 occlusion or organomineral interactions). A method commonly used to assess C protection in soil is to
268 fractionate the soil and analyze the C in each fraction (Fig. 5). These soil C pools rely on different
269 protection mechanisms, and the degree of stability increases with decreasing fraction size. These pools
270 are defined as: i) coarse particulate organic matter (coarse POM, soil fraction > 200 μ m) (Fig. 5a), that is
271 free in the soil at different levels of degradation ii) fine POM (soil fraction 50-200 μ m) (Fig. 5b), that
272 comprises organic C occluded in soil aggregates. These two pools are mostly derived from the
273 decomposition of roots and shoots (Kögel-Knabner, 2002) and their C protection from microbial
274 consumption relies mainly on the recalcitrance of their lignocellulose C structures (Six *et al.*, 2002). Finally,

275 iii) C protected in the coarse silt (20-50 μm) (Fig. 5c) and iv) in fine silt+clay pools (<20 μm) (Fig. 5d). C
276 protected in these pools is mostly derived from labile rhizospheric and microbial compounds (Cotrufo *et*
277 *al.*, 2013, Vidal *et al.*, 2018). C in these pools is highly degraded via decomposition and mineralized by
278 microbial metabolism, and it is protected from microbial consumption via occlusion in microaggregates
279 and through organo-mineral complexation with clay particles and metals.

280

281 *1.1.12. Short – term changes in soil organic carbon mineralization due to vegetation: the*
282 *priming effect*

283 Although the positive effect of revegetating soils in terms of C input and soil C accumulation potential is
284 well established, the influence of plants on the C-cycle can also have negative impacts on soil C
285 sequestration. As already mentioned, the potential of a soil system in respect to C sequestration is
286 determined by the balance between input of photosynthetically absorbed C in soil and output of CO₂ via
287 soil heterotrophic respiration (Smith *et al.*, 2000; De Deyn *et al.*, 2008). The soil heterotrophic respiration
288 is determined mainly by microbial communities and their activity, and their consumption of C in soil (Jones
289 *et al.*, 2009; Kuzyakov and Larionova, 2006). One of the effect of plant C input is to influence the microbial
290 communities structure and activity and the consequent consumption of pre-existent soil C, that is
291 commonly known as ‘priming effect’ (Broadbent and Nakashima, 1974; Sørensen, 1974; Wu *et al.*, 1993;
292 Kuzyakov *et al.*, 2000). The priming effect is defined as strong short – term changes in C mineralization due
293 to vegetation (Kuzyakov *et al.*, 2000). We talk about ‘positive priming effect’ when the input of labile C
294 increases the activity of microbial communities and the mineralization of pre-existent C in soil (Fontaine
295 *et al.*, 2003). The positive priming effect has an adverse effect on soil C storage. However, if microbial
296 communities in soil switch from consuming pre-existent C to mineralizing fresh C input, the mineralization
297 of soil C will decrease (Kuzyakov *et al.*, 2000). In this case we talk about a ‘negative priming effect’,
298 beneficial to C storage in soil. The magnitude of the priming effect and its direction (positive or negative)

299 results from a complicated series of interactions between soil, plants and microbial communities (Cheng
300 and Kuzyakov, 2005). The first mechanism is known as the 'competition hypothesis' (Jackson et al., 1989;
301 Schimel et al., 1989; Kaye and Hart, 1997; Hodge et al., 2000; Cheng and Kuzyakov, 2005) and postulates
302 that competition for mineral N will determine the direction of priming. If the soil is poor in N, then the
303 priming effect is negative due to competition between plants and microbes. In the long run, plants have a
304 higher efficiency for N mining, and they will reduce the nutrient sources for microbial communities,
305 decreasing their C consumption (Cheng and Kuzyakov, 2005). Instead, when mineral nutrients are not
306 limiting and there is no competition between plants and microbes, rhizodeposition will increase microbial
307 activity resulting in increased soil C consumption and a positive priming effect (Cheng and Kuzyakov, 2005).
308 These mechanisms hold when microbial communities need to mine C for nutrients and energy, and are
309 usually observed in studies involving poor soils (pine forests and dry grasslands) (Ehrenfeld et al., 1997;
310 Schimel et al., 1989; Cheng and Kuzyakov, 2005).

311 When mineral nutrients are not limiting and the input of labile C is high, the priming effect might be
312 controlled by the preference of microbes for labile root derived C compared to nutrient rich soil C (Cheng
313 and Kuzyakov, 2005). If no nutrient limitation is present, microbes will prefer labile derived C as an
314 abundant and ready available source of energy (Cheng, 1999; Cheng and Kuzyakov, 2005). In this case, a
315 switch of substrate utilized will decrease the C consumption and result in a negative priming effect,
316 favouring soil C storage (Cheng and Kuzyakov, 2005, De Graaf et al. 2010). These effects are regulated by
317 microbial metabolism (Cheng and Coleman, 1990). Increased microbial biomass is linked with positive
318 priming, while negative priming is usually correlated to decreased microbial biomass (Cheng and Coleman,
319 1990; Reid and Goss, 1982; 1983; Sallih and Bottner, 1988). However, De Graaf et al. (2010) showed how
320 different levels of labile C input can influence microbial dynamics and consequent priming effect. Low
321 input of labile C ($\cong 0.7 \text{ mgC g}^{-1} \text{ soil}$) will increase microbial activity and soil C mining, resulting in a positive

322 priming effect. Instead, high labile C input ($> 7.2 \text{ mgC g}^{-1}$) increases microbial biomass but induce microbes
323 to switch preference of substrate consumption, from old C present in the soil to the fresh C inputted from
324 plants substrate utilization switch, decreasing old C consumption compared to unvegetated soil and
325 resulting in a negative priming effect (De Graaf et al., 2010).

326

327 *1.1.13. Possible impacts of revegetating geotechnical soils on the priming effect*

328 In geotechnical works, soils are often heavily managed and revegetated. Environmental conditions are
329 perturbed and it is not uncommon that subsoil is excavated, brought to the surface and revegetated.
330 Subsoils have a high C stability given by i) low microbial biomass (Taylor et al. 2002; Andersen and Domsche
331 1989; Ekklund et al. 2001) and activity (Fang and Moncrieff 2005), ii) oxygen limitation (Rumpel and Kögel-
332 Knabner, 2010), iii) energy limitation due to reduced labile C inputs (Fontaine et al. 2007) and iv) spatial
333 heterogeneity of organic C in subsoil and consequent separation from microbes (Von Lützow et al. 2006;
334 Holden and Fierer 2005). Fontaine et al. (2007) showed how a supply of fresh C in deep soil can decrease
335 the stability of pre-existent old C and increase positive priming. However, to our knowledge, no in vivo
336 experiment has been implemented on this topic, and, more importantly, no studies are available on the
337 effects of excavating and revegetating subsoil on the priming effect. Excavating, crushing, mixing, and
338 revegetating soil will have a major impact on the factors determining the stability of C in subsoil, and
339 possibly a high priming effect.

340

341 **1.2. GENERAL KNOWLEDGE GAPS**

342 As stated above, soil embankments represent an interesting structure for C sequestration due to two
343 features: 1) plants can be chosen to vegetate the embankments, and therefore the C input in the system,
344 and 2) soil can be managed and chosen to optimize C sequestration. Embankments are constituted of a

345 core of compacted soil, usually excavated from a depth of >1m and with a high percentage of clay, and
346 they can be covered by a layer of stripped topsoil to be revegetated. The choice of revegetating organic
347 topsoil (down to 30 cm depth) stripped and used to cover the embankment, or directly on an uncompacted
348 surface layer of mineral subsoil (>1 m depth), will deeply influence the soil C storage potential of the
349 geotechnical structure. However, no studies have been developed in depth on the effects of revegetating
350 subsoil brought to the surface on C storage, and their potential as C sink. There is a need of comparing C
351 storage potential of different plants and soils to design the most efficient C storage system in geotechnical
352 soils, a potential that have been hypothesized before but never adressed (Dejong et al., 2011).

353 The study of two soils showing diverse characteristics (fertility, microbial communities, C saturation levels),
354 and the use of plant species that have contrasting root traits connected with higher recalcitrance or lability,
355 allows to tackle fundamental knowledge gaps regarding the actors and mechanisms driving C
356 sequestration in soil. The next paragraphs give an overview of the knowledge gaps addressed in each
357 chapter of the thesis.

358

359 *1.2.1. Plant carbon input: influence of root traits and carbon accumulation in different soil*
360 *C pools*

361 Rhizosphere is considered as the main pathway for C to enter the soil, however few studies have tackled
362 the relationships between root traits and C storage. The studies that have indeed explored the effect of
363 the root economics spectrum on C storage (e.g. De Deyn et al., 2008; Bardgett et al., 2014; Poirer et al.,
364 2018; Henneron et al., 2019) considered the C storage in bulk soil, without exploring the effect of root
365 traits on C quality, i.e. the accumulation of C in different pools. Moreover, among the different explored
366 root traits, the root elongation rate has never been studied in relationship to C storage. We state that root
367 economics spectrum is lacking an important trait, since changes in root elongation rate affect the

368 production and the spatial distribution of root exudates, the main precursor of C stored in SILT+CLAY pool
369 (Cotrufo et al., 2013; Holz *et al.*, 2018).

370 *1.2.2. Subsoil brought to the surface: effect on C fluxes and actors involved in C-cycle*
371 Soil C stock within a defined time frame is the balance between input and transformation of newly
372 photosynthesized C from plants to soil (new C) and losses of existing soil organic C (old C) (Kuzyakov and
373 Domansky, 2000; Fontaine et al., 2004). Moreover, the balance between new C and old C is far from being
374 the whole story, as increasing studies have highlighted the equal importance of quality of soil C, as C stored
375 in different C pools (Cardinael et al., 2015). To our best knowledge, no study has ever bridged the link
376 between C pools and the fates of new C and old C. Besides the exploration of the fates of soil new C and
377 old C, as well as their associations with C pool, another significant knowledge gap comes to the
378 predictability of the fates of soil new C and old C using plant and soil features. More specifically, no studies
379 investigated the effect of root traits, microbial communities and soil characteristics (with an eye of
380 attention to the C saturation theory) on new input and old c changes in different C pools.

381
1.2.3. Subsoil brought to the surface: what is the effect of revegetation on the priming effect
382 Revegetating subsoil could have a high impact on pre-existent old C stability and the priming effect. C in
383 subsoil is highly stable, and perturbation of the environmental conditions could deeply influence the
384 stability and protection of this pool. Studies on priming of subsoil have been conducted (Fierer et al., 2003;
385 Fontaine et al., 2007; Wang et al. 2014), however no studies investigated the effect of bringing subsoil to
386 the surface. Understanding the priming effect at soil fraction level may also bring us new insight on the
387 vulnerability of soil C pools to fresh C input.

390

391 **1.3.STRUCTURE OF THE THESIS: OBJECTIVES AND HYPOTHESES**

392 Figure 6 shows the different research questions tackled in the research and discussed in each chapter, plus
393 their link with the main factors and processes discussed in each chapter. In this theses I and the research
394 team collaborating in this project aim to tackle the following general objectives:

395 i - Understanding the effect of plant and soil features on soil C sequestration in terms of quantity and
396 quality (fundamental objective)

397 ii - Identifying possible plant and soil practices that can be implemented to increase soil C storage in
398 embankments and, possibly, in grey soils from geotechnical work (applied objective)

399 The above two objectives regarding the fundamental mechanisms of C-cycle will be tackled in every
400 chapter of the thesis.

401

402 ***1.3.1. Chapter II: Pathway to persistence: plant root traits alter C accumulation in different***
403 ***soil carbon pools through microbial mediation***

404 i - Objective 1: Understand what are the relationships between root traits and C accumulation in
405 different soil C pools for 12 different herbaceous species commonly used in embankment
406 revegetation (Fig. 7).

407 Hypothesis 1: We hypothesize that traits related to labile C input (root elongation rate, hemicellulose
408 content, root biomass) promote C accumulation in the protected coarse silt and fine silt + clay C pools,
409 since these traits are expected to favor rhizodeposition and microbial activity, whereas root traits related
410 to recalcitrance (high lignin and cellulose content, high C:N ratio) promote C accumulation in the
411 unprotected coarse POM pool.

412 ii - Objective 2: What is the effect of species selection on the C sequestration in different soil C pools

413 Hypothesis. 2: We hypothesize that N₂-fixing species favor C accumulation in the protected fine silt+clay
414 pools since they have traits more related to labile C input, while non N₂-fixing species will favor C
415 accumulation in the POM fraction.

416

417 *1.3.2. Chapter III: The fates of fresh new carbon and old soil carbon differ in topsoil and*
418 *newly exposed subsoil and are explained by root, microbial, and soil particle size*

419 i - Objective 1: Quantify the fluxes of new C and old C in different soil pools;

420 Hypothesis 1: We hypothesize that soil particle size fractions associated C pools can regulate the fates of
421 old C and new C in the C sequestration process;

422 ii - Objective 2: Examine the pattern of covariation between new C input and changes of old C in
423 different C pools

424 Hypothesis 2: The fate of new C and old C will show independent patterns

425 iii - Objective 3: Investigate if the different actors involved in C storage, and the influence that plant
426 and soil have on them, can explain the patterns of new C and old C fluxes in different soil C pools

427 Hypothesis 3: We hypothesize that plant traits related to chemical composition and recalcitrance will be
428 driving POM accumulation in new C and consumption in old C, while traits related with high C input will
429 drive storage in protected fractions via microbiological consumption and deposition. We expect aggregate
430 stability to be positively correlated with new C and old C accumulation in fine POM and coarse silt fractions
431 due to physical protection of aggregates. We expect that soil N content positively correlates with new C
432 input. Fine fraction in soil is believed to be positively correlated with the new C storage in fine silt+clay
433 fraction due to organomineral interactions, and new C storage in fine silt+clay is expected higher in subsoil
434 than in topsoil due to lower soil C saturation levels. Finally we expect microbial activity, diversity and
435 abundance to be strongly linked with the amount of new C deposited in the protected coarse silt and

436 silt+clay fractions, and with the consumption and transformation of old C in the unprotected coarse POM
437 and fine POM fractions due to mineralization from microbial communities.

438

439 *1.3.3. Chapter IV: Soil quality drives the priming effect and plant species refine it*

440 i - Objective 1: Quantify the changes in C and the input of new C in soil to determine the losses of old C
441 in revegetated topsoil and subsoil brought to the surface (Fig. 8) and the priming effect of revegetating
442 with N₂-fixing (*Medicago sativa*) and a non N₂-fixing species (*Lolium perenne*) species (Fig. 9).

443 Hypothesis 1: Our hypothesis is that topsoil will have higher losses of old C due to higher microbial biomass
444 and activity. However, due to the higher protection of old C in subsoil and the changes in environmental
445 conditions given by revegetation, we hypothesize that subsoil will have higher old C losses compared to
446 bare soil, meaning a higher positive priming effect compared to topsoil.

447 ii - Objective 2: Quantify the priming effect in different C pools related to granulometric soil fractions.

448 Hypothesis 2: Given the higher protection of C in the finer soil fraction (silt and silt +clay fractions) we
449 hypothesis that the priming will occur in the unprotected particulate organic matter fractions (POM and
450 finePOM).

451 iii - Objective 3: Study the evolution over time of the sources of respired C in the system (represented
452 by the abundance of ¹³C) and its correlations with old C losses, new C input and priming.

453 Hypothesis 3: We hypothesize that the source of respiration in the system will switch more towards
454 labelled plant inputs over time, along with plant development. We believe new C input to be positively
455 correlated with the abundance of ¹³C in respired CO₂ (A¹³C). However, we expect different behaviours in
456 the two soils regarding the old C losses. In topsoil we suggest that A¹³C will be negatively correlated with
457 old C losses, due to switch in microbiological substrate preference, while in subsoil A¹³C will be positively

458 correlated with old C losses, due to increased microbial activity. In the same way, priming will be negatively
459 correlated to A¹³C in topsoil, while being positively correlated in subsoil.

460

461 *1.3.4. Chapter V: general discussion, guidelines and prospective for carbon storage in*
462 *geotechnical embankments*

463 In Chapter V I intend to delineate a more comprehensive view on the effect of soil and plant selection on
464 C storage in embankments based on the results of this study. I want to discuss the potential benefits of
465 embankments for C storage and propose guidelines for embankments revegetation, more specifically: i)
466 possible management options to increase C storage in these geotechnical soils and ii) perspectives for
467 future studies on C sequestration.

468

469 *1.3.5. Annex I: Perspectives: the influence of vegetation on soil microstructure and its*
470 *implications on soil carbon sequestration: a geotechnical approach*

471 Annex one is an overview of an ongoing research with UNICAS regarding soil structure. More specifically
472 we investigate the influence of vegetation on soil microstructure and its implications on soil C storage and
473 protection. I propose a multidisciplinary approach including geotechnical engineering and soil
474 science/ecological methods to investigate soil structure in terms of i) soil porosity and void ratio, ii)
475 aggregate stability and C protection, and iii) new C input in different aggregate classes. These results will
476 allow a more comprehensive view on aggregate formation and C protection in revegetated topsoil and
477 subsoil brought to the surface, and understand the role of porosity and void ratio in relation to C
478 protection. Research questions, methodology and preliminary results are outlined in Annex I.

479

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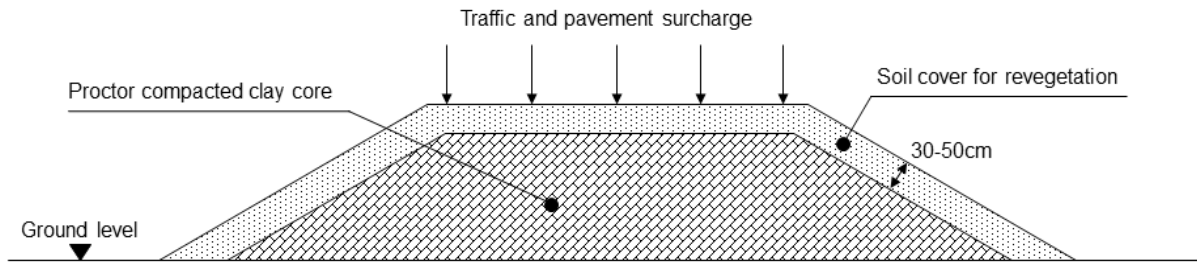
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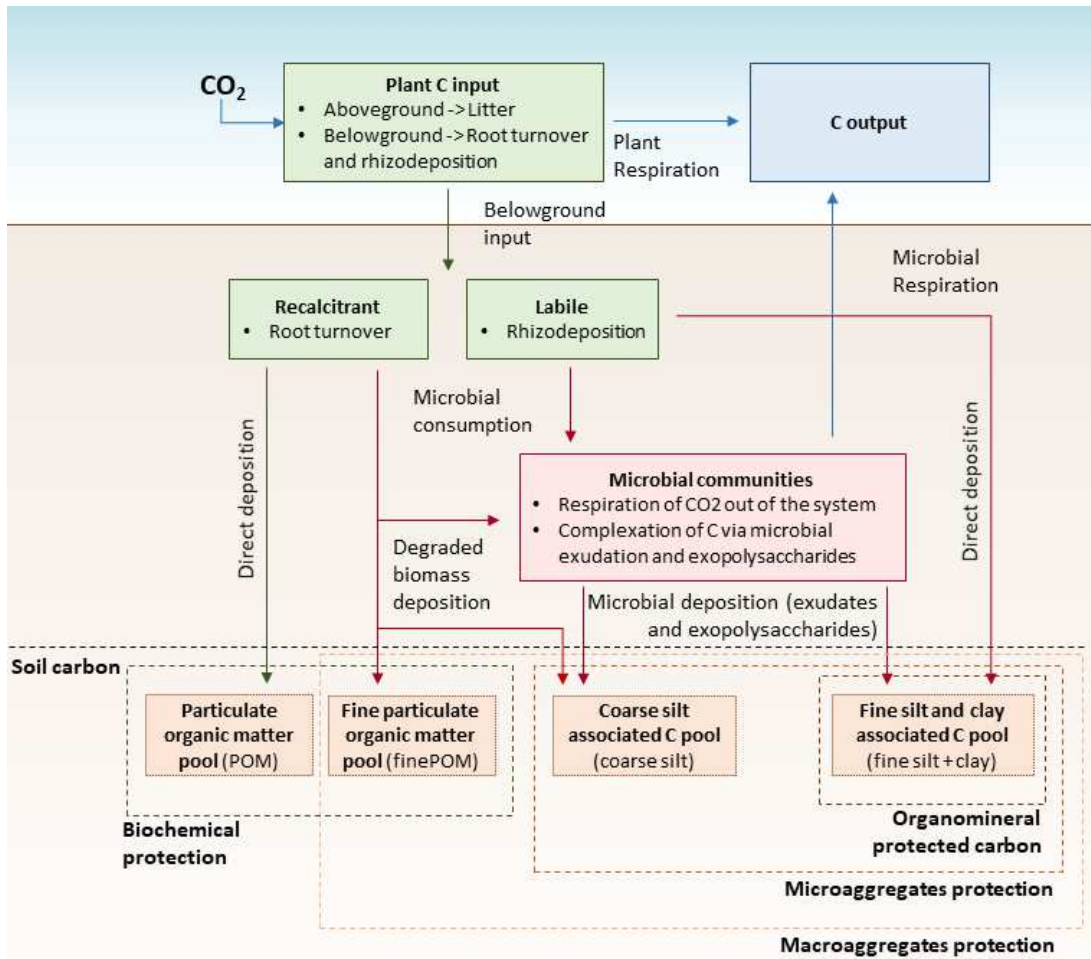
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715 FIGURES AND TABLES

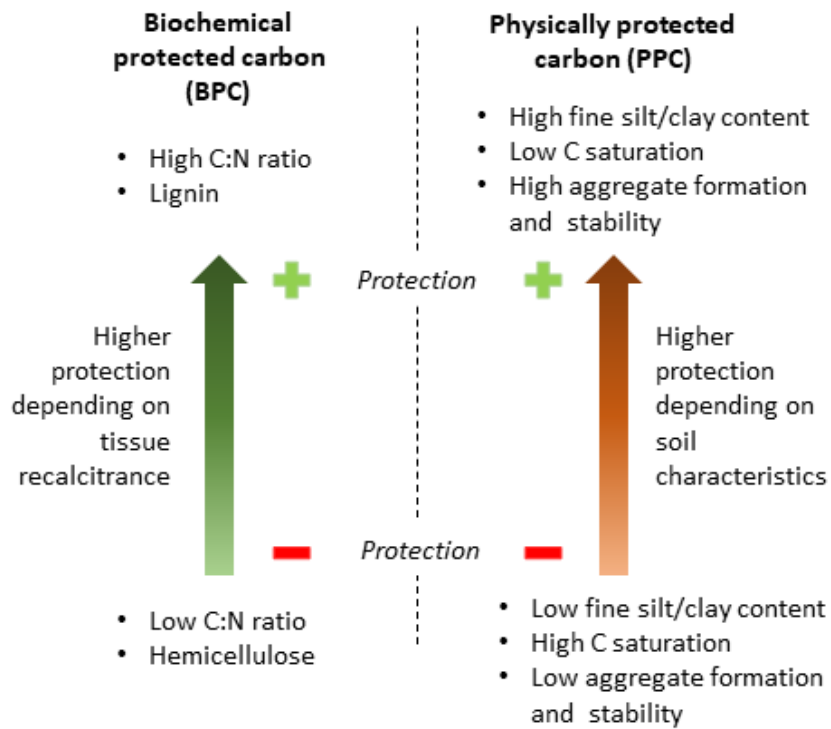


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717 **Figure 1:** Section of a geotechnical embankments comprehensive of proctor compacted clay core and cover of soil for
718 revegetation purposes

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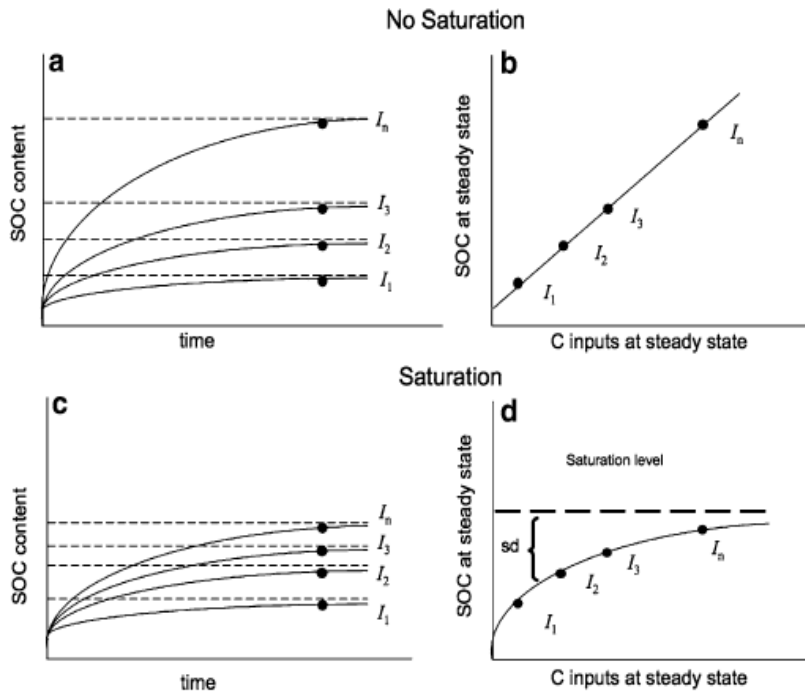
720
721 **Figure 2:** Flow chart of soil carbon (C) cycle and deposition/complexation in different soil fractions. In green square
722 boxes, C input from plants is depicted, and green arrow signifies deposition of direct plant inputs. The red box
723 symbolizes the microbial communities and the red arrows the consumption and deposition of C by microbial
724 communities. Brown boxes represent the different soil C pools. The green dashed line represents the biochemical
725 protection of free particulate organic matter (POM) in soil, while the brown dotted lines the soil protection via aggregate
726 complexation and organomineral absorption on fine silt and clay minerals.



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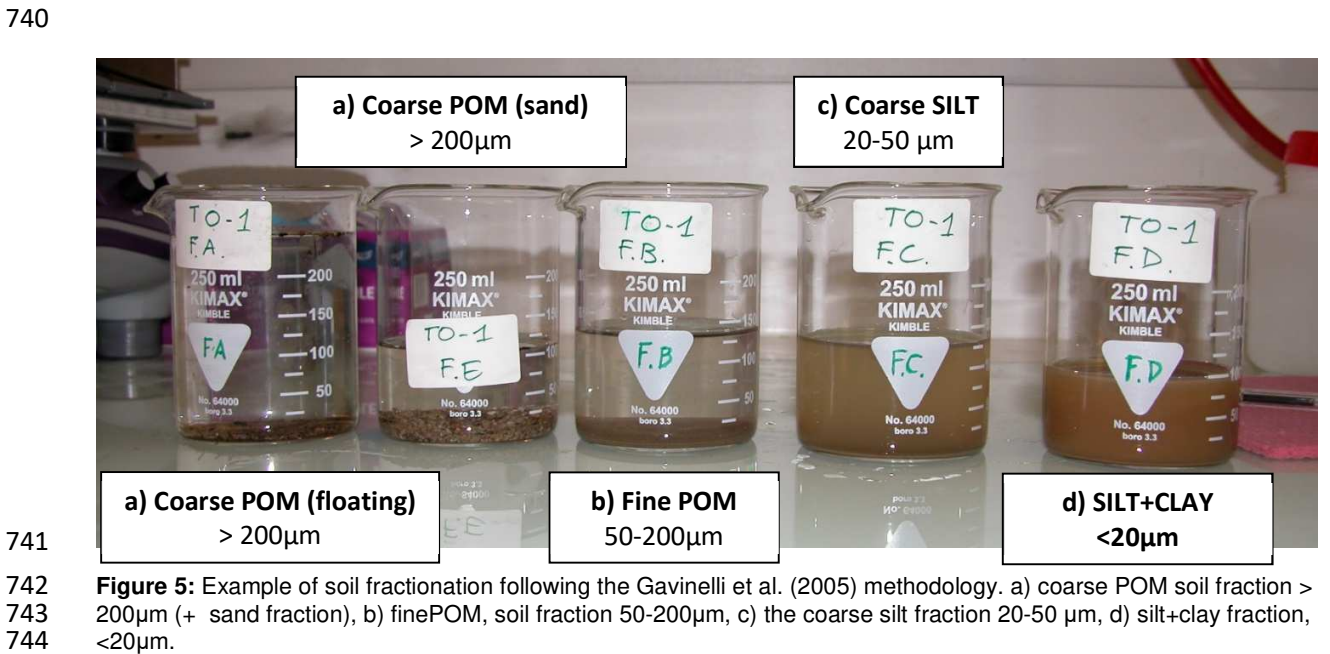
728 **Figure 3:** Representation of different types of carbon (C) protection in soil. Biochemically protected carbon (BPC, left)
 729 protection depends on the chemical composition of plant tissues, with recalcitrant C having a lower turnover. Physically
 730 protected carbon (PPC) depends on soil properties, more specifically on the aggregate formation and the resulting
 731 stability, the fine soil fraction in soil and soil C saturation levels.

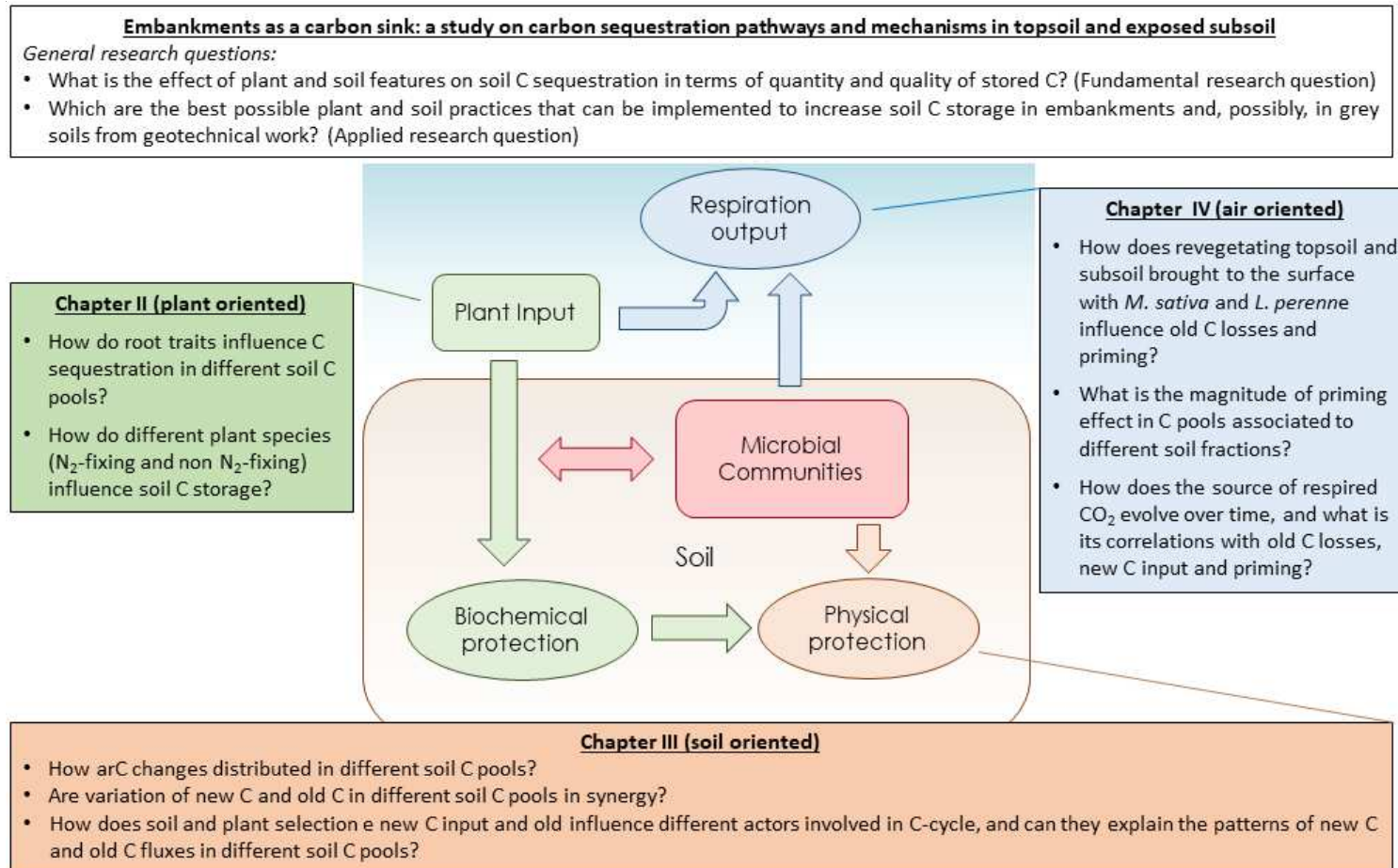
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734 **Figure 4 :** Different soil organic C (SOC) evolutions with a constant carbon (C) input for two conditions: a,b, not
 735 considering the effect of C saturation, and c, d including the C saturation effect. Under unsaturated conditions, (a)
 736 a steady state soil C accumulation over time will express a linear relationship if expressed (b) over C input. If the
 737 relationship between SOC and time is analyzed for (c) a C saturated soil it will not be proportional, meaning that a C
 738 input increase will not result in a linear SOC accumulation over time, but (d) in a asymptotic relationship (after Stewart
 739 et al. 2007).





744

Figure 6: scheme of thesis structure and related research questions (RQ). Applied RQ are presented in the first box, together with the title of the thesis. Fundamental RQ are displayed in the different boxes related to the different chapters of the thesis. The scheme in the middle represents a simplified version of Figure 1. The squared boxes represent the main actors in C-cycle (green plants, red microbes, brown soil) while the circles the pools of C: soil carbon (biochemically and physically protected) and the atmospheric C in CO₂.

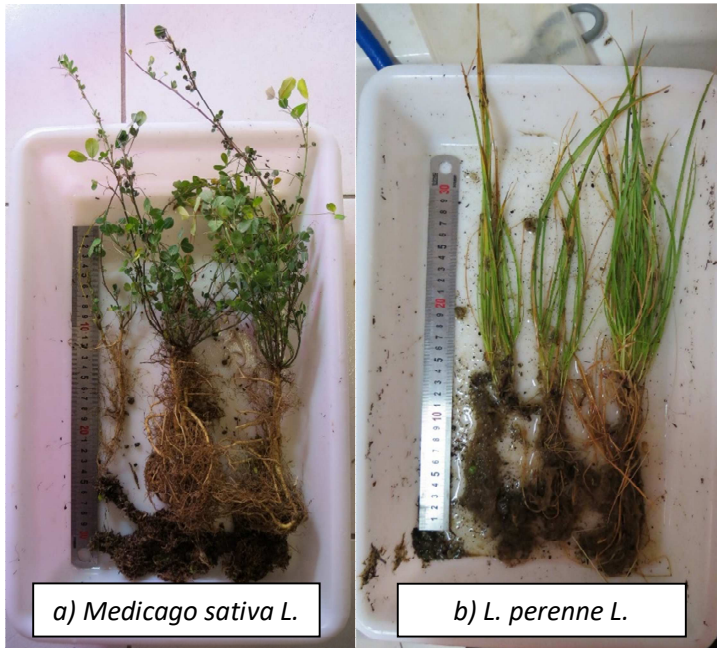


745

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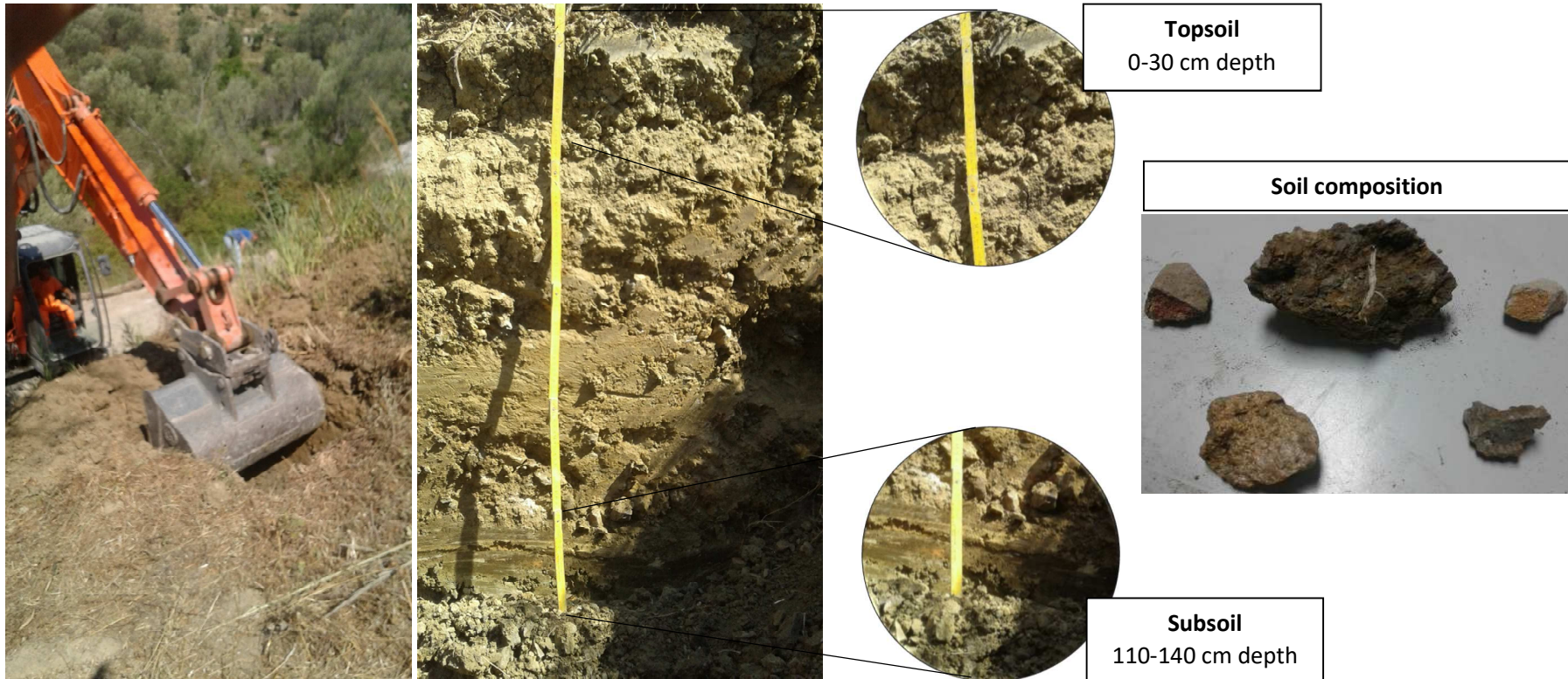
747

Figure 7: Experimental set up with 12 species grown in monoculture in grow-boxes. The picture shows two of the three blocks of growboxes present in the experiment (Chapter II)



748

749 **Figure 8:** Species grown in ^{13}C constant labelling experiment sampled after 6 months for root traits assessment
750 (Chapter III and IV).



751

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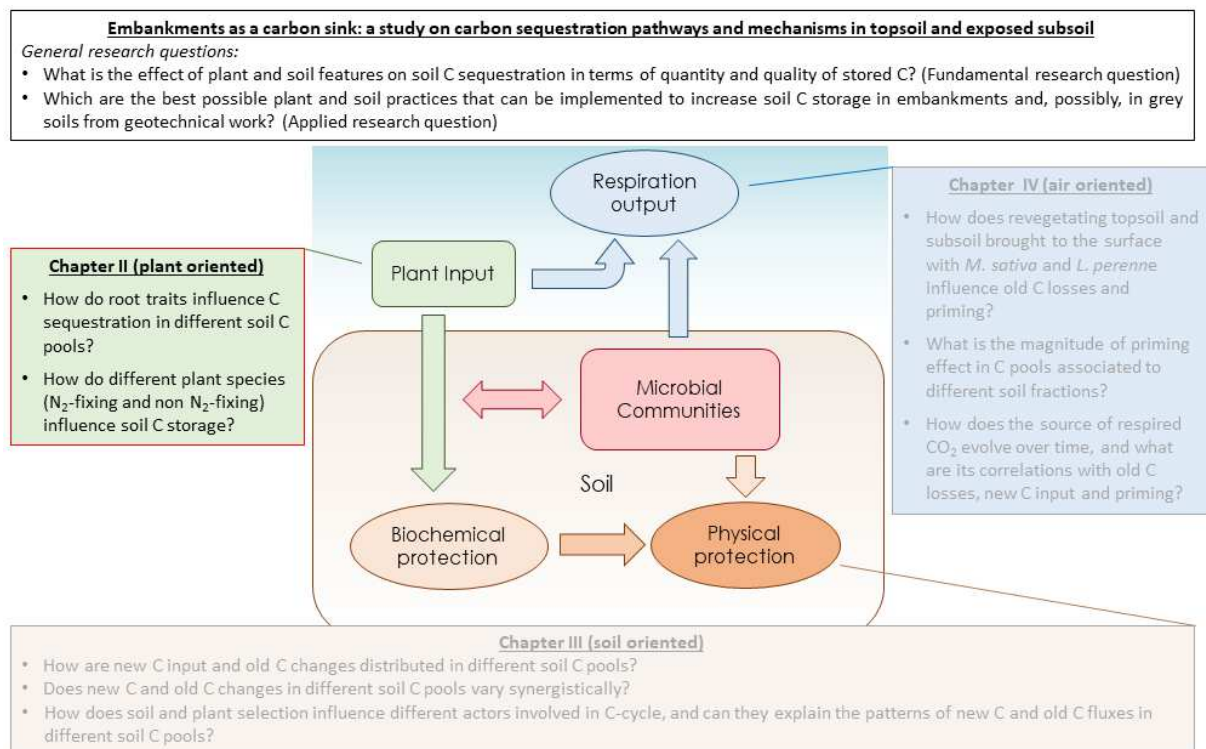
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Figure 9: Excavation in Pisciotta (SA), Italy, to collect the soil for the experiment described in Chapter III and IV. Topsoil excavated from 0-30cm depth, subsoil from 110-140 cm depth. On the far right a picture with the main components of soil structure visually isolated.

CHAPTER II: Pathway to persistence: plant root traits alter carbon accumulation in different soil carbon pools through microbial mediation



Plant input is the first step in soil C sequestration. Plant choice influences the final C storage in soil by providing different amounts and quality of C input. In this chapter, we aim to quantify this effect by assessing the C changes in different soil C pools associated with different soil size particle fractions, and relating them to contrasting root traits characterizing 12 different herbaceous species used for embankment revegetation in south of France.

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15 **ABSTRACT**

16 The persistence of soil organic carbon (C) depends primarily on its origin. Mineral-associated C derived
17 from microbial compounds and the rhizosphere of elongating plant roots is better protected and persists
18 longer in soil compared to unprotected particulate organic matter (POM). Originating from decomposing
19 root and shoot litter, POM is fast-cycling and persists mainly because of its high level of recalcitrance.
20 Theory suggests that plant root traits and growth dynamics should affect C accumulation into different
21 pools, but the specific traits that drive this accumulation have not been identified.

22 We measured root chemical traits and elongation rate (RER) in 12 herbaceous species, of which five
23 comprised the dinitrogen (N₂) fixing trait. Along with microbial activity, C was measured in four pools
24 (fractions), based on soil structure.

25 N₂-fixing species had fast-growing, hemicellulose-rich roots with low lignin and C:N, that increased C input
26 in the silt fraction, mediated by high microbial activity. Non N₂-fixing species possessed lignin-rich roots
27 with high C:N that grew slowly and with input of C in the POM fraction.

28 Our results demonstrate that specific root traits determine C input into different soil pools, mediated
29 primarily by microbial activity, thus determining the fate of soil organic C. We also highlight that C in
30 different soil pools, and not total soil organic C, should be measured in future studies.

31 **Keywords:** *protected and unprotected soil organic carbon, physical soil fractionation, root elongation rate,*
32 *rhizodeposition, recalcitrant, labile carbon input*

2.1. INTRODUCTION

33
34 In the current context of climate change and several international initiatives (e.g. COP21, 4p1000,
35 Stolbovoy *et al.*, 2017), understanding the interactions between vegetation and soil has become primordial
36 for sequestering atmospheric carbon (C) into anthropogenically disturbed soil, e.g., agricultural fields,
37 mining waste soil, road embankments and technosols (Paustian *et al.*, 2016; Griscom *et al.*, 2017; Minasny
38 *et al.*, 2017). Plants act as a major conduit for transferring C into soils via litterfall, root mortality and
39 exudation (Six *et al.*, 2004; Derrien *et al.*, 2016; Sokol *et al.*, 2019). Some C is transformed by soil microbes
40 and released back into the soil by respiration (Jones *et al.*, 2009; Kuzyakov and Larionova, 2005), but C
41 can also be stabilized in soil, increasing its residence time (Besnard *et al.*, 1996; Lal, 2004; Rasse *et al.*,
42 2005; Bardgett *et al.*, 2014 ; Vidal *et al.*, 2018; Sokol *et al.*, 2019). Carbon stabilization occurs through
43 three main mechanisms: selective preservation due to recalcitrance to decomposition, spatial
44 inaccessibility to decomposers due to occlusion in soil aggregates and complexation with minerals and
45 metal surfaces (Sollins *et al.*, 1996; von Lutzow *et al.*, 2006; Poirier *et al.* 2018). These mechanisms are
46 influenced by abiotic and biotic factors and especially by plant roots that contribute more than aerial parts
47 to C stabilization (Balesdent and Balabane, 1996; Rasse *et al.*, 2005). In a recent review, Poirier *et al.*, (2018)
48 have argued that the root traits that most influence C stabilization are those related to chemical
49 composition, root exudation and the presence of symbionts (mycorrhizas and nitrogen-fixing *Rhizobium*),
50 whereas the role of morphological traits is not yet clear. More specifically, root traits increasing
51 recalcitrance promote short-term C stabilization by slowing root decomposition rates, but traits that
52 reduce recalcitrance contribute to long-term C stabilization via the reaction of microbial products with
53 mineral surfaces. Although several studies have analyzed the link between plant functional traits, microbial
54 activity and C accumulation (Chapin 2003; Lavorel *et al.*, 2007; De Deyn *et al.*, 2008; Fujisaki *et al.* 2018;
55 Poirier *et al.* 2018), as yet, no study has focused on how root growth and specific traits can alter the

56 accumulation and potential persistence of different soil C pools, which are linked to the physical structure
57 of soil itself.

58

59 Through differences in physiological activity and chemical composition, specific root traits should affect C
60 accumulation into different pools based on soil structure. These pools are defined as: i) coarse particulate
61 organic matter (coarse POM, fraction > 200 μ m), that is free in the soil at different levels of degradation
62 and ii) fine POM (fraction 50-200 μ m), that comprises organic C occluded in soil aggregates and can be
63 occluded in macroaggregates. These two pools are mostly derived from the decomposition of roots and
64 shoots (Kögel-Knabner, 2002) and their C protection from microbial consumption relies mainly on the
65 recalcitrance of their lignocellulose C structures (Six *et al.*, 2002). Finally there is iii) C protected in the
66 coarse silt and fine silt+clay pools (fraction 20-50 μ m and <20 μ m, respectively), that is mostly derived from
67 labile rhizospheric and microbial compounds (Cotrufo *et al.*, 2013, Vidal *et al.*, 2018). Carbon in these soil
68 C pools is highly processed and protected from microbial consumption via occlusion in microaggregates
69 and through organo-mineral complexation with clay particles and metals. It is now generally accepted that
70 these low molecular weight, chemically labile carbon forms, persist in soil longer than chemically
71 recalcitrant C structures, when protected by organo-mineral complexation (Mikutta *et al.*, 2006; King *et*
72 *al.* 2019; Robertson *et al.*, 2019; Sokol *et al.*, 2019). The stability of sequestered C in soil is therefore linked
73 with the fraction of soil to which it is associated, with a higher stability of C pools associated with finer
74 fractions due to physical protection in microaggregates and via organo-mineral complexes.

75

76 Carbon accumulation into the coarse POM pool is related to the amount of recalcitrant matter present
77 and so should be higher in soils containing roots with high cellulose, lignin and carbon:nitrogen (C:N)

78 content (Poirier *et al.*, 2018). To increase long-term C persistence in coarse silt and fine silt+clay fractions,
79 root traits related to physiological activity and rhizodeposition stimulating microbial activity should have
80 the greatest influence. Rhizodeposition is the release of organic compounds by living roots, mainly through
81 i) sloughing-off of root border cells, ii) secretion of mucilage, iii) root exudation and iv) the senescence of
82 root epidermis (Poirier *et al.*, 2018). Root exudation is the largest producer of C, releasing up to 100 times
83 more C than border cells and mucilage, and three orders of magnitude more C than senescence of the root
84 epidermis (Poirier *et al.*, 2018). Root exudates are mainly released at the root apex (Jones *et al.*, 2009;
85 Canarini *et al.* 2019). Changes in root elongation rate (RER) thus affect the production and the spatial
86 distribution of root exudates, and therefore C input into soil (Holz *et al.*, 2018). Root elongation rate is
87 affected principally by local abiotic soil conditions such as soil temperature, moisture, and compaction,
88 and can differ among species, although most data relate to woody species (Kaspar and Bland 1992; Kirby
89 and Bengough 2002; Steinaker *et al.*, 2011; Heinze *et al.*, 2017). As the link between root morphological
90 traits and C accumulation in soil is tenuous (Poirier *et al.*, 2018), it is of interest to study RER as a new and
91 more powerful predictor of C deposit in coarse silt and fine silt+clay C pools. RER is usually higher in roots
92 with a low diameter, high specific root length and nitrogen uptake rate and is a good predictor of plant
93 growth rate (Larson *et al.*; Funk; 2016). We ask, therefore, if root chemical and morphological traits and RER
94 of diverse plant species can be linked to C accumulation in different soil C pools, thereby altering the
95 persistence of C in soil.

96 Root exudates are an important substrate for microbial communities, Dennis *et al.* (2010), argue that their
97 influence on structuring microbial communities will be greatest around the elongating root apex and
98 emerging laterals. The C substrates that are assimilated by microorganisms at the root apex are utilized
99 rapidly for respiration and growth, or lost as microbial exudates or exopolysaccharides that are used as a
100 substrate for subsequent microbial communities. Certain microbial species, such as *Rhizobium*, present in

101 nodules of dinitrogen fixing (N_2 -fixing) species, produce large amounts of exopolysaccharides (Downie,
102 2010) that can also be adsorbed onto fine silt and clay particles (Fehrmann and Weaver, 1978). Greater
103 soil C contents have been reported in soil beneath N_2 -fixing species (Binkley, 2005; Fornara and Tilman,
104 2008), however whether N_2 -fixing species promote a higher C accumulation in the fine silt and clay soil
105 fractions, remains unknown. Also, N_2 -fixing species have roots that are easily degradable with high
106 hemicellulose (Hernández *et al.*, 2017) and low C:N ratios (Warembourg *et al.*, 2003; Roumet *et al.*, 2005),
107 therefore enhancing microbial activity (Poirer *et al.*, 2018).

108

109 We explored the effects of root traits on C accumulation into different soil C pools beneath 12 herbaceous
110 species that possessed diverse root traits in terms of morphology, chemical composition, and root
111 elongation rate. Of the 12 species, five belonged to N_2 -fixing Fabaceae, five to Poaceae, one to Rosaceae
112 and one to Plantaginaceae. Our main hypothesis is that soil C accumulation into different soil C pools is
113 driven by root traits that affect microbial activity. More specifically, we hypothesize that (i) traits related
114 to RER promote C accumulation in the coarse silt and fine silt + clay C pools since these traits are expected
115 to favor rhizodeposition and microbial activity, whereas (ii) root traits related to recalcitrance (high lignin
116 and cellulose content, high C:N ratio) promote C accumulation in the unprotected coarse POM pool, (iii)
117 N_2 -fixing species favor C accumulation in the fine silt and clay pools. Results should enable us to
118 disentangle the relationships between root growth, traits and the accumulation and stabilization of C in
119 different soil C pools, between N_2 - and non N_2 -fixing species.

120

121 2.2. MATERIALS AND METHODS

122 2.2.1. *Experimental setup*

123 The experiment was set up in the experimental garden of CEFE-CNRS Montpellier, France (43.6389° N°,
124 3.864125° E and lasted 37 weeks (from t0: Sept-2016 to t36: July-2017). Twelve herbaceous species were
125 grown as monocultures in steel boxes (0.7 m length x 0.7 m width x 0.3 m depth): five N₂-fixing species
126 from the Fabaceae family and seven non N₂-fixing species, including five Poaceae, one Plantaginaceae and
127 one Rosaceae species (Table 1).

128 Seventy-eight boxes were prepared: six replicate boxes per species and six additional boxes of bare soil
129 used as controls. Boxes were organized in three blocks with two rows of 13 boxes in each block; each row
130 comprised 12 monocultures (one per species) and a bare soil randomly arranged in each row (see support
131 information, Fig. S1). Boxes of the first row were used for destructive plant and soil sampling, while the
132 boxes of the second row were equipped with temperature and humidity sensors (Section 1.1, Fig. S1) and
133 with rhizotrons for the study of root elongation. These boxes were undisturbed for the duration of the
134 experiment. Rhizotrons consisted of a 0.2 m width x 0.3 m depth x 0.005 m thick pane of transparent PVC
135 set into the lower walls of the boxes, through which root elongation rate (RER, in mm root⁻¹ day⁻¹) and root
136 length production (RLP, in mm mm⁻² day⁻¹) were observed (Fig. S2).

137 Boxes were inclined at 20° relative to the horizon (Fig. S2) to encourage the positive geotropism of roots
138 when they come into contact with rhizotron windows (Huck and Taylor, 1982). Soil boxes were filled with
139 0.113 m³ of soil sieved to 8 mm and manually compacted, to a total of 190 kg of soil per box (bulk density
140 = 1.70 ± 0.02 g cm⁻³). The soil, excavated in Villefort (France; 44°26'25" N, 3°55'58" E), was a sandy-loam
141 (62.6 % sand, 26.1 % silt, 11.3 % clay); with 1.36 g kg⁻¹ of total N, 16.9 g kg⁻¹ of total C, 0.069 g kg⁻¹ of
142 phosphorus (P Olsen), pH in water was 7.06, and cation-exchange capacity (CEC) was 7.98 cmol_c kg⁻¹.

143 On 17-19 October 2016, 72 boxes were sown as monocultures (12 species x 6 replicates with one replicate
144 species per row). Seeds of each species were sown in lines, the distance between lines was 7.5 cm and the
145 distance between plants within a line was 7.5 cm leading to a final plant density of 155 plant m⁻². During
146 the experiment, mean air temperature was 13C° (Fig. S3) and the cumulative precipitation was 349 mm
147 (Fig. S3 for additional information on climatic conditions over the 37 weeks of the experiment). In addition,
148 each box was watered with sprinklers to provide suitable conditions for plant development and boxes
149 were carefully weeded by hand. More information on experimental setup can be found in Support
150 information, Methods S1.

151

152 2.2.2. Analysis of carbon content in different soil fractions

153 Soil carbon content was measured before filling the boxes, as a reference for time 0 (t₀), on three samples
154 from the initial homogenized soil batch, and at the end of the experiment, i.e. at 37 weeks (t₃₇) after
155 sowing. At t₃₇, soil samples were taken at 0-20 cm depth using a soil corer (7.5 cm in diameter) in each
156 box of rows installed for soil sampling in the three blocks. All soil samples were separated into two depths
157 (0-10 cm and 10-20 cm), air dried and separately sieved to 2 mm. A subsample of 40 g of soil was collected
158 at a depth of 0-10 cm for subsequent fractioning into POM fractions (coarse POM: 200-2000 μm and fine
159 POM: 50-200 μm), coarse silt (20-50 μm), fine silt + clay (<20 μm) fractions. Soil fractioning was carried
160 out using the method from Gavinelli *et al.*, (1995) (see also Support information, Methods S2). The dried
161 fractions were weighed to check that the sum of the fraction's weight did not differ from +/-5% the initial
162 40 g total weight. The quality of the soil particle dispersion was checked and did not differ +/-5% compared
163 to soil texture analysis, being 62.6% in the sand fraction and 37.4% in the fine silt + clay fraction.

164 Carbon content in each of the soil fractions (C_{POM} - carbon in the coarse 200-2000 μm fraction ; C_{finePOM} -
165 carbon in the fine 50-200 μm fraction ; C_{SILT} - carbon in the 20-50 μm coarse silt fraction and C_{SILT+CLAY} -

166 carbon in the fine silt+clay <20 μm fraction) was analyzed using an elemental analyzer (CHN model EA
167 1108; Carlo Erba Instruments, Milan, Italy) to represent different soil C pools. A subsample of 0.1 g was
168 taken from each 40 g sample and analyzed without fractioning to determine the total C in the bulk sample.
169 The difference between total C in bulk soil and the sum of C in the different soil fractions was used to
170 assess the correctness of the fractionation (+/-5%) and this was equal to 97.3%.

171 We calculated soil C changes (ΔC) in each soil fraction as the difference between C concentration in mg C
172 g^{-1} soil at 0 and 37 weeks ($\Delta\text{C} = \text{C}_{t37} - \text{C}_{t0}$). The sum of ΔC in each soil fraction ($\Delta\text{C}_{\text{SUM}} = \Delta\text{C}_{\text{POM}} + \Delta\text{C}_{\text{finePOM}} +$
173 $\Delta\text{C}_{\text{SILT}} + \Delta\text{C}_{\text{SILT+CLAY}}$) was also calculated to investigate the variation in the totality of the soil fractions. Note
174 that ΔC can be either positive (accumulation) or negative (depletion).

175

176 **2.2.3. Measurement of root elongation rate (RER) and root length production (RLP)**

177 As soon as the first root was visible in a rhizotron, the rhizotron of one box for each species was scanned
178 every two weeks using a smartphone scanner application CamScanner (INTSIG Information Co., Ltd,
179 Shanghai, China; version 3.9.5). It was not possible to scan every box due to the time consuming
180 methodology used to assess RER and RLP. A smartphone (Samsung Xcover3, Samsung Electronics, Korea)
181 was kept at a fixed distance from the rhizotron (30 cm) and a ruler was included in the picture to set the
182 scale (Mohamed *et al.*, 2017). Images were then analyzed with the SmartRoot software (Lobet *et al.*, 2011),
183 a freeware plugin of ImageJ software (Schneider *et al.*, 2012). The images acquired were converted into 8-
184 bit grey scale and, when necessary, color- inverted, so that roots were dark on a lighter background.
185 SmartRoot allows the semi-automatic tracing of roots by clicking on the basal point of each root (Fig. S4a).
186 Data extracted include the length and diameter of the roots. The resulting traced image of roots could
187 then be imported and superimposed onto a new image, allowing analysis of subsequent images and
188 creating a time-dependent dataset acquiring root length at different time steps.

189 Root elongation rate (RER; mm d^{-1}) is indicative of primary root growth and is defined as the difference in
190 individual root length measured between two dates. RER is a frequent but punctual observation of root
191 dynamics over time. As soil C storage is a cumulative process, root length production (RLP; m) after 37
192 weeks was also calculated. RLP is the total length of all roots produced in a specific period of time
193 (Mommer *et al.*, 2015), and we estimated RLP for up to 60 roots (randomly chosen) per rhizotron when
194 available. Of these 60, 30 were selected from the 'new roots', i.e., the roots that were not present in the
195 previous scan, and so had formed in the previous two weeks. Of the 30 'new roots', 20 were primary and
196 first order roots and ten were second order laterals (Fig. S4a, according to the developmental centrifugal
197 protocol of root topology, Berntson, 1997). Then, 30 'old roots' were selected at each subsequent sampling
198 date. The 'old roots' were the roots already present in the previous scan (again, 20 primary axes and ten
199 second order lateral roots). Fig. S5 shows an example of rhizotron analysis for new and old roots. To have
200 a more representative sample of the 'old roots', ten primary roots were selected from the 20 primary 'old
201 roots' of the previous scan, ten were selected from the 20 newly emerged roots of the previous scan, five
202 were chosen from the ten second order lateral 'old roots' of the previous scan and five were selected from
203 the newly emerged second order laterals of the previous scan. This method was used to select roots at
204 each subsequent sampling date. If one or more roots had: 1) reached the boundaries of the rhizotron, or
205 2) were in a bundle and not distinguishable (Fig. S4b), or 3) could not be analyzed for any other reason
206 (e.g. soil masking the root), they were discarded and different roots were then selected. This methodology
207 might have hindered the selection of fast growing roots that reach the rhizotron boundaries faster than
208 slower growing roots.

209 The mean daily RER was calculated by subtracting from the length of a root (L_{t2}) the length of the root
210 acquired at the previous sampling date (L_{t1}). This result was then standardized dividing by the number of
211 days between the two sampling dates to have the mean elongation rate of a single root:

212 $RER(t) = (L_{t2} - L_{t1})/[t2 - t1]$ [1]

213 Root length production (RLP) of roots over the 37 weeks was chosen as a cumulative indicator for root
214 dynamics, adapted from Mommer *et al.*, (2015):

215 $RLP = \sum_{t=1}^T \sum_{r=1}^R (RER_{r,t} * \frac{R_{30}}{R})$ [2]

216 Where t represents the different observation intervals of two weeks each ($t = 1, 2, \dots, T$); $T = 14$, i.e. $(37+1)/2$;
217 $RER_{r,t}$ is the RER of a root r at the specific time interval t ; R the real number of roots analyzed in that
218 interval. Since the number of analyzed roots varied depending on dates and species, we decided to
219 standardize the analysis of RLP for $R_{30} = 30$ roots.

220 To refine the understanding of root dynamics, the RER and RLP were calculated separately for the new
221 roots (RER_{NEW} and RLP_{NEW} , i.e. roots initiated during the 2 weeks interval between measurements), old
222 roots (RER_{OLD} and RLP_{OLD} , i.e. roots older than 2 weeks), and also the total root system, regardless of root
223 age. For all species, RER was high during the first two samplings after their initiation and then decreased
224 rapidly or stopped. Therefore, mean RER could be biased by the development of new roots, justifying our
225 decision to separate roots based on age and order for the statistical analysis.

226

227 **2.2.4. Analysis of root traits**

228 After 37 weeks, a soil core (7.5 cm diameter, 20 cm depth) centered on one individual plant per species
229 and per box was collected. In each core, roots were separated from the aboveground part and washed.
230 Roots were sorted into absorptive roots, typically the first, second and third root orders (defined as the
231 most distal root orders), and transport roots, that were higher order roots (all orders above third order
232 roots), (McCormack *et al.*, 2015). A subsample of absorptive roots (0.1g dry mass on average) was

233 selected, stained with a solution of methyl violet (0.5 g L⁻¹), spread into a transparent water filled tray and
234 scanned at 800 dpi (Epson Expression 1680, Canada).

235 The software Winrhizo Pro (Regent Instruments, Quebec, Canada) was used to determine the root
236 diameter (from 0 to 2 mm, with a 0.1 mm diameter interval) of absorptive roots. Roots were then oven
237 dried at 40°C for 3 days and weighed to determine the total root dry mass for each core.

238 For each species, determination of root chemical composition was conducted on three subsamples of
239 absorptive roots reserved for chemical analyses. C and N concentrations were determined on ground
240 material using an elemental analyser (CHN model EA 1108; Carlo Erba Instruments, Milan, Italy).
241 Concentrations of water-soluble compounds + hemicellulose, cellulose and lignin were obtained following
242 the Van Soest method (Van Soest, 1963, Roumet *et al.*, 2016) and using a fiber analyser (Fibersac 24;
243 Ankom, Macedon, NJ, USA).

244

245 *2.2.5. Soil microbial activity*

246 Substrate-induced respiration (SIR) was determined as a proxy of the soil microbial respiration potential,
247 according to Beare *et al.* (1990). Briefly, 20 g air-dried 2 mm sieved soil samples were incubated in 150 mL
248 sealed serum flasks with 1.5 mg C-glucose g⁻¹ soil, at 80% field capacity and at 25°C. A 200 µL aliquot of
249 the flask headspace was analyzed for CO₂ concentration after 2 and 6 hours using a microcatharometer
250 (MicroGC Serie S, SRA Industries, Marcy l'Etoile, France) equipped with a PoraPlot column (Agilent, Santa
251 Clara, United States). Substrate induced respiration rates were calculated as the mass of C-glucose
252 converted to C-CO₂ per g of soil dry weight and per hour (in µg C-CO₂ g⁻¹ soil h⁻¹).

253

2.2.6. *Statistical analysis*

254 The normal distribution of residues was verified using a Shapiro-Wilk test ($\alpha p = 0.05$). A one-way analysis
255 of variance (ANOVA) and post-hoc Tukey HSD tests were performed to test the effects of species on RER,
256 RLP, root traits and ΔC sequestration in soil C pools. To test the effect of N_2 -fixing and non N_2 -fixing species,
257 the same analyses were conducted to test the differences between Fabaceae and Poaceae. As only one
258 species per family were available for Rosaceae and Plantaginaceae, these families were excluded from the
259 analysis. If the data were not normally distributed, ANOVA was substituted with a Kruskal-Wallis test.
260

261 A principal component analysis (PCA) was performed on 12 variables (six root traits, four ΔC of each soil C
262 pool plus their sum, and SIR) using the mean for three replicate boxes. RER and RLP were not included in
263 the PCA since they were measured on one replicate box per species. Pearson's correlations coefficients
264 were calculated to study the relationships between root traits and ΔC in each soil C pool. To investigate
265 the effect of abiotic factors on root growth dynamics, Pearson's correlation coefficients were calculated
266 between RER, RLP, soil and air temperature, soil humidity and solar irradiation.

267 All the statistical analyses were performed in the open-source statistical environment R, version 3.4.3 (R
268 Development Core Team, 2017) using the packages *Hmisc* (Harrel 2007) and *vegan* (Oksanen *et al.* 2019).

269

270 2.3. RESULTS

271

272 2.3.1. *Effect of species identity on soil carbon accumulation (ΔC) in different C pools* 273 *associated with soil fractions*

274 Species did not significantly influence the accumulation of C in different pools, nor in the sum of C pools
275 (Fig. 1). The mean ΔC_{SUM} increase was 1.72 ± 1.45 mg C g^{-1} soil, and was highest in soil beneath *L.*
276 *corniculatus* (3.60 ± 0.70 mg C g^{-1} soil) compared to the bare soil control (0.21 ± 3.87 mg C g^{-1} soil) (Fig. 1a).
277 The mean increase in the unprotected pool ΔC_{POM} was 0.58 ± 0.34 mg C g^{-1} soil (Fig. 1b) and in the $\Delta C_{finePOM}$

278 was 1.21 ± 0.74 mg C g^{-1} soil (Fig. 1c). In the protected C_{SILT} pool, the ΔC mean increase was 0.57 ± 0.34 mg C
279 g^{-1} soil (Fig. 1d), while the $\Delta C_{SILT+CLAY}$ decreased by -0.50 ± 0.77 mg C g^{-1} soil (Fig. 1e). However, no significant
280 differences were found between any species and bare soil with regard to any C pool (Fig. S6, raw C data in
281 different soil C pools for each species at t37).

282 At the family level (Fig. 2a,b), the only significant differences in ΔC were found with regard to C_{POM} and
283 C_{SILT} . C_{POM} was significantly higher in soil beneath Poaceae species (ANOVA, $p = 0.024$, Fig. 2a) whilst C_{SILT}
284 was significantly higher in Fabaceae species (ANOVA, $p = 0.060$, Fig. 2b), and no significant differences
285 were found in C_{SILT} between Poaceae and bare soil.

286

287 2.3.2. Root elongation rate (RER) and root length production (RLP)

288 More than a threefold variation in mean daily RER_{TOT} occurred among species, ranging from 0.23 mm d^{-1}
289 (*F. rubra*) to 0.75 mm d^{-1} (*T. repens*) (Table 1). Mean daily RER_{TOT} did not differ between Fabaceae
290 (0.57 ± 0.08 mm d^{-1} on average) and Poaceae (0.42 ± 0.13 mm d^{-1}) (ANOVA, $p = 0.221$, Table 1). Mean daily
291 RER_{TOT} peaked at 0.75 mm d^{-1} in mid-February for Poaceae and then decreased, attaining a value of 0.4
292 mm d^{-1} from April to June 2017 (Fig. S7, S8). For Fabaceae species, RER_{TOT} peaked at 1.1 mm d^{-1} in May
293 2017, before decreasing sharply in June 2017 (Fig. S7 average RER_{TOT} for Fabaceae and Poaceae species;
294 Fig. S8 RER_{NEW} and RER_{OLD} for each species analyzed).

295 The RER for new roots (RER_{NEW} , 0.83 ± 0.22 mm d^{-1}) was significantly higher than that of old roots (RER_{OLD} ,
296 0.17 ± 0.09 mm d^{-1} , ANOVA, $p < 0.001$). RER_{NEW} ranged from 0.32 mm d^{-1} (*F. rubra*) to 1.13 mm d^{-1} (*D.*
297 *glomerata*) whereas RER_{OLD} ranged from 0.05 mm d^{-1} (*P. pratensis*) to 0.40 mm d^{-1} (*T. pratense*). RER_{NEW} did
298 not differ in Fabaceae than Poaceae while RER_{OLD} was higher in Fabaceae (0.25 ± 0.09 mm d^{-1} on average)
299 than Poaceae (0.13 ± 0.03 mm d^{-1}) (ANOVA, $p = 0.020$) (Table 1).

300 After 37 weeks, we measured the highest cumulative RLP_{TOT} , combining RLP_{NEW} and RLP_{OLD} , in *O. viciifolia*
301 (3.62 m) and the lowest in *F. rubra* (1.19 m) (Table 1, one value per species). Fabaceae species possessed
302 a higher RLP_{TOT} (3.37 ± 2.32 m) compared to Poaceae (2.32 ± 0.70 m) (ANOVA, $p = 0.032$). Root dynamics
303 of certain Fabaceae species were correlated with climate factors. In *L. corniculatus*, mean daily RER_{TOT} ,
304 RER_{OLD} , RER_{NEW} , RLP_{TOT} , RLP_{OLD} and RLP_{NEW} were all positively correlated with soil and air temperature and
305 solar irradiation (Table S1, S2). In *T. repens*, RER_{TOT} and RER_{NEW} , RLP_{NEW} were significantly and positively
306 correlated with soil and air temperature (Table S1, S2). With regard to Poaceae species, RER_{NEW} of *D.*
307 *glomerata* was negatively correlated with soil and air temperature and solar irradiation (Table S1). In *O.*
308 *viciifolia*, RLP_{TOT} was slightly and positively correlated with solar irradiation (Table S2). In *D. glomerata*,
309 RLP_{NEW} only, was negatively correlated with soil and air temperature (Table S2).

310 2.3.3. Root biomass, diameter and chemical composition

311 At 37 weeks, *M. sativa* had significantly greater root biomass (4.23 ± 0.42 g) compared to all other species
312 (Table 1) and in general, Fabaceae species had a significantly higher mean root biomass (2.08 ± 1.33 g)
313 compared to Poaceae (0.62 ± 0.11 g). The diameter of absorptive roots differed significantly between
314 species, with *O. viciifolia* having the thickest absorptive roots and *D. glomerata* the thinnest (0.21 ± 0.14
315 mm; Table 1). Species from the Fabaceae family had significantly thicker absorptive roots (0.39 ± 0.11 mm)
316 compared to Poaceae (0.23 ± 0.03 mm).

317 The chemical composition of absorptive roots strongly varied among species and between Fabaceae and
318 Poaceae (Table 1). Absorptive roots of Fabaceae possessed more hemicellulose + water-soluble
319 compounds (705 ± 74 mg g^{-1}) than Poaceae (543 ± 33 mg g^{-1}), a lower lignin content (Fabaceae: 173 ± 56
320 mg g^{-1} and Poaceae: 302 ± 59 mg g^{-1}), and a lower C:N ratio (Fabaceae: 19.15 ± 3.07 and Poaceae: $58.67 \pm$
321 6.34). The cellulose content did not differ either among species or families (Table 1).

322

2.3.4. Soil Substrate Induced Respiration (SIR)

SIR for the soil microbial community ranged from $2.47 \pm 0.34 \mu\text{g C-CO}_2 \text{ g}^{-1} \text{ soil h}^{-1}$ (beneath *B. erectus*) to $6.41 \pm 0.56 \mu\text{g C-CO}_2 \text{ g}^{-1} \text{ soil h}^{-1}$ (*M. sativa*) and was higher in Fabaceae compared to Poaceae (Table 1; ANOVA, $p < 0.001$).

327

2.3.5. Relationships between C accumulation, root growth dynamics, root traits, and microbial activity

The PCA conducted on the ΔC in the different C pools, SIR and root traits explained 64.6% of the variance (Fig. 3). The first PCA axis (horizontal) accounting for 44.4% of the variation governed $\Delta\text{C}_{\text{POM}}$ (negative) and $\Delta\text{C}_{\text{SILT}}$ (positive), while the remaining C pools, as well as the sum of C pools, were fairly orthogonal to $\Delta\text{C}_{\text{POM}}$ and $\Delta\text{C}_{\text{SILT}}$ and more related to the second PCA axis (vertical), that accounted for 20.2%. SIR and root biomass, diameter, and hemicellulose + water soluble compounds content of absorptive roots all went along the 1st axis (positive) together with $\Delta\text{C}_{\text{SILT}}$. Root traits linked with recalcitrance, lignin, cellulose and C:N ratio, went along the 1st axis (negative) together with $\Delta\text{C}_{\text{POM}}$. Hull polygons reflecting intraspecific variations generally had small areas and were segregated over the biplot (Fig. 2). The PCA strongly discriminated Poaceae from Fabaceae. Poaceae were all on the negative end of the first axis and were characterized by high lignin and cellulose contents, high C:N and accumulation of C in the coarse POM fraction. At the opposite end, Fabaceae were mostly located on the right end of the first axis and were characterized by a higher biomass and thicker roots that were rich in hemicellulose, favoring accumulation of C in the coarse silt fraction. The two other species were situated in intermediate positions on the axis.

$\Delta\text{C}_{\text{POM}}$ was significantly and negatively correlated with the diameter of absorptive roots and to hemicellulose + water-soluble compounds content (Table 2). $\Delta\text{C}_{\text{SILT}}$ was significantly and positively correlated with the diameter of absorptive roots, root biomass, hemicellulose + water-soluble compounds, RLP_{OLD} and mean daily RER_{OLD} (Table 2; Fig. S9), and negatively correlated with lignin and C:N ratio.

347 Variations in ΔC_{SUM} , $\Delta C_{finePOM}$ and $\Delta C_{SILT+CLAY}$ were not explained by any variables. Soil SIR was significantly
348 and positively correlated to ΔC_{SILT} , root biomass, RER_{OLD} and RLP_{OLD} , whereas hemicellulose + water-soluble
349 compounds was negatively correlated with lignin content and C:N ratio (Table 2).

350

351 2.4. DISCUSSION

352 Total C accumulation in soil was not affected by plant species, but in line with our hypotheses, the
353 accumulation of C into different soil C pools, specifically coarse POM and coarse silt, was indeed dependent
354 on root traits. C_{POM} was associated with the recalcitrance of absorptive roots, while $C_{SILT+CLAY}$ was affected
355 by more labile compounds and the higher RER and RLP of older roots. Poaceae species favored C
356 accumulation in the C_{POM} pool, but Fabaceae in the $C_{SILT+CLAY}$ pool. Total soil organic carbon (SOC) is,
357 therefore, a poor indicator of C sequestration and persistence in soils, and studies of C sequestration
358 should focus on defining C input into different C pools associated with different textural fractions
359 (Wiesmeier *et al.*, 2019).

360

361 2.4.1. Hypothesis 1: Root elongation rate and root N content are expected to favor C 362 accumulation in the coarse silt and fine silt + clay soil C pools

363 We hypothesized that a fast RER would promote C sequestration in coarse silt and fine silt+ clay soil
364 fractions, through an increase in exudation and microbial activity along newly initiated roots. Interestingly,
365 RER_{OLD} and RLP_{OLD} were significantly and positively correlated with microbial activity (SIR) and ΔC_{SILT} , but
366 not with the RER and RLP of newly initiated roots, that had very high rates of growth. Dennis *et al.* (2010)
367 hypothesized that rapidly elongating root tips grow quickly out of the main zone of microbial activity, that
368 is established once root exudates have been consumed. These established microbial communities then
369 consume rhizodeposits from mucilage and cell senescence as well as root exudates from growing roots in

370 proximity. Therefore, slower growing older roots would be maintained in this zone of high microbial
371 activity, and C sequestration in the coarse silt fraction would be higher, especially in N₂-fixing species with
372 populations of bacteria distributed along roots, away from the apex. N₂-fixing *Rhizobium* bacteria also
373 increase root elongation (Garrido-Oter *et al.*, 2018), likely inducing a feedback mechanism whereby a
374 stimulated RER results in a higher exudation rate (Garcia *et al.* 2001), acting as a substrate for newly
375 colonizing *Rhizobium* communities. Although the role of microbial communities is of utmost importance
376 for C input into the soil, differences in the use of C within plants could also explain the lack of a relationship
377 between RER_{NEW}, RLP_{NEW}, SIR and C_{SILT}. In fast-growing, newly initiated roots, we suggest that C in the form
378 of non-structural carbohydrates (NSC, produced during photosynthesis), will be used preferentially for cell
379 production and expansion, as found in a recent seasonal study of root elongation and NSC fluxes (Wang *et al.*,
380 *et al.*, 2018). In older roots with lower RER, less NSC is required for growth, and excess NSC would be freely
381 exuded, reflected in the high microbial activity that we observed. Accordingly, this also explains the lack
382 of relationships between RER_{TOT}, RLP_{TOT} and C_{SILT}, probably due to the mixed effects of new and old roots.

383 Contrary to our hypothesis, RER and RLP did not promote accumulation of C in the C_{SILT+CLAY} pool.
384 Surprisingly, the C_{SILT+CLAY} pool was the only pool where C was actually lost over the 37 weeks, in both bare
385 soil and beneath all plant species, and this mineralisation of C is not explained by microbial activity or by
386 any root traits. Although the origin of C in the fine silt+clay pool is also from plant and microbial exudates
387 and exopolysaccharides (Dungait *et al.*, 2012; Cotrufo *et al.*, 2013), disturbance (during soil preparation)
388 will result in an accelerated mineralisation of C, as it is released from disrupted soil aggregates
389 (Franzluebbers, 1999). This increase in C mineralisation is higher in clays, as organic matter that was highly
390 protected within the clay fraction will be released during disruption, providing a new pool of C available
391 to microorganisms (Hassink, 1992).

392

393 2.4.2. Hypothesis 2: more recalcitrant root traits are expected to favor the unprotected
394 coarse POM fraction
395 Root traits linked to recalcitrance (high cellulose, lignin and C:N) were not significantly correlated to C
396 accumulation in the coarse POM fraction, but the PCA showed that this suite of traits was linked to C_{POM}
397 on the first axis, suggesting a relationship with C accumulation in the coarse POM fraction, in agreement
398 with our second hypothesis. Recalcitrant compounds, such as lignin, cellulose and the C:N ratio, have all
399 been reported to decrease root decomposition rates (Poirier *et al.*, 2018). Lignin-carbohydrate complexes
400 prevent polymer-hydrolyzing enzymes access to substrates, thus reducing the degradability of plant
401 organic matter (Cornu *et al.*, 1994, Malherbe and Cloete, 2002). Microbial activity was also strongly and
402 negatively correlated to C:N and lignin content, probably because lignin reduces the accessibility of
403 polysaccharides to microorganisms through the formation of links between lignin and polysaccharides
404 (Bertrand *et al.*, 2006). Products of lignin degradation can also react with ammonia or amino acids to form
405 further recalcitrant complexes that are less available to microorganisms (Nömmik and Vahtras, 1982).
406 Interestingly, C accumulation in the coarse silt fraction was negatively correlated with recalcitrant traits
407 (lignin and C:N) but positively with hemicellulose content and root diameter. Hemicelluloses comprise
408 polysaccharides soluble in alkali and are easily degradable by microorganisms (Dekker, 1985).
409 Hemicelluloses are usually produced to the detriment of lignin and favor tissue degradability through
410 higher accessibility to amorphous phases in the lignocellulose structure (Malherbe and Cloete, 2002).
411 Microorganisms will use this easily degradable C for growth and respiration and the production of exudates
412 and exopolysaccharides, that are used as a substrate for subsequent microbial communities (Dennis *et al.*,
413 2010). These low molecular weight compounds are the main precursors of C in the coarse silt pool (Cotrufo
414 *et al.*, 2013, Vidal *et al.*, 2018), explaining the high C_{SILT} we found beneath N_2 -fixing species with high
415 hemicellulose + water soluble compounds and low lignin contents. Absorptive roots were negatively
416 correlated with C accumulation in the coarse POM C pool and positively correlated with C accumulation in

417 the coarse silt C pool. Absorptive roots may have a higher turnover rate and undergo rapid transformation
418 through microbial degradation, explaining the low accumulation in the coarse POM C pool and the positive
419 correlation with the coarse silt C pool. However, this relationship may also be an artefact because Poaceae
420 roots are inherently very fine compared to Fabaceae roots (Roumet *et al.*, 2006; 2016), highlighting that
421 the understanding of relationships between C sequestration and morphological traits is challenging
422 because of their inherent nature and plasticity (Poirier *et al.*, 2018).

423 Our results show therefore that the C_{POM} pool is largely driven by the slow decomposition of recalcitrant
424 plant tissues, as suggested by De Deyn *et al.*, (2008) and Poirier *et al.* 2018. These processes were not
425 observed in the C_{finePOM} pool. As this C pool is a transitional pool between C_{POM} and C_{SILT} , variability in
426 processes would be high and measurable effects could have been canceled out over the course of the
427 experiment.

428 *2.4.3. Hypothesis 3: Fabaceae and Poaceae strongly differ in their influence on* 429 *accumulation of C into different soil fractions*

430 Contrary to what has been observed in previous studies (Fornara and Tilman, 2008; Binkley 2005), we did
431 not find evidence of a greater accumulation of total C (ΔC_{SUM}) in soil beneath N_2 -fixing species, because
432 variability was high within Fabaceae. However, we showed that Poaceae and Fabaceae strongly differed
433 in their effect on the accumulation of C into different soil C pools. Roots of Poaceae, as compared to
434 Fabaceae, have a lower RER and RLP. Poaceae produce thinner roots, rich in lignin and cellulose with a
435 higher C:N ratio. These more recalcitrant tissues slow down microbial activity and their decomposition
436 rate as shown previously (Roumet *et al.* 2016; Freschet *et al.* 2017). Due to their particular chemical
437 composition, Poaceae promote C accumulation in the unprotected coarse POM C pool, but impede C
438 accumulation in the more stable coarse silt C pool (Fig. 4). However, roots of Fabaceae grow faster and
439 produce thick roots that are easily degradable, since they are rich in N (low C:N ratio) and hemicellulose.
440 These traits favor microbial activity shown by the SIR that was 40% higher beneath Fabaceae. The

441 production of labile C from microbial communities, that boost the accumulation of C in the protected
442 coarse silt fraction, also reduce losses in the fine silt+clay pool. A scheme illustrating the mechanisms of
443 soil C accumulation in different soil C pools for Poaceae and Fabaceae can be found in Fig. 4.

444

445 2.5. CONCLUSION

446 Our findings show that specific plant root traits influence the accumulation of C into different pools, largely
447 through the mediation of microbial activity, shaping the C pathway in soil and, finally, its persistence. Root
448 elongation rate and length production promote microbial activity in slower-growing older roots only,
449 suggesting either a spatial influence of root exudate accessibility on microbial communities, or a
450 relationship between NSC use in roots and available exudates for microbial consumption. Although root
451 growth was affected by temperature in several species, it was not possible to determine if temperature
452 could affect microbial activity and C sequestration in this experiment. Nevertheless, the link between soil
453 temperature, root growth and microbial activity remains a promising line of inquiry for future soil C studies
454 (Žifčáková *et al.*, 2015). Recalcitrant traits (lignin, cellulose and high C:N) promoted accumulation of C in
455 the unprotected coarse POM fraction. Root traits associated with high degradability, labile C input and
456 microbial activity (hemicellulose + water soluble compounds, RER_{OLD} , RLP_{OLD}) stimulated C accumulation in
457 the protected coarse silt fraction. Therefore, due to their root characteristics, Poaceae species increased
458 C in the C_{POM} pool, and Fabaceae in the C_{silt} pool. However, studies on long term C dynamics are needed to
459 understand the microbial evolution and consequent C accumulation in different pools. Moreover, the
460 influence of different soils and associated microbial communities need to be taken into consideration for
461 a broader understanding of C pool dynamics. Our results will not only be useful for identifying plant species
462 capable of enhancing long-term C sequestration in soil, but will also contribute significantly to the
463 understanding of mechanistic processes within the carbon cycle.

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643 FIGURES AND TABLES

644 **Table 1:** Plant root traits and microbial activity for the 12 herbaceous species. Mean data are also given for species from Fabaceae and Poaceae families.

645

Family Species Acronym code	Fabaceae					Poaceae					Rosaceae	Plantaginaceae	Effect of species (ANOVA)			Fabaceae	Poaceae	Effect of families (ANOVA)		
	<i>Lotus corniculatus</i> Lc	<i>Trifolium repens</i> Tr	<i>Trifolium pratense</i> Tp	<i>Onobrychis viciifolia</i> Ov	<i>Medicago sativa</i> Ms	<i>Bromus erectus</i> Be	<i>Festuca rubra</i> Fr	<i>Dactylis glomerata</i> Dg	<i>Poa pratense</i> Pp	<i>Lolium perenne</i> Lp	<i>Sanguisorba minor</i> Sm	<i>Plantago lanceolata</i> Pl	Df	F	p			Df	F	p
RERTOT (mm d ⁻¹)	0.55±1.1 (a)	0.75±1.5 (a)	0.59±1 (a)	0.53±10 (a)	0.57±0.9 (a)	0.42±0.6 (a)	0.23±0.50 (a)	0.58±0.9 (a)	0.66±2.90 (a)	0.42±0.8 (a)	0.39±0.5 (a)	0.5±0.7 (a)	11,175	18.9	0.06 ¹	0.57±0.08	0.42±0.13	2,9	1.8	0.22
REROLD (mm d ⁻¹)	0.17±0.07 (a)	0.2±0.08 (a)	0.4±0.11 (a)	0.21±0.05 (a)	0.26±0.08 (a)	0.12±0.03 (a)	0.09±0.04 (a)	0.14±0.06 (a)	0.05±0.02 (a)	0.11±0.04 (a)	0.13±0.04 (a)	0.16±0.04 (a)	11,178	17.9	0.08 ¹	0.25±0.09	0.13±0.03	2,9	6.3	0.02
RERNEW (mm d ⁻¹)	0.99±0.27 (a)	0.91±0.21 (a)	0.73±0.15 (a)	0.81±0.16 (a)	0.97±0.12 (a)	1.06±0.29 (a)	0.32±0.08 (a)	1.13±0.25 (a)	0.96±0.31 (a)	0.67±0.11 (a)	0.66±0.08	0.76±0.11	11,178	17.4	0.1 ¹	0.88±0.11	0.79±0.33	2,9	0.37	0.7
RLPTOT (m)	3.03	3.26	3.36	3.62	3.61	2.55	1.19	2.76	2.16	2.95	2.31	3.04	-	-	-	3.37±2.32	2.32±0.7	2,9	5.17	0.03
RLPOLD (m)	4.51	4.91	4.26	4.96	5.23	3.47	1.05	3.30	2.76	4.29	3.32	4.50	-	-	-	1.33±0.61	0.61±0.2	2,9	7.5	0.01
RLPNEW (m)	0.89	1.05	1.78	1.35	1.60	0.78	0.53	0.81	0.32	0.62	0.47	0.99	-	-	-	4.78±2.97	2.97±1.21	2,9	5.12	0.03
Root biomass (g)	1.53±2.6 (bc)	0.55±1.13 (c)	2.01±0.62 (b)	2.06±1.1 (b)	4.23±0.42 (a)	0.65±0.08 (c)	0.70±1.6 (c)	0.58±0.51 (c)	0.60±0.83 (c)	0.57±0.57 (c)	0.91±2.13 (c)	0.49±0.60 (c)	11,24	27.3	<0.001	2.08±1.33	0.62±0.11	1,28	10.9	<0.001¹
Diameter absorptive roots (mm)	0.47±0.05 (b)	0.28±0.05 (de)	0.32±0.04 (cd)	0.55±0.18 (a)	0.35±0.18 (c)	0.27±0.02 (def)	0.22±0.05 (fg)	0.21±0.14 (g)	0.22±0.19 (g)	0.22±0.05 (fg)	0.26±0.09 (efg)	0.28±0.14 (de)	11,23	98.7	<0.001	0.39±0.11	0.23±0.03	1,27	31.7	<0.001
Hemicell. +H₂O soluble compounds (mg g ⁻¹)	779.5±58.1 (a)	612±8.3 (bcde)	674.5±44.6 (abc)	704.1±78.8 (ab)	755.1±31 (a)	533.9±12.1 (de)	572.3±27.3 (cde)	530.4±57.6 (de)	562.4±31.2 (cde)	520.5±31.7 (e)	703.2±NA (ab)	639.4±12.7 (bcd)	11,21	17.4	<0.001	705.11±74.39	543.51±33.56	1,25	51.5	<0.001
Cellulose (mg g ⁻¹)	101.6±6.3 (a)	163.2±8.7 (a)	102.5±17.7 (a)	120±69.3 (a)	123.9±27.7 (a)	177.9±76.5 (a)	160.7±44.8 (a)	137.5±7.7 (a)	89.1±6 (a)	181.6±32.3 (a)	140±NA (a)	151.2±54 (a)	11,21	1.7	0.13	122.39±33.61	154.89±50.8	1,25	3.9	0.06
Lignine (mg g ⁻¹)	118.9±57.8 (c)	224.8±0.4 (abc)	223±33.7 (abc)	175.9±9.4 (abc)	120.9±3.5 (c)	288.3±88.5 (ab)	267±39.5 (ab)	332.1±65.3 (a)	348.4±25.1 (a)	297.9±62.1 (ab)	156.8±NA (bc)	209.3±66.7 (abc)	11,21	6.7	<0.001	172.5±56.53	301.6±59.02	1,25	33.7	<0.001
C:N ratio	15.4±0.8 (c)	21.2±1.1 (c)	17.6±3.20 (c)	20.5±3.2 (c)	21.4±2 (c)	50.8±7.6 (b)	61.3±2.6 (ab)	62.5±6.4 (ab)	59.1±4.1 (ab)	61±4.4 (ab)	69.8±3.2 (a)	68.9±3.1 (a)	11,21	96.4	<0.001	19.15±3.07	58.67±6.34	1,25	436	<0.001
SIR (µg C-CO ₂ g ⁻¹ soil h ⁻¹)	5.37±0.46 (ab)	4.43±0.48 (bc)	6.11±0.35 (a)	4.07±0.18 (bcd)	6.41±0.56 (a)	2.47±0.34 (d)	3.22±0.11 (cd)	3.58±0.28 (cd)	3.15±0.15 (cd)	3.17±0.1 (cd)	3.16±0.23 (cd)	2.99±0.11 (cd)	9,20	16	<0.001	5.28±1	3.12±0.4	1,28	45.9	<0.001

646 ¹ Distribution not normal, Kruskal-Wallis test instead of ANOVA

647 For each species and for Poaceae and Fabaceae, mean and standard deviation of the following variables are given. Abbreviations: RER_{TOT} – root elongation
 648 rate of the entire root system; RER_{OLD} – of old roots older than 2 weeks; RER_{NEW} – of new roots younger than 2 weeks; RLP_{TOT} root length production of the
 649 entire root system; RLP_{OLD} – of old roots; RLP_{NEW} – of new roots; Root biomass – total root biomass of a core sampled at t37; Diameter absorptive –mean

650 diameter of absorptive roots at t37; Hemicellulose + H₂O – concentration of hemicellulose and water soluble compounds in absorptive roots, Cellulose, Lignin
651 – concentrations of cellulose and lignin in absorptive roots; C:N – ratio of carbon and nitrogen in absorptive roots; SIR – microbial substrate induced respiration.
652 Different letters next to the average value indicate statistically significant differences ($p < 0.05$) between species or families according to Tukey HSD tests. DF
653 – degree of freedom (number of species - 1, number of observations). Statistically significant values ($p < 0.05$) are indicated in bold text.

654 **Table 2:** Pearson's correlation coefficients (*r*) showing relationships between root variables and soil variables.

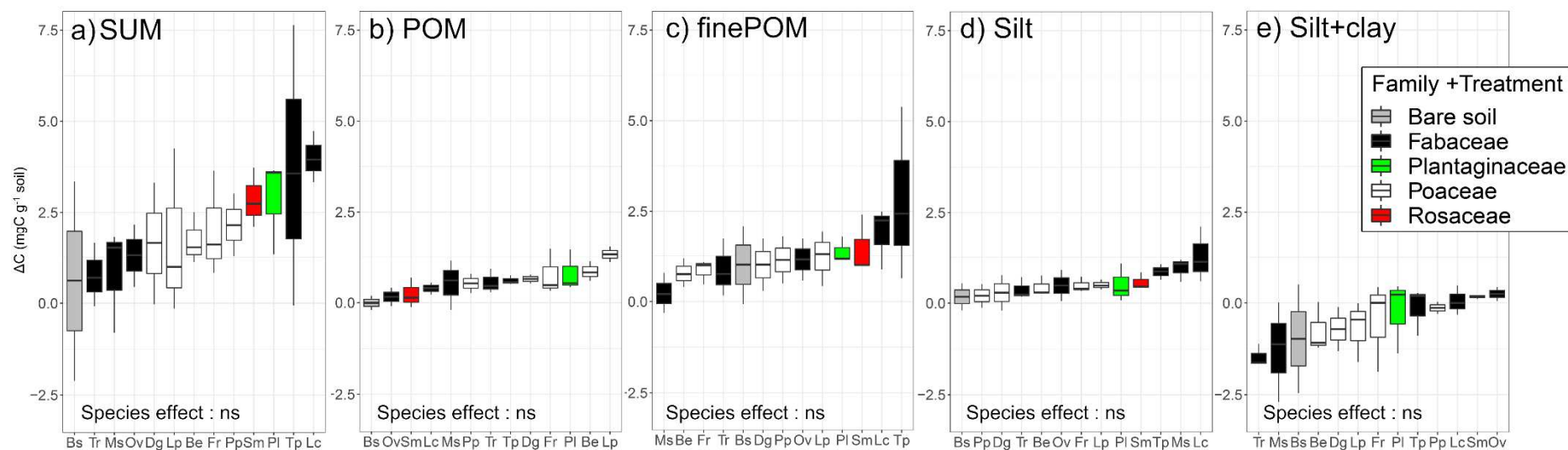
		ΔC_{SUM}	ΔC_{POM}	$\Delta C_{\text{finePOM}}$	ΔC_{silt}	$\Delta C_{\text{silt+clay}}$	SIR
Root dynamics	RER _{TOT}	-0.32	-0.20	-0.06	-0.10	-0.17	0.45
	RER _{OLD}	-0.17	-0.25	-0.04	0.72**	0.37	0.74**
	RER _{NEW}	-0.24	-0.13	-0.20	-0.12	-0.55	0.18
	RLP _{TOT}	-0.34	-0.33	-0.18	0.37	0.34	0.51
	RLP _{NEW}	-0.30	-0.30	-0.15	0.33	0.17	0.43
	RLP _{OLD}	-0.32	-0.25	-0.22	0.66*	0.34	0.70*
Root morphology	Root biomass	-0.20	-0.46	-0.37	0.78**	0.30	0.80**
	Absorptive roots diameter	-0.20	-0.72**	-0.17	0.79**	0.14	0.56
Root chemical traits	Hemicell. +H ₂ O soluble compounds	-0.06	-0.61*	-0.13	0.82**	-0.3	0.68*
	Cellulose	-0.18	0.47	-0.14	-0.22	0.19	-0.53
	Lignin	0.15	0.56	0.26	-0.84***	0.49	-0.60*
	Root C:N ratio	0.26	0.37	0.09	-0.68*	0.29	-0.86***
	SIR	-0.16	-0.19	-0.11	0.65*	-0.57	

655 In bold, significant *r* values: * significant at 0.05 > *p* ≥ 0.01 level, ** significant at 0.01 > *p* ≥ 0.001 level, and *** significant at *p* < 0.001 level

656 Abbreviations: RER_{TOT} – root elongation rate of the entire root system; RER_{OLD} – of old roots older than two weeks; RER_{NEW} – of new roots younger than 2
657 weeks; RLP_{TOT} – root length production of the entire root system; RLP_{OLD} – of old roots; RLP_{NEW} – of new roots; Root biomass – total root biomass of a core
658 sampled at t37; Diameter absorptive – mean diameter of absorptive roots at t37; Hemicellulose + H₂O – concentration of hemicellulose and water soluble
659 compounds in absorptive roots, Cellulose, Lignin – concentrations of cellulose and lignin in absorptive roots; C:N – ratio of carbon and nitrogen in absorptive
660 roots; SIR – microbial substrate induced respiration.

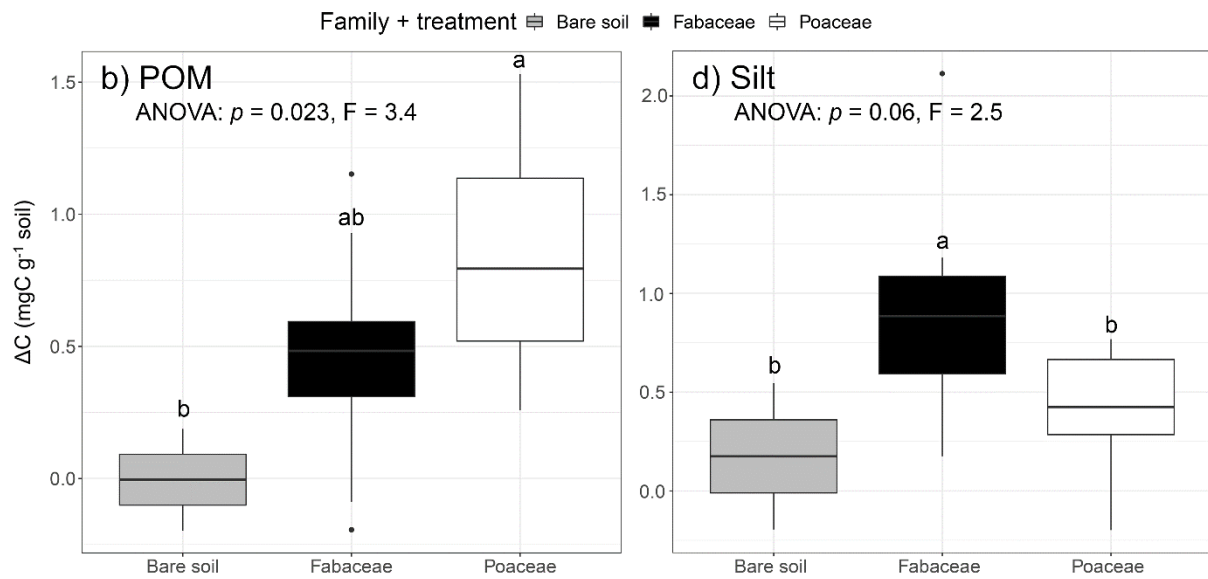
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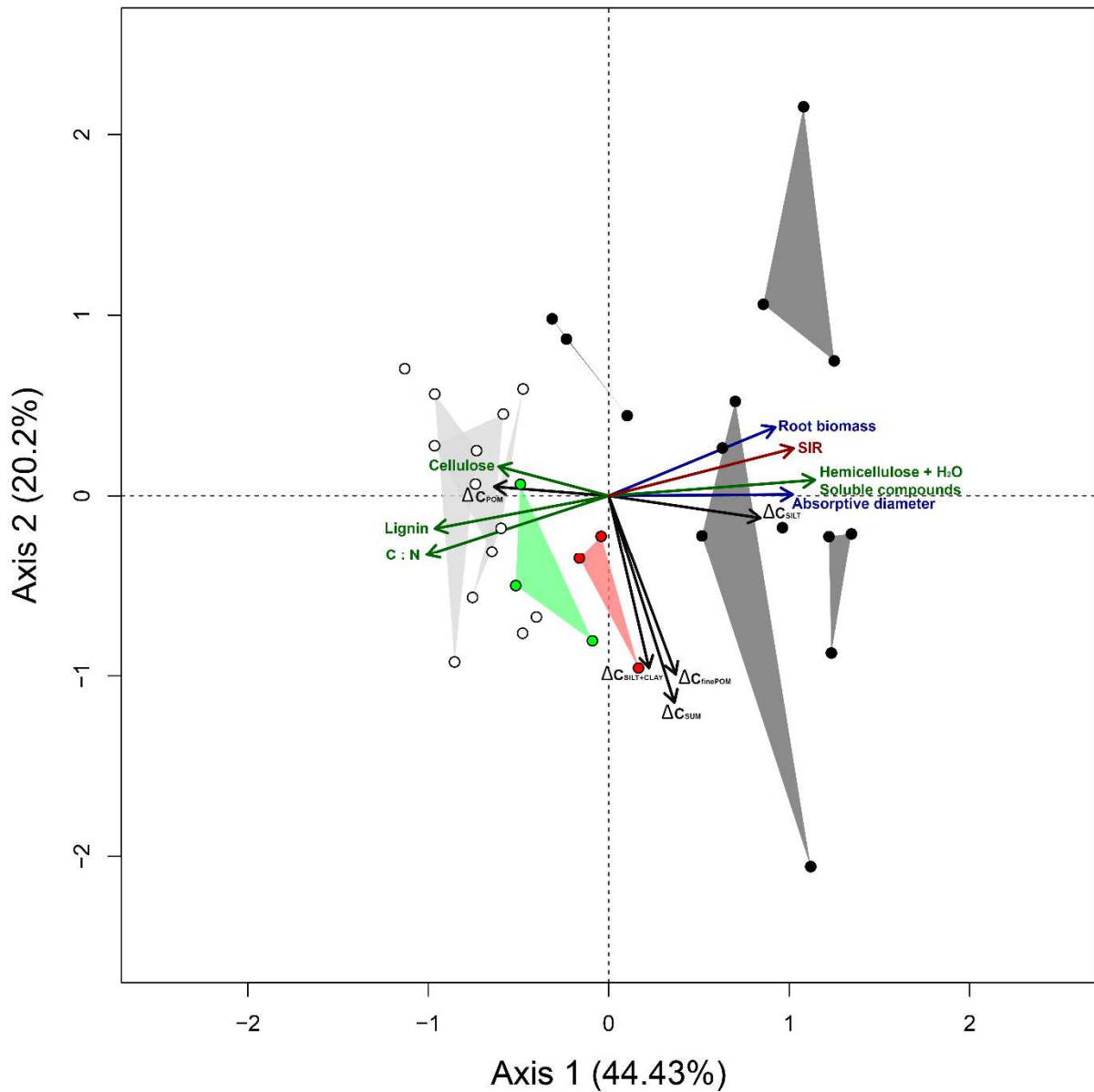
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664 **Figure 1:** Comparison of the difference ($\Delta C = C_{137} - C_{10}$) in carbon (C) after 37 weeks between different soil fractions for each species. a) total C (ΔC_{SUM}), b) C in the coarse POM
 665 fraction (ΔC_{POM}), c) C in the fine POM fraction ($\Delta C_{\text{finePOM}}$), d) C in the coarse silt fraction (ΔC_{SILT}) and e) C in the fine silt+clay fraction ($\Delta C_{\text{SILT+CLAY}}$). In each boxplot, the lower
 666 edge of the box corresponds to the 25th percentile data point, while the top edge of the box corresponds to the 75th percentile data point. The line within the box represents the
 667 median.



668

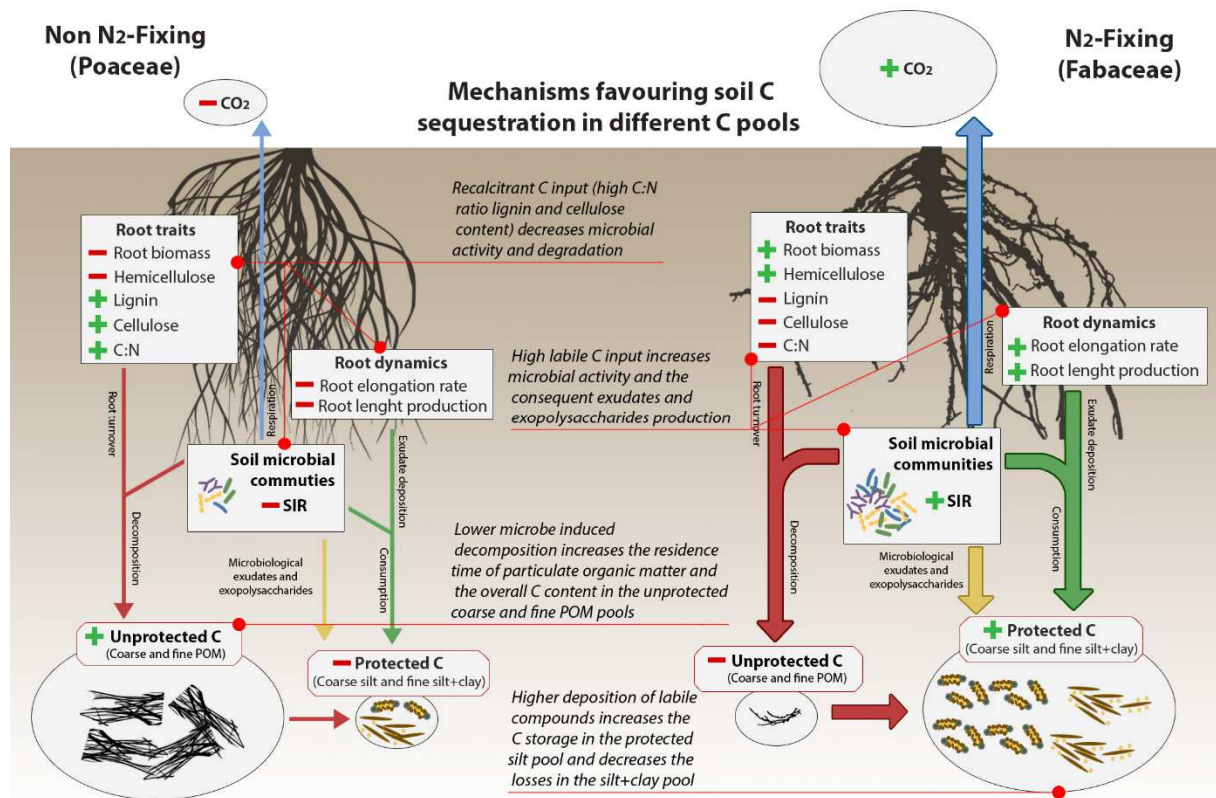
669 **Figure 2:** Comparison of the difference ($\Delta C = C_{137} - C_{10}$) in carbon (C) after 37 weeks among families and control.
 670 a) C in the coarse POM fraction (ΔC_{POM}) and b) C in the silt fraction (ΔC_{silt}). No significant differences were found
 671 in total C (ΔC_{SUM}), C in the fine POM fraction ($\Delta C_{\text{finePOM}}$), or in C in the silt+clay fraction ($\Delta C_{\text{silt+clay}}$). In each boxplot,
 672 the lower edge of the box corresponds to the 25th percentile data point, while the top edge of the box corresponds
 673 to the 75th percentile data point. The line within the box represents the median and black dots indicate outliers.
 674 Different letters above the boxplots indicate statistically significant differences ($p < 0.05$) among families and control
 675 according to a Tukey HSD test. Note the graduation differences in y-axis between a) and b).



676

677 **Figure 3:** Principal Component Analysis of six soil variables (five carbon pool changes and SIR) and six root
 678 variables measured on 12 species. Black dots are Fabaceae, white dots are Poaceae, red dots are *Sanguisorba*
 679 *minor*, and green dots are *Plantago lanceolata*. The Hull polygons unify the different replicates for the same species.
 680 Abbreviations: SIR – microbial substrate induced respiration; Hemicellulose + H₂O – concentration of hemicellulose
 681 and water soluble compounds in absorptive roots, Cellulose, Lignin – concentrations of cellulose and lignin in
 682 absorptive roots; C:N – ratio of carbon and nitrogen in absorptive roots; ΔC_{POM} – difference ($\Delta C = C_{137} - C_{10}$) in
 683 carbon (C) after 37 weeks for the coarse POM C pool; $\Delta C_{finePOM}$ – for the fine POM C pool; ΔC_{SILT} – for the silt C
 684 pool; $\Delta C_{SILT+CLAY}$ – for the silt + clay f C pool; ΔC_{SUM} – sum of different fraction ΔC as the total changes in C
 685 concentration in soil.

686



687

688 **Figure 4:** Conceptual scheme of carbon (C) sequestration mechanisms into different soil pools under N₂-fixing
 689 (Fabaceae) and non N₂-fixing (Poaceae) species. Square boxes refer to the major actors affecting C sequestration.
 690 Ellipses show the destination of C into unprotected pools (POM and finePOM) and protected pool (coarse silt and
 691 fine silt+clay). Text in the central column describes the mechanisms favoring C sequestration into soil C pools. The
 692 arrows symbolize the processes of transformation or transport of C into different pools. Arrows colors represent the
 693 nature of the C: red C deriving from root turnover, green C from root rhizodeposition, yellow C from microbial
 694 exudates and exopolysaccharides and blue the C respired back in atmosphere as CO₂. The thickness of the arrows
 695 is qualitative, with wider arrows reflecting higher C fluxes. The signs: "+" (in green) means an increase and "-" (in
 696 red) means a decrease.

697

698

699 SUPPLEMENTARY MATERIALS

700 **Table S1:** Pearson’s correlation coefficients (r) showing relationships between mean root elongation rate and climate. Analysis include: Daily mean soil temperature, daily mean
 701 air temperature, daily minimum air temperature, daily maximum air temperature, mean daily solar radiation and a) RER_{TOT}: mean root elongation rate of a single root, b) RER_{OLD}:
 702 mean root elongation rate of a single roots that was already present at the previous sampling date (older than 14 days), c) RER_{NEW}: mean root elongation rate of a single newly
 703 initiated root (aged 1 to 14 days). The correlations have been calculated for each RER sampling dates, every two weeks for each species of the experiment over the 0-10 months
 704 period.

705

a - Correlation between climate factors and RER of the total root system						
		Soil temperature	Average air temperature	Minimum air temperature	Maximum air temperature	Average solar radiation
Poaceae	<i>Dactylis glomerata</i>	-0.23	-0.21	-0.1	-0.18	-0.1
	<i>Lolium perenne</i>	0.01	-0.04	-0.03	0.09	0.38
	<i>Festuca rubra</i>	-0.21	-0.27	-0.34	-0.2	0.15
	<i>Bromus erectus</i>	-0.02	-0.05	-0.01	0.02	0.29
	<i>Poa pratensis</i>	0.1	0.08	0.34	0.1	0.43
Fabaceae	<i>Trifolium repens</i>	0.76***	0.81***	0.45	0.68**	0.33
	<i>Trifolium pratense</i>	0.02	-0.02	-0.02	0.09	0.4
	<i>Lotus corniculatus</i>	0.70**	0.70**	0.44	0.68**	0.72**
	<i>Onobrychis viciifolia</i>	0.13	0.09	0.3	0.17	0.54*
	<i>Medicago sativa</i>	0.01	0.01	-0.05	0.02	0.22
Rosaceae	<i>Sanguisorba minor</i>	-0.06	-0.06	-0.06	-0.04	0.16
Plantaginaceae	<i>Plantago lanceolata</i>	0	-0.04	0	0.04	0.3

706

b - Correlation between climate factors and RER of the old roots						
Family	Species	Soil temperature	Average air temperature	Minimum air temperature	Maximum air temperature	Average solar radiation
Poaceae	<i>Dactylis glomerata</i>	-0.27	-0.26	-0.17	-0.22	-0.12
	<i>Lolium perenne</i>	0.14	0.13	0.31	0.09	0.21
	<i>Festuca rubra</i>	-0.25	-0.3	-0.41	-0.25	0.05
	<i>Bromus erectus</i>	0.17	0.28	-0.07	0.2	0.29
	<i>Poa pratensis</i>	-0.03	0	0.4	0.04	0.21
Fabaceae	<i>Trifolium repens</i>	0.28	0.28	0.15	0.4	0.45
	<i>Trifolium pratense</i>	0.18	0.25	-0.07	0.24	0.38
	<i>Lotus corniculatus</i>	0.73**	0.71**	0.51	0.74**	0.89***
	<i>Onobrychis viciifolia</i>	0.04	0.02	0.26	0.07	0.36
	<i>Medicago sativa</i>	0.03	0.01	-0.05	0.03	0.23
Rosaceae	<i>Sanguisorba minor</i>	0.13	0.2	-0.08	0.1	-0.33
Plantaginaceae	<i>Plantago lanceolata</i>	-0.05	-0.07	-0.05	-0.03	0.21

c - Correlation between climate factors and RER of new roots						
		Soil temperature	Average air temperature	Minimum air temperature	Maximum air temperature	Average solar radiation
Poaceae	<i>Dactylis glomerata</i>	-0.55*	-0.54*	-0.47	-0.60*	-0.66**
	<i>Lolium perenne</i>	0.03	0	-0.05	0.1	0.29
	<i>Festuca rubra</i>	-0.11	-0.16	-0.04	-0.15	0.26
	<i>Bromus erectus</i>	-0.06	-0.07	-0.08	-0.05	0.17
	<i>Poa pratensis</i>	0.09	0.08	0.36	0.06	0.38
Fabaceae	<i>Trifolium repens</i>	0.59*	0.67**	0.17	0.57*	0.17
	<i>Trifolium pratense</i>	0.11	0.1	0.37	0.08	0.41
	<i>Lotus corniculatus</i>	0.76***	0.75**	0.48	0.76***	0.76**
	<i>Onobrychis viciifolia</i>	0.13	0.1	0.3	0.15	0.51
	<i>Medicago sativa</i>	0.04	0.06	-0.35	0.04	0.21
Rosaceae	<i>Sanguisorba minor</i>	0.09	0.16	0.04	0.06	-0.05
Plantaginaceae	<i>Plantago lanceolata</i>	-0.05	-0.06	-0.06	-0.03	0.19

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710 **Table S2:** Pearson's correlation coefficients (r) showing relationships between cumulative root length production and climate. Analysis include: daily mean soil temperature, daily
 711 mean air temperature, daily minimum air temperature, daily maximum air temperature, mean daily solar radiation, and a) RLP_{TOT}: cumulative root length production of all the
 712 analyzed 30 roots b) RLP_{OLD}: cumulative root length production of roots that were already present at the previous sampling date (older than 14 days), c) RLP_{NEW}: cumulative root
 713 length production of newly initiated roots (aged 1 to 14 days). The correlations have been calculated for each RLP sampling dates, every two weeks for each species of the
 714 experiment over the 0-10 months period.

715

a - Correlation between climate factors and RLP of the total root system						
		Soil temperature	Average air temperature	Minimum air temperature	Maximum air teperature	Average solar radiation
Poaceae	<i>Dactylis glomerata</i>	-0.4	-0.41	-0.46	-0.45	-0.38
	<i>Lolium perenne</i>	-0.02	-0.05	-0.08	0.04	0.29
	<i>Festuca rubra</i>	-0.19	-0.25	-0.32	-0.18	0.18
	<i>Bromus erectus</i>	-0.01	-0.05	0.01	0.03	0.31
	<i>Poa pratensis</i>	0.12	0.09	0.35	0.12	0.46
Fabaceae	<i>Trifolium repens</i>	0.36	0.37	-0.01	0.33	0.38
	<i>Trifolium pratense</i>	0.03	0	0.03	0.07	0.34
	<i>Lotus corniculatus</i>	0.72**	0.72**	0.46	0.70**	0.74**
	<i>Onobrychis viciifolia</i>	0.13	0.09	0.3	0.17	0.54*
	<i>Medicago sativa</i>	0.01	-0.01	-0.04	0.02	0.24
Rosaceae	<i>Sanguisorba minor</i>	-0.06	-0.07	-0.05	-0.04	0.18
Plantaginaceae	<i>Plantago lanceolata</i>	0	-0.04	0	0.04	0.31

716

b - Correlation between climate factors and RLP of the old roots						
		Soil temperature	Average air temperature	Minimum air temperature	Maximum air teperature	Average solar radiation
Poaceae	<i>Dactylis glomerata</i>	-0.29	-0.34	-0.38	-0.28	0
	<i>Lolium perenne</i>	0	-0.07	-0.02	-0.08	0.24
	<i>Festuca rubra</i>	-0.16	-0.22	-0.31	-0.15	0.19
	<i>Bromus erectus</i>	0.18	0.25	0	0.23	0.38
	<i>Poa pratensis</i>	0.02	0.03	0.4	0.1	0.31
Fabaceae	<i>Trifolium repens</i>	0.28	0.26	0.2	0.41	0.51*
	<i>Trifolium pratense</i>	0.22	0.26	0.02	0.29	0.47
	<i>Lotus corniculatus</i>	0.72**	0.69**	0.52	0.75***	0.90***
	<i>Onobrychis viciifolia</i>	0.06	0.03	0.28	0.1	0.42
	<i>Medicago sativa</i>	0.04	0	0.02	0.08	0.33
Rosaceae	<i>Sanguisorba minor</i>	-0.04	-0.05	-0.06	-0.02	0.18
Plantaginaceae	<i>Plantago lanceolata</i>	0	-0.04	0.02	0.05	0.33

c - Correlation between climate factors and RLP of new roots						
		Soil temperature	Average air temperature	Minimum air temperature	Maximum air teperature	Average solar radiation
Poaceae	<i>Dactylis glomerata</i>	-0.52*	-0.51*	-0.51	-0.54*	-0.48
	<i>Lolium perenne</i>	-0.25	-0.3	-0.43	-0.27	0.04
	<i>Festuca rubra</i>	-0.08	-0.14	-0.03	-0.1	0.33
	<i>Bromus erectus</i>	0.01	-0.03	0.01	0.05	0.33
	<i>Poa pratensis</i>	0.13	0.1	0.34	0.12	0.46
Fabaceae	<i>Trifolium repens</i>	0.51*	0.57*	0.11	0.49	0.26
	<i>Trifolium pratense</i>	0.17	0.14	0.36	0.17	0.51*
	<i>Lotus corniculatus</i>	0.69**	0.69**	0.41	0.70**	0.77***
	<i>Onobrychis viciifolia</i>	0.06	0.01	0.02	0.13	0.43
	<i>Medicago sativa</i>	0.03	0.02	-0.03	0.05	0.27
Rosaceae	<i>Sanguisorba minor</i>	0.17	0.14	0.19	0.22	0.46
Plantaginaceae	<i>Plantago lanceolata</i>	0.01	-0.03	0.01	0.05	0.32

717

727 **Figure S2:** Growth boxes used in the experiment. Above a gravel layer, soil was homogeneously compacted into
 728 growth boxes. Seeds were sown at a density of 155 plants m⁻². Panes of plexiglass on the front of the box allowed
 729 root elongation to be observed over the 37 week long experiment.

730

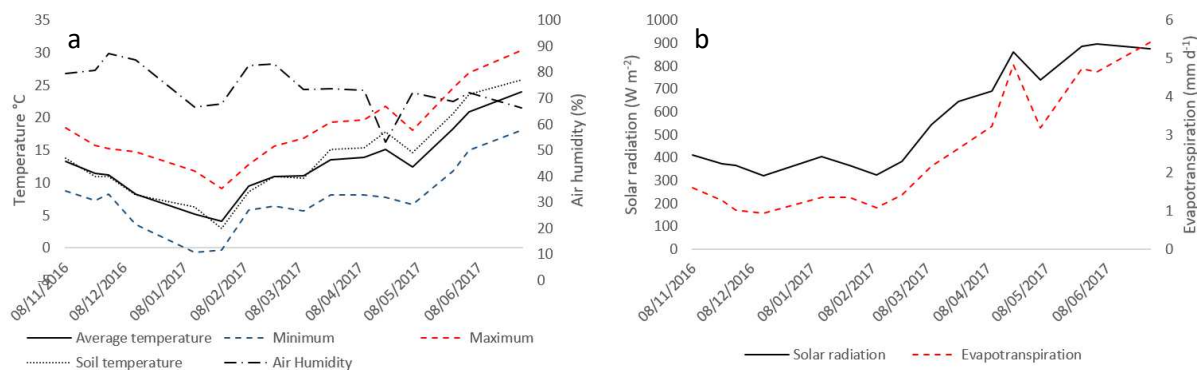


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732

733 **Figure S3 :** Climatic conditions over the 37 weeks of experimentation. a) mean air temperature (black solid line),
 734 minimum air temperature (blue segmented line), maximum air temperature (red segmented line), soil temperature
 735 (black dotted line) and air humidity (black segmented-dotted line). In b), mean solar irradiation (black solid line) and
 736 evapotranspiration (red segmented line).

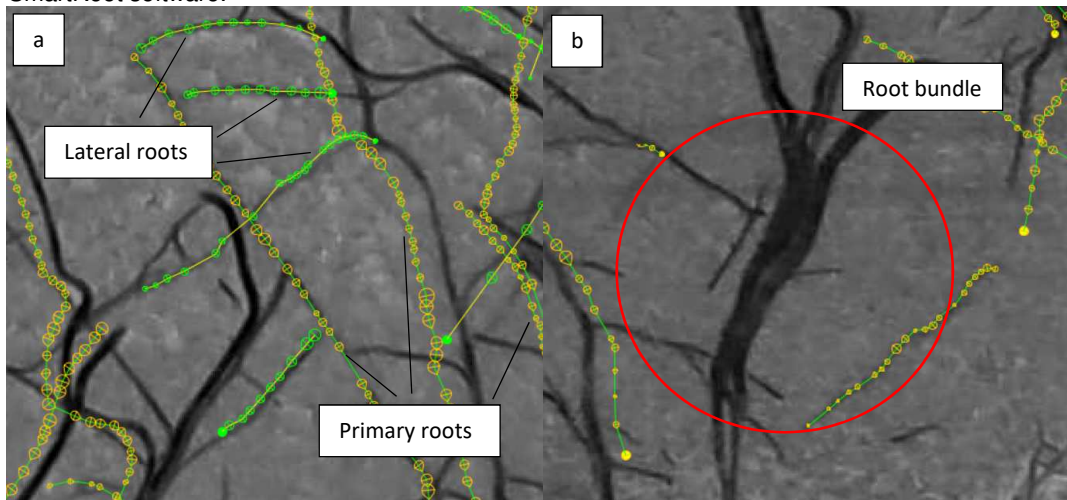
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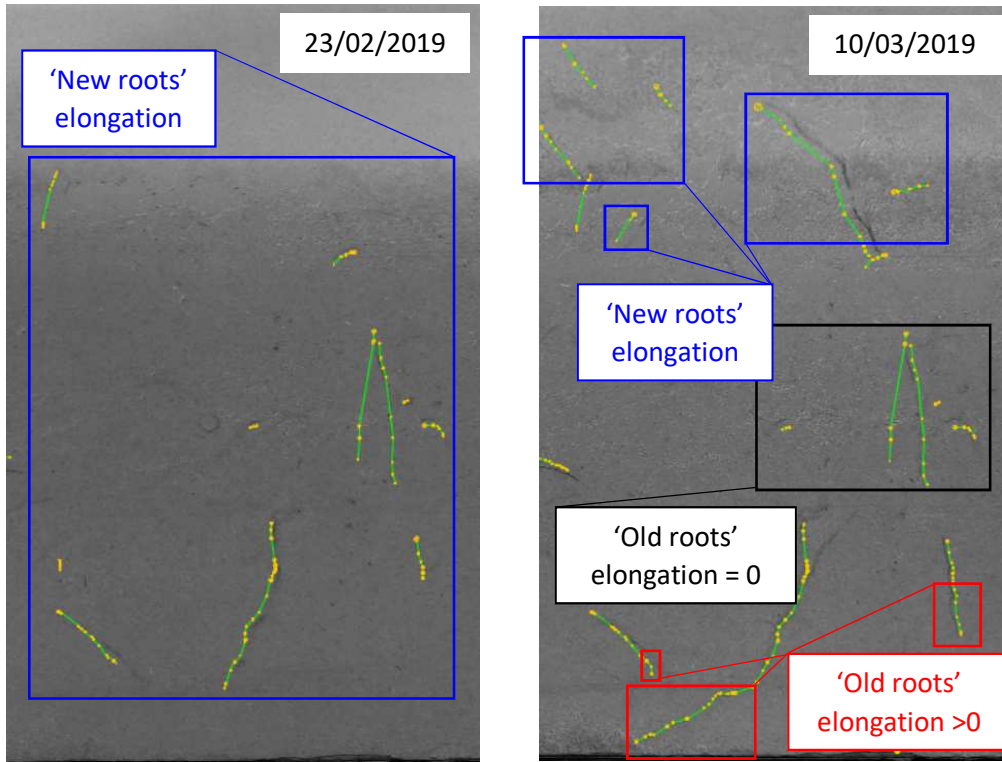
740 **Figure S4:** Example of an image analyzed using SmartRoot software. a) primary roots (in orange) from which
741 lateral roots are initiated (in green). Each orange circle along the root axis represent a single 'mouse click'
742 selection. b) a bundle of roots growing close together, where the number and diameter of roots cannot be
743 recognized by the SmartRoot software.



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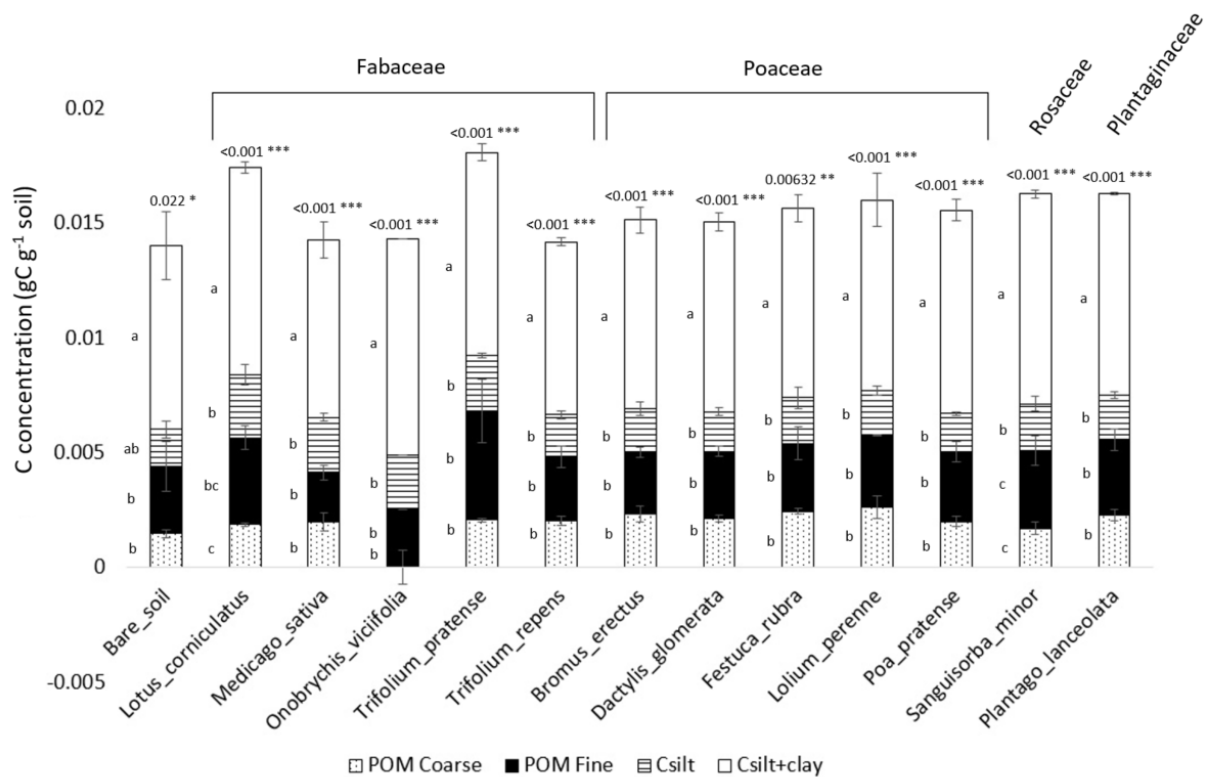
745

746 **Figure S5:** Example of two subsequent images of roots of *B. erectus* taken on 23/02/2019 and 10/03/2019. The
747 figure on the left show newly initiated roots that will be analyzed to calculate the RER_{NEW} and RLP_{NEW} on
748 23/02/2019. At the next date for image analysis (10/03/2019), some of the previously analyzed roots remained the
749 same length ($RER=0$, middle of the rhizotron), whereas other elongated ($RER>0$, bottom of the rhizotron), and were
750 used to calculate RER_{OLD} and RLP_{OLD} . On the top part of the rhizotron, some new roots were initiated, and analyzed
751 to calculate the RER_{NEW} and RLP_{NEW} on 10/03/2019.



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753

754 **Figure S6:** Carbon (C) content in each soil C pool 37 weeks after sowing and beneath 12 species and the control
 755 bare soil. The C pools analyzed are C_{POM} in the coarse particulate organic matter $>200\mu\text{m}$), C_{finePOM} (C in the fine
 756 particulate organic matter $200\text{-}50\mu\text{m}$), C_{SILT} (in the coarse silt fraction $50\text{-}20\mu\text{m}$) and $C_{\text{SILT+CLAY}}$ (C in the fine silt +
 757 clay fraction $<20\mu\text{m}$). The letters on the left hand of the fraction bars indicate significant differences (Tukey HSD, P
 758 < 0.05) between C pools within species.

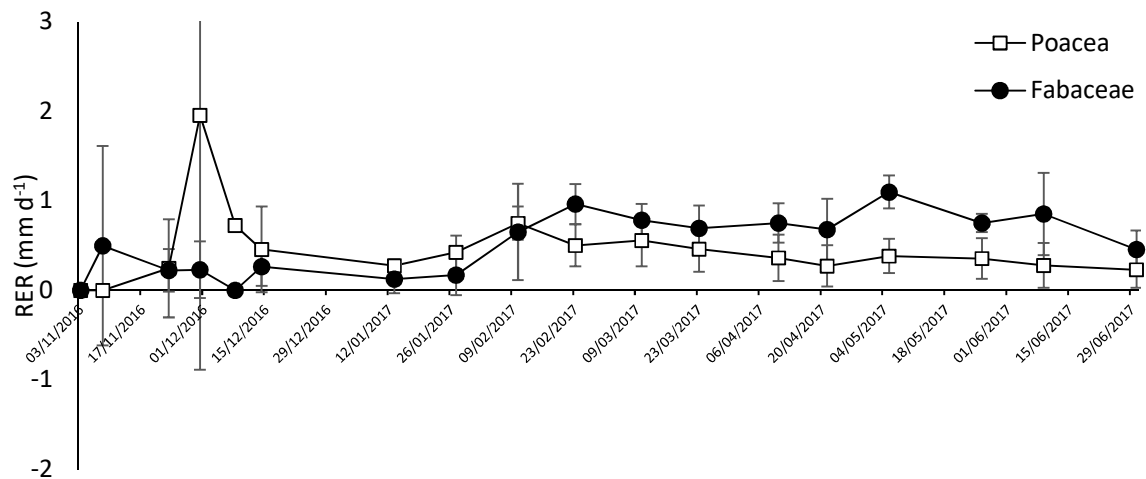


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761 **Fig. S7:** Mean daily root elongation rate (RER_{TOT}) for all the roots analyzed in the rhizotrons (without distinguishing
 762 between old and new roots) in Fabaceae (N₂-fixing, black circles) and Poaceae (N₂-fixing, white squares) family. Mean
 763 daily RER_{TOT} in Fabaceae peaked in May - June, whereas in Poaceae, mean daily RER_{TOT} was fairly constant between
 764 February and June, with no marked peaks. Data are means \pm standard error of the RER data in the 2 weeks prior to
 765 the measurement of root elongation.

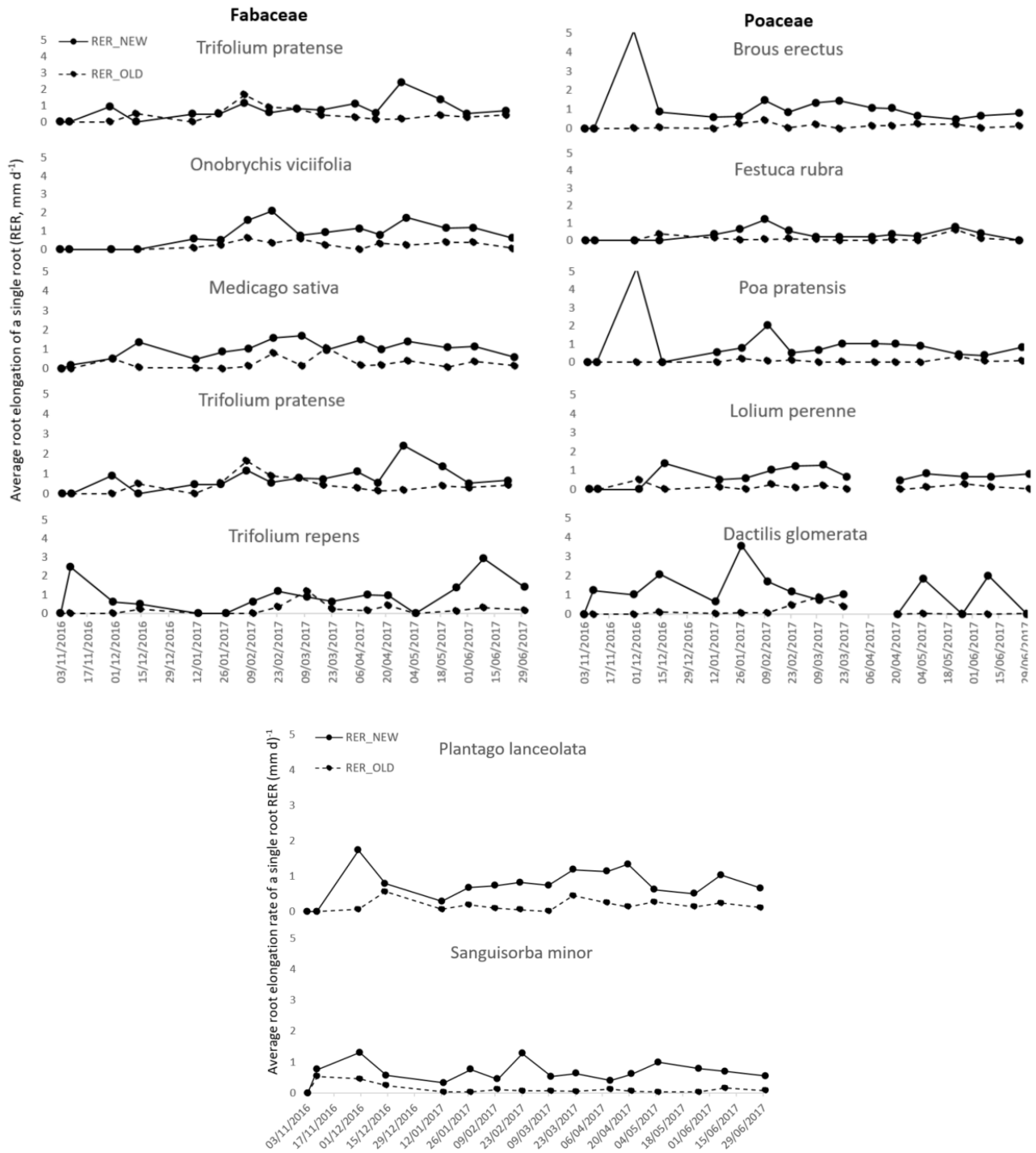
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769 **Figure S8:** Mean root elongation rate of individual roots (RER, measured every two weeks) for each of the 12 specie.
 770 The solid black line is the RER_{NEW} of the roots that were newly initiated (aged 1 to 14 days), and were not present at
 771 the previous sampling date. The dotted black line represents the RER_{OLD} of the roots that were already present at the
 772 previous sampling date, and so were older than 14 days.



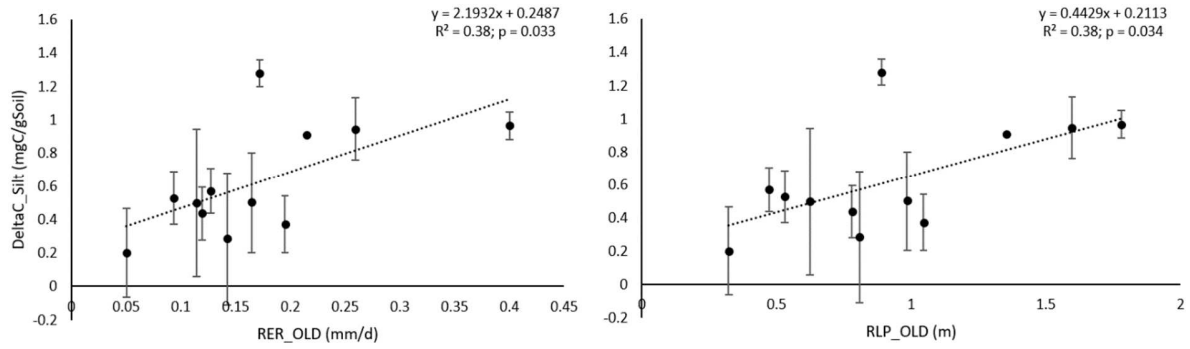
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777 **Fig. S9:** Significant, positive linear regression relationships of ΔC_{SILT} and root dynamics. (a) mean daily root elongation
778 rate of old roots (RER_{OLD} , older than 14 days), and (b) root length production of old roots (RLP_{OLD} , older than 14 days)
779 and the increase in carbon in the silt fraction (ΔC_{SILT}) after 37 weeks, for all species combined. Data are means \pm
780 standard error.



781

782 **Methods S.1:** Additional information on experimental setup. Five layers of soil were successively added
783 and therefore soil in all boxes was compacted similarly. All the soil layers came from the same
784 homogeneous mixture of soil excavated in Villefort, France (44°26'25" N, 3°55'58" E). Boxes were laid
785 out in three blocks, each block consisted of two rows of adjacent boxes at a distance of 0.5m between
786 each box (Figure S1). Each box was netted to avoid birds disturbing seeds. As soil was poor in terms of
787 N, it was inoculated with a solution of local Rhizobium bacteria strains (Incolum Valorhiz™, France)
788 once seeds had germinated. This is a technique typically used on roadsides in Southern France to
789 enhance seedling survival of N₂-fixing species, and comprises a collection of local Rhizobium strains
790 and their purification in the laboratory. Half of the boxes were also equipped with soil moisture sensors
791 (Waterscout SM100, Spectrum Technologies Inc.) and a datalogger (WatchDog weather station 200
792 series, Spectrum Technologies Inc.) that recorded soil humidity at a depth of 100mm every hour for
793 the duration of the experiment. In the same boxes, one i-button (iButtonLink, Wisconsin, USA) was
794 placed at a depth of 100 mm, to monitor soil temperature every 4 hours. A weather station is set up
795 permanently in the experiment garden, and air humidity, air temperature (minimum, maximum and
796 mean daily) and solar irradiation (measured daily) were monitored throughout the experiment. Mean
797 air temperature over the 37 weeks of the experiment was 13 °C, with a maximum of 30 °C and a
798 minimum of -0.4 °C (Figure S4a). Soil temperature in the boxes followed closely the air temperature
799 over the 37 weeks period, with a mean of 13.5 °C, a maximum of 25.9 °C and a minimum of 3 °C (Figure
800 S4a). Air humidity ranged from 53 – 87%, with a mean value of 74% (Figure S4a) and mean solar
801 irradiation ranged from 320 - 897 W m⁻² with a mean value of 568 W m⁻² (Figure S4b). During the
802 experiment, plants were cut to ground level every 4 months to simulate the management of vegetation
803 on local road embankments and maintain a regular plant density.

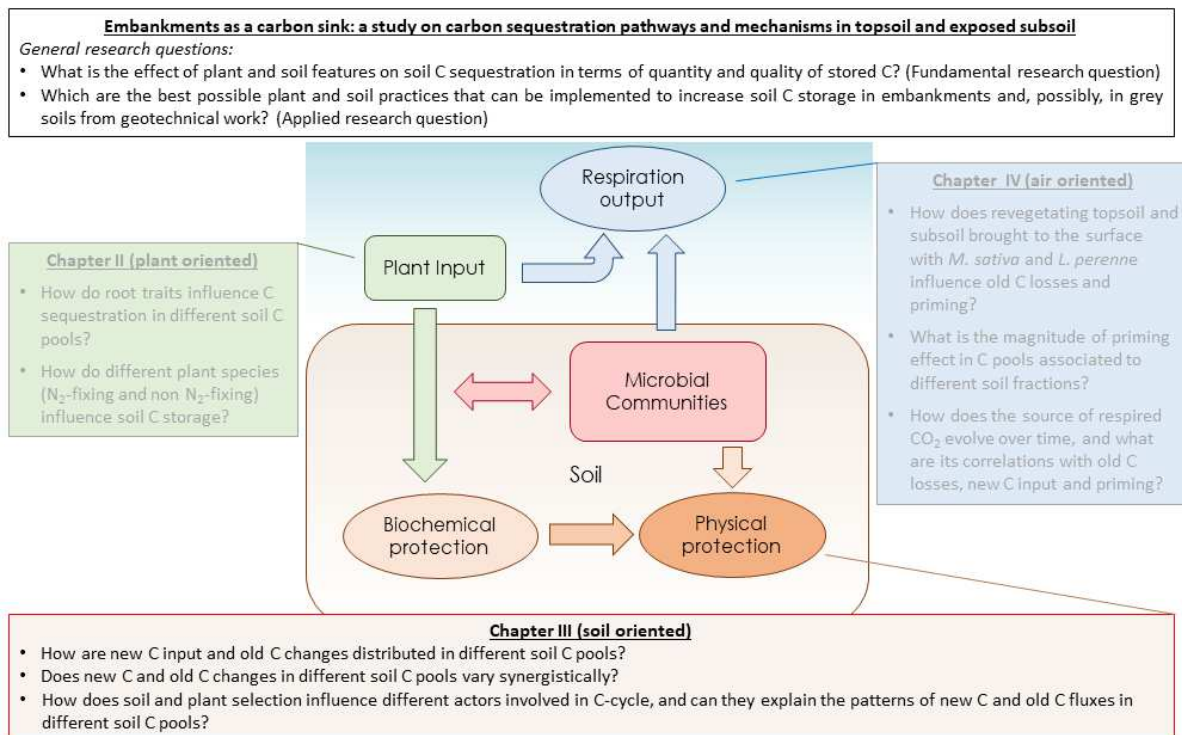
804 **Methods S2:** Analysis of carbon content in different soil fractions. Soil samples were presoaked
805 overnight in 300 ml of deionized water at 4°C with 0.5 g of hexametaphosphate to enhance
806 disaggregation. Soil was then shaken at 300 rpm (Digital orbital shaker, Intertek) with five agate
807 marbles for 2 h (i.e., the time suggested for sandy soils, to avoid the transfer of C into finer fractions,
808 Gavinelli et al. 1995). The soil was wet sieved with a 200 µm sieve, and the resulting 2000-200 µm
809 fraction was then transferred into a separate container and soaked in deionized water. The floating
810 coarse particulate organic matter (POM) was then carefully collected. The remaining 2000-200 µm
811 fraction represented the coarse sand fraction in soil and was carefully collected by washing the content
812 of the sieve in a beaker using deionized water. Then, the remaining fraction was sieved with a 50 µm
813 sieve, to separate and collect the fine sand fraction and the fine POM fraction (50-200 µm). The
814 remaining fraction <50 µm was sonicated with a 1510E-MT Branson sonicator for 10 minutes to break

815 microaggregates before sieving at 20 μm . The 20-50 μm fraction (coarse silt) was collected and the
816 resulting solution of deionized water and <20 μm fraction collected in a beaker and filled up to 1.0 L.
817 This solution was tumbled 30 times to homogenize it and an aliquot of 100 mL was collected with the
818 aid of a syringe, representing the fine silt + clay fraction. All the fractions were oven dried at 40°C until
819 all the water evaporated

820

821

Chapter III: The fates of fresh new carbon and old soil carbon differ in topsoil and newly exposed subsoil and are explained by root, microbial, and soil particle size



In Chapter II we studied the effect of revegetation on C storage in different soil C pools. In Chapter III we aim to refine the understanding of C pathways in soil by selecting two species with contrasting root characteristics (among the species in Ch. II) and sowing them on two soils showing contrasting characteristics (fertile topsoil and poor subsoil) in a ^{13}C constantly enriched environment. We will differentiate the input of fresh new C and changes in preexistent old C in the different soil C pools and study their relationships with root traits, microbiological and soil characteristics.

Abbreviation	Definition	Corresponding symbol
C	Carbon; soil carbon refers to soil organic carbon in this study	
C pool	Soil carbon contents associated with different soil particle fraction sizes: particulate organic matter fraction (50-200 μm), fine particulate organic matter fraction (20-50 μm), coarse silt fraction (20-50 μm) and fine silt and clay fraction (<20 μm)	$X = \text{POM}, \text{finePOM}, \text{SILT}$ and SILT+CLAY
C content	Soil carbon concentration (in $\text{mgC g}^{-1}\text{soil}$) per unit weight of soil for each C pool or pool summed	$C_{\text{POM}}, C_{\text{finePOM}}, C_{\text{SILT}}, C_{\text{SILT+CLAY}}, C_{\text{SUM}}$
C change	Difference in soil carbon contents (in $\text{mgC g}^{-1}\text{soil}$) between the end and beginning of the experiment	$\Delta C_{\text{POM}}, \Delta C_{\text{finePOM}}, \Delta C_{\text{SILT}}, \Delta C_{\text{SILT+CLAY}}, \Delta C_{\text{SUM}}$
C quality	Proportion (in %) of soil carbon content belonging to each carbon pool	$\%C_{\text{POM}}, \%C_{\text{finePOM}}, \%C_{\text{SILT}}, \%C_{\text{SILT+CLAY}}$
new C	Fresh soil carbon due to plant inputs	$\Delta \text{New } C_x$ (new C gain for the pool X)
old C	Existing soil carbon before plant growth	$\Delta \text{Old } C_x$ (old C change for the pool X)
t_0	Time zero, beginning of the experiment	
t_6	Time after 6 months, end of the experiment	
POM	Particulate organic matter	
C:N	Carbon – nitrogen ratio in plant tissue	
GMA	Global Metabolic Activity of microbial communities	
SIR	Substrate induced respiration	
ANOVA	Analysis of variance	
PCA	Principal component analysis	
H	Shannon metabolic diversity index	

1 3.1.INTRODUCTION

2 3.1.1. *General context in soil organic carbon sequestration on embankments: can subsoil* 3 *brought to the surface be used as a C sink?*

4 Soil is the largest terrestrial carbon (C) reservoir and soil organic C (SOC) exchanges rapidly with C in the
5 atmosphere and biosphere (Torn et al. 2009). In the context of global warming, knowing the fate of SOC is
6 essential for greenhouse gasses mitigation. So far, national and supranational programs have been
7 developed to maintain soil organic C stability and promote C sequestration in soil (e.g. 4p1000).
8 Appropriate soil and vegetation management that favors C transfer from air to soil via plants has been
9 shown to be a promising way to increase the soil C sink (Rees et al., 2005; Minasny et al., 2017). Most of
10 the research have been carried on agricultural and 'natural' soils, while heavily disturbed antropized soils,
11 i.e.. soils related to geotechnical operations. In this work we focus on the revegetation of geotechnical
12 road and railroad embankments, and their potential for soil C storage. Topsoil has often been used for the
13 revegetation of embankments, however, subsoil can be brought to the surface and revegetated directly
14 (REF). We argue that revegetating subsoil brought to the surface have a high influence on the C-cycle due
15 to the different characteristics of subsoil compared to topsoil: lower fertility levels, different aggregate
16 characteristics, microbiological communities and dynamics (Taylor et al. 2002, Murray et al. 2004, Chabbi
17 et al. 2009, Jones et al. 2018) and, most notably, C saturation (Lorenz and Lal 2005; Rumpel and Kögel-
18 Knabner 2011, Beare et al. 2016). It has been hypothesized that soil has a C saturation level associated to
19 its fine particle size partition (i.e., clay content) and the initial old C content (Six et al. 2002; Stewart et al.
20 2007). Protection via organomineral interactions relay on surface area of soil particles, and after the
21 available areas and reactive surfaces are occupied by C, further C input will not be adsorbed anymore and
22 therefore will not be protected (Six et al. 2002). The potential amount of C protected via organomineral
23 complexation depend on the amount of the <20µm fraction and the initial amount of C in the associated
24 soil C pool. We argue that subsoil have a lower C saturation compared to topsoil due to higher clay content

25 and lower C content (Lorenz and Lal 2005; Rumpel and Kögel-Knabner 2011, Lawrence-Smith et al. 2015)
26 and, therefore, could store stable C more efficiently via organomineral interactions in the finer silt and
27 silt+clay soil fractions (<20µm). For this reason, subsoil brought to the surface could be an interesting
28 feature for C storage, and we aim to investigate the influence of excavating and revegetating subsoil on
29 the main actors involved in C-cycle and their influence on C storage in different soil C pools associated to
30 granulometry of soil fractions.

31

32 *3.1.2. New and old carbon in soil*

33 The soil C stock within a defined time frame is the balance between input and transformation of newly
34 photosynthesized C from plants to soil (new C) and losses of existing soil organic C (old C) through microbial
35 respiration (Kuzyakov and Domansky, 2000; Fontaine et al., 2004). Although total soil C sequestration is
36 increasingly measured as an important ecosystem service, few studies have quantified the proportions of
37 new C input from plants and the losses of old soil C during respiration. It is unclear how the input of new
38 C and the losses of old C participate to the final soil C sequestration and if trade off or synergetic patterns
39 exist between new C input in soil and old C losses. The underlying mechanisms behind these processes are
40 poorly understood, but are crucial if we wish to improve soil C sequestration.

41

42 *3.1.3. Soil organic carbon quality: carbon pools are associated to different soil granular* 43 *fractions*

44 More and more studies have highlighted the importance of C quality in soil (Chapter II, this thesis;
45 Cardinael et al.,2015). High quality soil C refers to organic C compounds that have long mean residence
46 time and good stability against mineralization because of their physical or physiochemical associations
47 with soil particles. Characterizing absolute and relative sizes of soil C pools associated to soil particle size
48 fractions is a powerful instrument to evaluate soil C quality. The commonly used classification of soil pools

49 in literature refers to (i) C in particulate organic matter (POM) (C_{POM} , 2000-200 μm), (ii) C in fine POM
50 ($C_{finePOM}$, 200-50 μm), (iii) C in coarse silt (C_{SILT} , 50-20 μm) and (vi) C in fine silt+clay ($C_{SILT+CLAY}$, <20 μm). In the
51 POM and finePOM pools, C is usually supposed originate from plant litter debris at different levels of
52 degradation, and is more exposed to decomposers (Kögel-Knabner, 2002), whereas C in the SILT and
53 SILT+CLAY pools are considered more stable due to their organomineral binding with fine soil particles
54 (Sollins et al., 1996; von Lützow et al., 2006; Cotrufo et al. 2013). Although recent studies have quantified
55 soil C in different pools instead of that in total soil C (Cardinael et al., 2015; Chapter II, this thesis), no study
56 to our knowledge has bridged the link between C pools and the fates of new C and old C. Speculating such
57 an association is reasonable, as fates of new C and old C should have different sensitivities to fresh plant
58 C inputs, that has been shown to have significant impact on the relative size of soil C pools, i.e., soil C
59 quality (Cardinael et al., 2015). To differentiate the inputs of new C and the changes of old C in different C
60 pools, stable isotopic labelling have proven to be a powerful methodology. Growing plants in an
61 atmosphere with increased % of ^{13}C in the CO_2 allows to differentiate the new C inputted in soil from the
62 preexistent old C (Staddon 2004). Being able to differentiate old C is very interesting, since it allow to
63 quantify even the changes in the old C pool in a determined timeframe, other than the input of new C.

64

65 *3.1.4. New old carbon distribution in different soil pools: drivers and mechanisms*

66 Besides understanding the fates of new and old soil C in different soil fractions, we also need to determine
67 how plant and soil characteristics affect the trajectory of new and old C. The dynamics of new C in soil is
68 assumed to be jointly determined by plant performance and soil C storage capacity. Plants transform
69 atmospheric C via litter decomposition or root exudation, therefore, traits related to decomposition and
70 exudation should be examined in soil C sequestration studies (De Deyn et al., 2008; Roumet et al. 2016;
71 Henneron et al., 2019). Roumet et al. (2016) suggested that species with contrasted growth strategies and

72 tissue quality, i.e., N₂-fixing fast-growing species with a low tissue carbon:nitrogen (C:N) ratio versus non
73 N₂-fixing slow-growing species with a high tissue C:N ratio, results in contrasted soil C sequestration.
74 However, the relative importance of such traits on soil C sequestration is not yet known. In this regard,
75 studying the effects of N₂-fixing and non N₂-fixing species is of particular interest, since they are placed at
76 the two opposite ends of the root economic spectrum (Roumet et al. 2016, Rossi et al., submitted, Chapter
77 II) and expected to significantly influence the C sequestration quantity and quality in different soil
78 fractions. The capacity of soil to influence new C storage is related, as already mentioned, to its C
79 saturation levels. New C input increase soil aggregation that, with a double feedback effect, in turn protect
80 the C from microbial mineralization via physical protection in it the aggregate structure (Tisdall and Oades
81 1979,1982; Chevallier et al., 2004). Aggregate stability, as a proxy for aggregation resistance to disruption,
82 might very well be associated with C protection in soil. N levels in soil will affect the soil fertility, and
83 therefore plant development and microbial biomass and activity (Sarker et al., 2017).

84 Studies over the last 20 years have greatly focused on the priming effect, i.e., the phenomenon that fresh
85 biomass may, in most cases, stimulate microbial activities and thus accelerate the loss of old C existing in
86 soil (i.e., positive priming) (Kuzyakov et al. 2000; Blagodatskaya & Kuzyakov, 2008). Positive priming can
87 offset the gain of new C in soil and result in a net negative C balance (Cheng et al., 2003, Fontaine et al.,
88 2004). Many factors, from fresh tissue recalcitrance to soil physical properties are found to influence the
89 loss of old C. In particular, fresh tissue recalcitrance greatly affects the proliferation rate of the microbial
90 population and subsequent soil respiration rate. However, soil aggregate and particle size determine the
91 ability of soil to protect old C from microbial mineralization (Six et al., 2002). For example, aggregates act
92 as a physical barrier that separate occluded C from microbes and enzymes (Besnard *et al.*, 1996; Rasse *et*
93 *al.*, 2005; Bardgett *et al.*, 2014; Sokol *et al.*, 2019

94 I ask therefore, if plant traits, and soil characteristics (as aggregate stability, N content, and particle size
95 fractions) alter new C and old C dynamics in different soil fractions through their direct effect and indirect
96 influence on microbial communities. Microbial abundance (in terms of biomass that can be calculated as
97 concentration of DNA in soil), the global metabolic activity as the amount of respired CO₂ per g of soil in a
98 specific timeframe (GMA), and the diversity in metabolic substrate consumption, often represented by the
99 Shannon index (H), will be deeply influenced by soil (Liang et al. 2017) and plant species (Cotrufo et al.
100 2013). Microbial communities are have a pivotal role determining the mineralization and losses of old and
101 new C, its subsequent transformation in degraded POM or the C input in the protected silt and silt+clay
102 fractions.

103

104 **3.1.5. Research hypothesis**

105 Using a microcosm experiment, coupled with stable isotopic (¹³C) labelling, we aim at characterizing the
106 fates of new C from plant roots (root debris and exudations) and old C (pre-existing C in soil), as well as
107 their interdependence, across different soil fractions under a fully crossed soil and vegetation treatment:
108 two types of natural soils (subsoil and topsoil) × three vegetation treatments (bare soil, *Medicago sativa*
109 and *Lolium perenne*). Soil and microbial community characteristics and plant root traits were measured to
110 disentangle the effects of different drivers on changes in new and old C. I hypothesize that:

111 i) Soil particle size can regulate the fates of old and new C within fractions we hypothesize that the input
112 of new C will be higher in the particulate organic matter (via root turnover) and in the SILT+CLAY fraction
113 due to exudation and microbial in vivo transformation of C. Old C is expected to be depleted from coarser
114 fractions (POM and finePOM) via microbial mineralization, and a transferred to finer fractions, increasing
115 the old C in the SILT and SILT+CLAY fractions.

116 ii) The fates of new and old C show independent patterns: old C losses are expected to be more related to
117 microbial characteristics than to input of new C, however the influence of new C on microbial activity might
118 show an indirect effect decreasing old C concentration

119 iii) the patterns of new C and old C fluxes in different soil C pools could be explained by plant, micro-
120 organism and soil characteristics. More specifically, root traits connected to high root biomass and labile
121 input (i.e. acquisitive resource strategies N₂-fixing species) are expected to increase new C in the soil,
122 especially in the SILT and SILT+CLAY pools. We suppose subsoil to have a higher new C accumulation in the
123 SILT+CLAY fraction due to higher fine fraction and lower initial C content, decreasing soil C saturation. In
124 this respect, we think FF will be positively correlated with new C in the SILT+CLAY fraction. Aggregate
125 stability (measured as mean weight diameter, MWD) is expected to be positively correlated with new and
126 old C accumulation in silt and silt+clay fractions due to C protection. We believe soil N content will overall
127 increase the input of new C in all the fractions due to its connection with soil fertility and biomass
128 production. Microbial characteristics (GMA, H and DNA concentration) are expected to be positively
129 correlated with the accumulation of new C in the SILT and SILT+CLAY fraction and decrease the old C
130 content due to metabolism and respiration of C.

131

132 3.2. MATERIALS AND METHODS

133 3.2.1. *Soil and plant preparation, experimental design and set-up*

134 The soil used in the experiment was excavated from Pisciotta (Italy, 40°07'N, 15°14'E, 178 m a.s.l.) at two
135 depths of the same soil profile: topsoil (0.0 – 0.3 m depth) and subsoil (1.1 – 1.4 m depth). The soil was a
136 clay loam soil (USDA) with a comparable granulometric texture between topsoil and subsoil (topsoil: 27.3%
137 clay, 31.1% silt, 41.6% sand; subsoil: 34.8% clay, 36.8% silt, 28.4% sand). Topsoil (7.0) had a lower pH than
138 subsoil (8.4).

139 Both top- and subsoil were sieved to 5 mm prior and then placed in containers (20 cm x 20 cm x 20 cm),
140 where it was packed manually to a depth to 10 cm. Pots were weighed to ensure that they contained the
141 same amount of soil (+/- 2.5 %). N₂-fixing *Medicago sativa* L. and non N₂-fixing *Lolium perenne* L. were
142 planted as monocultures with exactly the same pattern. In each pot, three seeds were put at six
143 equidistant spots. After germination, one seedling was removed with scissors at ground level, at each spot.
144 For each soil type (i.e. top- and subsoil) and species, six replicate containers were prepared and six bare
145 containers per soil type were used as controls (n = 36 in total).

146 Containers were placed into three identical microcosms at the Ecotron growth facility at Montpellier,
147 France (<http://www.ecotron.cnrs.fr/>). In each microcosm, two replicates of all treatments, i.e., 12 pots,
148 were placed randomly to avoid any effect of microcosm on plant growth and soil processes. Plants were
149 grown at a constant air temperature of 21°C and at 80% humidity (to reduce the soil water loss by
150 evapotranspiration). Artificial light was provided by three lamps (Gavita PRO 300 LEP 02, Netherlands) in
151 each microcosm with a 12h day/night cycle, shifted to allow air sampling at the same moment of the
152 plant's circadian rhythm (data not shown in this study, see Chapter 4). A shade was placed on the lamps
153 and the distance of the lamps from the plants was adjusted to achieve the most possible homogenous
154 light intensity on the foliage (300 $\mu\text{mol m}^{-2} \text{s}^{-1}$). Soil moisture was kept at $45 \pm 10\%$ of the soil water holding
155 capacity for the entire duration of the experiment. To minimize disturbances due to microcosm openness,
156 a system of plastic pipes was installed into the microcosm for watering. Every two weeks, pots were taken
157 out to assess their evapotranspiration rate and weight (data not shown). Each time pot position was
158 randomized when they were put back to the microcosm. Air enriched with enriched ¹³C (with a
159 concentration of 2%, approximately two times higher than the natural ¹³C abundance of 1.1%, in other
160 words $\delta^{13}\text{C}$ of CO₂ in the chamber was roughly +760, as compared to the ambient -8) was supplied into the
161 microcosms once the first emergence of seedlings was observed in any microcosm (approximately three

162 weeks). The air enrichment with ^{13}C was supplied only during the 12h day cycle and the ^{13}C infusion was
163 stopped during the night period. The whole experiment lasted for 183 days, or six months from 29
164 September 2017, (t_0) to 31 March 2018 (t_6). Experiment length enabled us to (i) track the changes in soil
165 C immediately after seeding emergence (ii) avoid the effect of plant leaf and flower litter on soil C, which
166 was not our study objective. Any plant litter was removed manually every 2 weeks from the soil surface.

167

168 *3.2.2. Soil fractionation and assessment of soil carbon and $\delta^{13}\text{C}$*

169 Before the experiment, three soil samples per soil type were sampled for the measurement of carbon
170 content in different fractions, mean weight diameter of aggregates, nitrogen content, microbial activity,
171 DNA concentration and Shannon metabolic diversity at t_0 . Each sample was mixed and divided into four
172 parts, and an equal amount of soil from each part was collected and mixed to obtain a homogenized
173 sample of soil.

174 At t_6 , soil was removed from each pot, weighed and then cut into two equal-size half blocks with a saw
175 and a ruler: one half was air dried and used for soil analyses and the other half was used for plant trait and
176 microbiological measurements. A mixed sample from each pot was collected from a depth of 3.5-10 cm
177 depth. The soil samples at t_0 and t_6 were then sieved at 2 mm and 40g were sub-sampled and fractioned
178 using the Gavinelli et al. (1995) method (Supplementary material, Method S1, Fig. S1). The resulting four
179 fractions (POM: $>200\mu\text{m}$, finePOM: $200\text{-}50\mu\text{m}$, SILT: $50\text{-}20\mu\text{m}$, SILT+CLAY: $<20\mu\text{m}$) were analyzed for both
180 C content and $\delta^{13}\text{C}$ using an elemental analyzer Isoprime100 coupled with an Elementar Varo Isotope Cube
181 (machine reference no). The sum of C in different fractions represents the total C in the sample. A
182 subsample of 0.1 g was taken from each 40 g sample and analyzed without fractioning to determine the
183 total C in the bulk sample. We checked the accordance between the mean difference between total C in

184 bulk soil and the sum of C in the different soil fractions and the results were satisfactory (mean 93.3% of
185 recovery).

186 To assess the changes in total C in each fraction, the differences between C content at t0 and t6 were
187 assessed:

$$188 \quad \Delta C_x = \Delta C_{x,t6} - \Delta C_{x,t0} \quad [1]$$

189 Where ΔC_x is the change in C content (mg C g⁻¹ soil) in a given soil C pool.

190

191 *3.2.3. Estimation of new and old carbon in soil fractions*

192 The increased atmospheric $\delta^{13}\text{C}$ signature in the microcosm allowed a calculation of the proportions of
193 new C in the different soil fractions. We used an isotope mixing model (Balesdent and Mariotti, 1996):

$$194 \quad \%NewC = \frac{\delta(t6) - \delta(t0)}{\delta B - \delta(t0)} \quad [2]$$

195 Where %New C is the percentage of new C in a specific fraction, $\delta(t6)$ is the $\delta^{13}\text{C}$ signature of C measured
196 in a specific fraction at t6, $\delta(t0)$ is the $\delta^{13}\text{C}$ signature of C of a specific soil fraction t0, δB is the $\delta^{13}\text{C}$ signature
197 of the new C input in the system (in our case the signature of the absorptive and transport root biomass).

198 The δB was specific for each pot based on the analysis of the root biomass, and the mean was 615 ± 38 . The
199 choice of root biomass as the $\delta^{13}\text{C}$ reference for C input was made because root material was considered
200 to be the main input of C, given that shoot litter was negligible.

201 The new C at t0 was zero. To calculate the gain of new C (mgC g⁻¹ soil) in a specific soil C pool X, we
202 multiplied %Cnew by the total amount of C at t6 (C_x) of the pool X:

$$203 \quad \Delta NewC_x = C_x(t6) \times \%NewC \quad [3]$$

204 To assess the changes in the old C in different soil C pools, we subtracted the new C from the ΔC of each
205 soil C pool.

$$206 \quad \Delta OldC_x = \Delta C_x - \Delta NewC_x \quad [4]$$

207

208 **3.2.4. Microbial global metabolic activity (GAM) and Shannon metabolic diversity index (H)**

209 To analyze functional diversity, precisely GAM and H from microbial communities, 20 g of soil were
210 collected immediately after sampling from each container at t_6 from the half of soil collected for chemical
211 analyses. We used a Microresp system that comprises a Deepwell plate (Fisher Scientific E39199) holding
212 soil subsamples saturated with a solution with different substrates, a detection plate containing the
213 detection gel, a rubber seal to connect the deepwell and the detection plate and metal clamp to keep the
214 two parts tightly together (MicroResp™, Aberdeen, UK) (Fig. S2). The output of Microresp is to assess the
215 respiration rate of soil saturated (at 80% of field capacity) with different substrates presenting different
216 levels of recalcitrance and biological properties. Detailed methodology is provided in supplementary
217 materials, Method S2. Substrates utilized for MicroResp are shown in Table S1.

218 To have a proxy of the global metabolic activity (GAM) of the microbial communities, the respiration rates
219 from the different 15 C substrates were summed ($\text{mg C-CO}_2 \text{ g}^{-1} \text{ soil h}^{-1}$; Frac et al. 2012, Ammar et al. 2017).
220 For each replicate, a Shannon metabolic diversity index was calculated as:

$$221 \quad H = -\sum_{i=1}^{15} \pi \times \log(\pi) \quad [5]$$

222 Where π is the standardized respiration rate for substrate (i) (Shinan et al. 2017).

223

224 **3.2.5. Microbial DNA concentration as proxy for microbial biomass**

225 To examine microbial biomass in different communities, 10g of soil was immediately frozen at -20°C after
226 sampling until samples were processed for DNA extraction. Total DNA was extracted from soil (0.5 g). DNA
227 extraction was performed using FastDNA® SPIN Kit for Soil Isolation Kit according to manufacturer's
228 instructions (MP Biomedicals, USA). An additional step to wash the DNA binding matrix with 500 µl of
229 guanidine thiocyanate 5.5M was added following Tournier et al. (2015). The concentration of extracted
230 DNA in solution (ng/µL) were measured using a Quant-iT™ PicoGreen™ dsDNA Assay Kit for DNA
231 quantification and used as a proxy for microbial biomass (Bohórquez et al. 2017).

232

233 **3.2.6. Percentage of fine fraction in soil, soil nitrogen and aggregate stability**

234 After the wet sieving and weighing of the different soil fractions, the percentage of fine fraction (FF, in %)
235 was determined as the ratio of the SILT+CLAY soil fraction weight (<20 µm) and the total mass of the
236 fractioned soil sample (in average 40g). When analyzing C content and $\delta^{13}\text{C}$ for each the bulk soil fraction,
237 the amount of nitrogen in soil (N; mg g⁻¹ soil) was also determined.

238 As a proxy for aggregate stability, mean weight diameter (MWD) of aggregates was assessed following the
239 conventional methodology according to Le Bissonnais et al. (2006). 20g of aggregates were collected from
240 the half pot for soil analysis, air dried and sieved first at 5 mm and after at 3 mm, to isolate the 3 - 5mm
241 aggregates fraction. Aggregates were put in the oven for 24h to reach the same water matrix potential.
242 First, 5g of 3-5mm fraction are weighed and gently immerse in a 250 cm³ beaker filled with 50 cm³ of
243 ethanol for 10 minutes. After ethanol was carefully sucked off with a pipette and the sample transferred
244 in a 250 cm³ Erlenmeyer flask containing 50 cm³ of deionized water, and brought to 200 cm³. The flask was
245 agitated 20 times and left 30 minutes for sedimentation of coarse particles. Water was sucked off with a
246 pipette and the mixture of soil and water transferred to a 50 µm sieve previously immersed in ethanol.

247 The disaggregated soil was sieve gently by moving 5 times the sieve in the ethanol to separate the <50 µm
248 soil fraction from the >50 µm. >50 µm aggregate fraction was collected from the 50-µm sieve, oven-dried
249 and gently dry-sieved by hand on a column of six sieves: 2000, 1000, 500, 200, 100 and 50 µm. The mass
250 percentage of each aggregate fraction was calculated, and subtracting the mass of soil collected on sieves
251 to the initial mass analyzed mass we calculated even the <50µm fraction. MWD is calculated as the sum
252 of the mass fraction of soil remaining on each sieve after sieving multiplied by the mean aperture of the
253 adjacent mesh:

$$254 \quad MWD = \frac{\sum_{f>2mm}^{f0.05-1mm} (A_f * df)}{100} \quad [6]$$

255 Where A_f is the aggregate fraction abundance in % of the total weight of the analyzed sample remaining
256 in a specific f aggregate fraction ($f = >2\text{mm}, 1-2\text{mm}, 0.5-1\text{mm}, 0.2-0.5\text{mm}, 0.1-0.2\text{mm},$ and $0.05-0.01\text{mm}$),
257 and df the diameter in mm of the smaller sieve characterizing the lower boundary of the f aggregate
258 fraction.

259

260 **3.2.7. Root traits**

261 For each species, three out of the six plants in each pot were sampled and their root system carefully
262 washed and separated from the aboveground part. Roots were water-rinsed in a plate container. A 0.1 g
263 composite subsample of roots was cut off from several parts of the root systems with scissors. After being
264 carefully washed, they were stained with a solution of methyl violet (0.5 g L^{-1}). Following McCormack *et*
265 *al.* (2015), we visually separated transporting (long, thick, high-order roots (>3) and absorptive roots
266 (short, thin, low-order roots 1 – 3). Both types of roots were separately extended over a transparent water
267 filled tray and scanned at 800 dpi (Epson® perfection V700 PHOTO, Canada). The software Winrhizo Pro®
268 (Regent Instruments, Quebec, Canada) was used to determine root length (L, in mm) per diameter class

269 stretching from 0 to 2 mm with a 0.1 mm interval. Roots were then oven dried at 60°C for three days and
270 weighed to determine the total root dry mass (in g) for each pot. Winrhizo Pro® also provides the mean
271 diameter of the analyzed root sample (Prieto et al., 2016). We then calculated the mean diameter of
272 adsorptive roots (D_{ad}) as the mean diameter between each replicate container, for in each soil and species
273 type.

274 Specific root length (SRL, $m\ g^{-1}$) was calculated as the ratio between root length and dry mass (Esseinstat,
275 1992). Only absorptive root data were then used in data analysis as they are most short-lived and active
276 in exudation, thus should be the main contributor of new C deposition to soil.

277 Following the same sampling and sorting manner, another sample of 0.1 g absorptive roots was obtained
278 and then finely ground. The ground samples were analyzed with an elemental analyzer Isoprime100
279 coupled with an Elementar Varo Isotope Cube to determine root C and N contents and root $\delta^{13}C$.

280 Finally the amount of new C moved into the SILT+CLAY pool by g of root was calculated by standardizing
281 the $\Delta New\ C_{SILT+CLAY}$ for the g of dry root weight per gram of soil (DRW; $g\ dry\ roots\ g^{-1}\ soil$). Also the amount
282 of new C moved into the SILT+CLAY pool by cm of root was calculated by standardizing the $\Delta New\ C_{SILT+CLAY}$
283 for the length of the root (L, $cm\ of\ roots\ g^{-1}\ soil$) per gram of soil. To calculate the root L, we multiplied SRL
284 per the DRW.

285

286 **3.2.8. Statistical analysis**

287 One way analysis of variance (ANOVA) was performed to test the effects of plant and soil treatments on

288 1) C content: ΔC_x , $\Delta New\ C_x$ and $\Delta Old\ C_x$ for the C pool X and all pool summed (*SUM*)

289 2) C quality: $\%C_x$ each C pool X

290 3) Soil, root and microorganism feature indicators, including initial SOC stock, FF, soil N content,
291 MWD, total root biomass, mean absorptive diameter, C:N ratio of absorptive roots, SRL of
292 absorptive roots, concentration of extracted DNA in ml⁻¹ of soil solution as proxy for microbial
293 biomass, microbial activity (GMA) and metabolic community diversity (H) .

294 The normality of distribution of residues was verified using a Shapiro-Wilk test ($\alpha_p = 0.05$). Principal
295 component analysis (PCA) and Pearson's correlations factors were used to study the relationships between
296 C sequestration indicators and soil type, root and microbial indicators. All the statistical analyses were
297 performed using the open-source statistical environment 'R', version 3.4.3 (R Development Core Team,
298 2017) using the packages *vegan* and *Hmisc*. (Oksanen et al. 2019, Harrel 2007).

299

300 3.3.RESULTS

301 3.3.1. Changes in total soil carbon

302 In general, a decrease in mean total soil C content occurred after 6 months in subsoils (Fig. 1a,b). The mean
303 negative ΔC in bare subsoil (-0.37 ± 0.18 mgC g⁻¹ soil) was not significantly different to that with either *L.*
304 *perenne* (-0.38 ± 0.11 mgC g⁻¹ soil) or *M. sativa* (-0.17 ± 0.25 mgC g⁻¹ soil) present. Although the presence of
305 vegetation did not significantly affect total C in subsoil (ANOVA, $p \geq 0.05$) (Figure 4.1), mean ΔC had a
306 significant increase in topsoil (ANOVA, $p < 0.05$), with the highest increase in C content under *M. sativa*
307 ($+0.68 \pm 0.36$ mgC g⁻¹ soil), followed by *L. perenne* ($+0.1 \pm 0.51$ mgC g⁻¹ soil). In bare soil, ΔC w (-0.47 ± 0.28
308 mgC g⁻¹ soil) (Figure 4.1a,b). The effect of soil type on ΔC_{SUM} was significant under *M. sativa* (ANOVA, p
309 < 0.05), but not under *L. perenne* (ANOVA, $p \geq 0.05$) due to the high variability in data. There was no effect
310 of soil type on ΔC_{SUM} in bare soil (ANOVA, $p \geq 0.05$) (Fig. 1b).

311 Mean old C decreased in all treatments (Fig. 1a), although a slight positive $\Delta\text{Old C}$ could be occasionally
312 found in some pots (Fig. 2a). In topsoil, the increase in new C was sufficient to compensate for the loss of
313 old C, but it was not the case in subsoil, where $\Delta\text{New C}$ was less than the $\Delta\text{Old C}$ (Figure 4.1a). As expected,
314 bare soil had a negligible input of new C, while topsoil under *M. sativa* had both the smallest lower negative
315 $\Delta\text{Old C}$ and the highest positive $\Delta\text{New C}$. Data were highly variable with regard to negative $\Delta\text{Old C}$ in topsoil
316 under *L. perenne* (Fig. 2a). The effect of species, $\Delta\text{Old C}$ and $\Delta\text{New C}$ were less pronounced in subsoil than
317 in topsoil. In vegetated soils, positive $\Delta\text{New C}$ was accompanied by a smaller loss of old C (lower $\Delta\text{Old C}$)
318 (Fig. 2a). However, there were no consistent relationships between changes in old C and in new C in either
319 soil type or plant species (Fig. 2a).

320 Over 6 months, the amount of active C (i.e., $|\Delta C_{\text{New}}| + |\Delta C_{\text{Old}}|$) took 9.1% and 6.1% of the total C contents
321 for subsoil and topsoil, respectively. The amount of active C in topsoil was 1.5 times higher than that in
322 subsoil.

323

324 3.3.2. Changes in soil carbon in different soil C pools associated to soil fractions

325 In subsoil, $\Delta C_{\text{finePOM}}$ was significantly lower than in the other pools, (ANOVA, $p < 0.05$). The ΔC between
326 pools was not significant for *M. sativa* and bare soil due to the high variability in data (Figure 4.3a). In
327 topsoil, mean $\Delta C_{\text{finePOM}}$ was not significantly different with that in the ΔC_{POM} or ΔC_{SILT} . $\Delta C_{\text{SILT+CLAY}}$ was usually
328 the highest among all the four C pools (Fig. 3a,b).

329 Regardless of soil type, plant species had a limited effect on $\Delta C_{\text{finePOM}}$ and ΔC_{SILT} (Figure 4.3a, b). In subsoil,
330 plants increased ΔC_{POM} (Fig. 3a), but not in topsoil. However, in topsoil, plants increased $\Delta C_{\text{SILT+CLAY}}$
331 ($1.28 \pm 0.63 \text{ mgC g}^{-1} \text{ soil}$ for *M. sativa*, $1.00 \pm 0.44 \text{ mgC g}^{-1} \text{ soil}$ for *L. perenne*), compared to bare soil
332 ($0.90 \pm 0.29 \text{ mgC g}^{-1} \text{ soil}$). In the remaining C pools, ΔC significantly decreased (Figure 4.3b).

333 The only significant difference in soil planted with either species, was the increase in $\Delta C_{\text{SILT+CLAY}}$ in the topsoil
334 compared to subsoil (ANOVA, $p < 0.01$).

335 $\Delta \text{New } C_{\text{finePOM}}$ $\Delta \text{New } C_{\text{SILT}}$ changed negligibly with regard to soil type and plant species. Compared to bare
336 soil, new C gain in planted subsoil soil was mainly located in the POM pool, while new C gain in planted
337 topsoil was located in both POM and CLAY+SILT C pools (Fig. 4a,c). Soil under *M. sativa* had significantly
338 higher $\Delta \text{New } C_{\text{POM}}$ and $\Delta \text{New } C_{\text{SILT+CLAY}}$ pools compared to *L. perenne* (Fig. 4a,c).

339 The effect of soil type on old C was more accentuated compared to New C. In subsoil, neither C pool and
340 plant species had no effect on old C loss, which was always negative (C depletion). In topsoil, instead, here
341 was a positive accumulation for $\Delta \text{Old } C_{\text{SILT+CLAY}}$, while decreased in all the other pools that did not differ
342 among each other (Fig. 4b,d). Plant species had no effect on old C loss in any C pool (ANOVA, $p \geq 0.05$).

343

344 3.3.3. Changes in carbon quality

345 *M. sativa* increases the % of C stored in the POM C pool over the total amount of C in the soil thanks to
346 high input of new c, while *L. perenne* and bare soil decrease the % of C in this pool (Fig. 5a). The % of C
347 stored in the finePOM and SILT pools decrease over the 6, with negligible input of new C (Fig. 5b,c). Finally,
348 every treatment increase the % of C stored in the SILT+CLAY pool over the 6 months every treatment
349 increase the C % compared to t0 (Fig. 5d).

350

351 3.3.4. Root, soil and microbial characteristics

352 Root biomass of both plant species was significantly higher in topsoil (*M. sativa* $17.53 \pm 2.03 \text{ g pot}^{-1}$, *L.*
353 *perenne* $4.09 \pm 1.43 \text{ g pot}^{-1}$) than in subsoil (*M. sativa* $5.13 \pm 1.36 \text{ g pot}^{-1}$, *L. perenne* $1.05 \pm 1.36 \text{ g pot}^{-1}$) (Tables
354 1, 2). In *M. sativa* soil type did not influence mean SRL, diameter or C:N ratio of absorptive roots. In *L.*
355 *perenne*, greater SRL (topsoil $+11.97 \text{ m g}^{-1}$) and C:N ratio ($+39.30$) in subsoil was found compared to topsoil

356 (Tables 1, 2). In *L. perenne*, absorptive roots were thinner in subsoil compared to topsoil (-0.01 mm; Tables
357 1, 2). *M. sativa* had greater root biomass (topsoil +13.44 g pot⁻¹, subsoil +4.08 g pot⁻¹) and mean diameter,
358 but lower SRL and C:N ratio than *L. perenne*, in both soils (Tables 1, 2). In topsoil, GMA, H and DNA mass
359 were all significantly greater compared to subsoil once plants had grown. In bare soil, GMA did not
360 significantly differ among topsoil and subsoil (Tables 4.1, 2). In topsoil, the presence of both species
361 significantly increased GMA (*M. sativa* +10.39±4.63 µgC-CO₂ g⁻¹ soil h⁻¹, *L. perenne* +3.01±2.19 µgC-CO₂ g⁻¹
362 soil h⁻¹) and DNA concentration (*M. sativa* +5.33±5.29 µgC-CO₂ g⁻¹ soil h⁻¹, *L. perenne* +4.73±6.37 µgC-CO₂
363 g⁻¹ soil h⁻¹) that did not differ between treatments (Tables 1, 2). In subsoil, GMA decreased over the 6
364 months (*M. sativa* -1.59±1.53 µgC-CO₂ g⁻¹ soil h⁻¹, *L. perenne* -0.16±2.73 µgC-CO₂ g⁻¹ soil h⁻¹) while DNA
365 concentration increased (*L. perenne* +2.00±1.41 ng µL⁻¹; *M. sativa* 4.20±1.64 ng µL⁻¹), and they did not
366 differ between plant species (Table 2). In subsoil H was significantly higher in soil planted with either
367 species compared to bare soil (Tables 1, 2).

368 Soil type significantly influenced the soil structural and characteristics and N content: topsoil had a higher
369 MWD and N content regardless of plant species (Table 1, 2). The fine fraction (FF) was higher in subsoil
370 compared to topsoil (Tables 1, 2). There was a significant increase in MWD with both the species in topsoil
371 (*M. sativa* +0.52±0.29 mm, *L. perenne* +0.62±0.20 mm), while in subsoil MWD was not significantly
372 different from t0 or between treatment (Table 1, 2). Soil N and FF were not significantly different among
373 treatments (Tables 1 and 2) and soil N was depleted during the 6 months, with subsoil showing an
374 homogeneous depletion among treatments (-0.1±0.8 mgN g⁻¹ soil) while in subsoil *L. perenne* had higher
375 decrease in soil N (-0.13±0.12 mgN g⁻¹ soil) compared to bare soil and *M. sativa*. In subsoil, no effect of
376 treatment was found in any of the soil characteristics.

377

378 **3.3.5. Relationship between changes in new C and old C and soil, microorganism and root**
379 **variables**

380 The PCA conducted on the Δ New C and Δ Old C in the different soil C pools, root traits, DNA mass, H and
381 GMA, and soil structural characteristics explained 83.9% of the total variance (Fig. 6). The first PCA axis
382 (horizontal) accounted for 63.3% of the variation. On the negative end, results were governed by the Δ New
383 C_{POM} , Δ New C_{finePOM} and Δ New $C_{\text{SILT+CLAY}}$, and Δ Old C_{finePOM} . At the positive end, results were driven by total
384 Δ Old C_{SUM} and Δ Old $C_{\text{SILT+CLAY}}$, while the remaining new C and old C pools were orthogonal and more related
385 to the second PCA axis (vertical), that accounted for 20.6%. Microbial traits (GMA, DNA and H), MWD, soil
386 N and root biomass, all went along the first axis (negative). Root traits linked with recalcitrance (C:N ratio
387 and SRL) and fine fraction percentage FF went along the 1st axis (positive) together with Δ Old C_{SUM} and
388 Δ Old $C_{\text{SILT+CLAY}}$.

389 The PCA strongly discriminated top- and subsoil at the two extreme of the first axis, with topsoil on the
390 negative end of the first axis, characterized by high GAM, H and DNA concentration, high MWD, soil N and
391 root biomass, correlated with Δ New C and Δ Old C_{finePOM} . Subsoil was on the negative end of the first axis,
392 with FF, root C:N and SRL, suggesting a loss of Δ Old C_{SUM} and Δ Old $C_{\text{SILT+CLAY}}$. Species were discriminated
393 mostly by the second axis, with *L. perenne* on the positive end of the axis together with higher C:N ratio,
394 SRL, and Δ Old C_{POM} and *M. sativa* on the negative end, with high root biomass and diameter of absorptive
395 roots, illustrating a positive Δ New C_{POM} pool and the Δ New C_{SUM} , and negative Δ Old C_{SILT} and Δ Old $C_{\text{SILT+CLAY}}$.

396 The gain in new C, regardless of total new C, or within each soil C pool, was better related to every analyzed
397 variable than the loss of old C (Table 3). The gain in Δ New C in every C pool was positively correlated with
398 microbial traits (GMA, AND, and H), except for Δ New C_{POM} and H (Table 3). Δ New C_{POM} , Δ New $C_{\text{SILT+CLAY}}$, and
399 Δ New C_{SUM} were significantly and negatively correlated with SRL and C:N ratio of absorptive roots. Apart
400 from Δ New C_{POM} , the gain in new C in every soil fraction was positively correlated with MWD and soil N
401 content (Table 3), but negatively correlated with FF. The negative Δ Old C was significantly and positively

402 correlated with H in $\Delta\text{Old } C_{\text{POM}}$ and with GMA in $\Delta\text{Old } C_{\text{finePOM}}$, but was negatively correlated with every
403 microbial trait (GAM, DNA concentration, H) in $\Delta\text{Old } C_{\text{SILT+CLAY}}$ and $\Delta\text{Old } C_{\text{SUM}}$. Root variables were poor
404 predictors of total old C losses, except for root biomass, which was positively correlated with the loss in
405 $\Delta\text{Old } C_{\text{finePOM}}$ and negatively with that in $\Delta\text{Old } C_{\text{SILT+CLAY}}$. Soil variables were all correlated with the losses in
406 $\Delta\text{Old } C_{\text{SILT+CLAY}}$ and $\Delta\text{Old } C_{\text{SUM}}$ demonstrating an opposing pattern compared with correlations with the gain
407 in new C. The losses in $\Delta\text{Old } C_{\text{finePOM}}$ and in $\Delta\text{Old } C_{\text{SILT}}$ were poorly correlated with most of the variables.
408 Finally, when standardized by dry root weight, $\Delta\text{New } C_{\text{SILT+CLAY}}$ in subsoil planted with either *M. sativa* and
409 *L. perenne* was significantly than that found in topsoil, but the difference was not significant (Fig. S5a;
410 ANOVA, $p > 0.05$). When the $\Delta\text{New } C_{\text{SILT+CLAY}}$ was standardized for every cm of root, no differences could
411 be observed between top- and subsoil planted with either species (Fig. S5b; ANOVA, $p > 0.05$).

412

413 3.4. DISCUSSION

414 Soil had the highest influence on C sequestration. New C accumulation, old C and total C changes in terms
415 of both absolute (C quantity) and relative (C quality) values significantly differ among soil C pools, thus
416 validating our Hypothesis 1. The most reactive pools were POM and SILT+CLAY for new C accumulation.
417 We examined the correlations between new C and old C for total soil C and each C pool and found
418 synergetic patterns in a generally consistent manner, thus rejecting our Hypothesis 2. Finally, we showed
419 that new C and old C changes could be partially explained by multiple soil, microorganism and root
420 variables despite their disparities in drivers, validating our Hypothesis 3. In general, the main drivers for C
421 storage were N content and microbial activity, which influenced soil quality. Biomass development was
422 the third driver showing high correlations with new C storage in soil fractions, but subdued to soil
423 characteristics. We did not observe a positive effect of lower C saturation on C storage in SILT+CLAY in

424 subsoil, due to the lower biomass development and microbial activity, resulting in a lower new C input.
425 Regarding the applied aspect, we found topsoil had relatively higher new C gain and lower old C loss
426 compared to subsoil and *M. sativa* had a better performance in gain of new C and limit of old C loss than
427 *L. perenne*, although such an effect of species was moderated by soil type. Understanding and assessing
428 the choice of plant and soil on C sequestration will help shape practical guidelines in revegetation and
429 restoration programs of geotechnical systems, notably road embankments.

430

431 *3.4.1. Importance of differentiating soil carbon origin and pools (Hypothesis 1)*

432 Here, we clearly confirm the importance of disentangling the C fates of different origins and pools. Taking
433 the vegetated topsoil as an example, we found that the increase in total C after six months was
434 mainly attributed to a high input of new C and to the high increase of new and old C in the most stable
435 SILT+CLAY pool. This result is in line with previous studies on either C origins (De Deyn et al., 2008; Cotrufo
436 et al., 2013; Liang et al., 2017; Vidal et al., 2018) or C quality by taking into account C pools (O'Brien and
437 Jastrow, 2013; Cardinael et al., 2015; Saenger et al., 2015; Chapter II this thesis).

438 In topsoil, the total old C change was close to zero, but was actually an offset between an active and high
439 gain in old C in the SILT+CLAY pool and an active and high loss in old C in the POM pool. As the fate of old
440 C was not estimated in the different C pools, it is possible to wrongly diagnose that old C was little active
441 during the whole revegetation process.

442 The correlations between total ΔC and diverse soil, root and microbial characteristics, did not reveal the
443 relationships in most of the C pools. (e.g. Cardinale et al., 2015, Rossi et al., submitted). Being able to
444 separate new C and old C fluxes thanks to isotopic enrichment have proved fundamental to investigate
445 correlations that are hidden when considering the total ΔC as the sum of new and old C changes in the
446 system. Jointly considering C origins and pools enabled us to better depict the pathways of C flux from

447 plant roots to soil and among soil C pools. We found that, once soil was vegetated (either topsoil or
448 subsoil), the increase in total soil new C was mainly due to the increase in the least stable POM pool. This
449 result confirms the key role of plant roots in supply of C to the POM pool via root turnover, and is in line
450 with results by e.g. De Deyn et al. (2008), Cotrufo et al. (2013) and Rees et al. (2005). Surprisingly, we found
451 a high $\delta^{13}\text{C}$ signal in the SILT+CLAY pool for both soil types, which corresponds to a minor, but non-
452 negligible amount of new C supply into the most stable pool. Given that this phenomenon is more
453 pronounced for *M. sativa* (N_2 -fixing species, lower tissue recalcitrance due to lower C:N ratio) than for *L.*
454 *perenne* (non N_2 -fixing species, higher tissue recalcitrance due to higher C:N ratio), we may partially
455 attribute this phenomenon to the higher mineralization rate of the POM pool, that supplies the SILT+CLAY
456 pool. However, in this case, we argue that POM is not the only cause of the new C increase in the SILT+CLAY
457 pool, as POM, consisting of plant residues rich in cellulous and lignin, has a mean residence time much
458 higher than six months, i.e., the experiment duration (Cotrufo et al., 2015). Instead, it would be more likely
459 that such new C increase in the SILT+CLAY pool in the short term be a consequence of the higher microbial
460 proliferation and activity induced by a higher root exudation / microbial symbiosis with *Rhizobium* in *M.*
461 *sativa* (Cotrufo et al., 2015). Such a mechanism is incorporated as a part of the entombing effect in the
462 recent “Soil Microbial Pump” hypothesis (Liang et al. 2017). As an alternative pathway to the routinely
463 characterized ex vivo C flux from plant tissue to soil C pools via decomposition, the entombing effect refers
464 to the in vivo C flux from triggered microbial necromass and metabolites to the very stable soil C pools
465 (Liang et al. 2017). Although the estimation of microbial necromass was not available in this study and still
466 remains a technical bottleneck (Liang et al. 2019), we may expect a higher level of microbial necromass
467 due to the observed high GAM, H, DNA indicators in *M. sativa*, compared to *L. perenne* and bare soil.
468 Accordingly, our observed new C enhancement in both POM and the very stable SILT+CLAY C pools in our
469 experiment could be considered as novel data supporting the importance of the entombing effect.

3.4.2. Generally a strong synergy exists between new and old carbon (Hypothesis 2)

470 We found that soil that gained new C usually had a significantly smaller loss in old C. In topsoil sowed
471 with *M. sativa*, that had more new C input, , more labile tissues (low C:N content) and higher microbial
472 activity, old C loss in soil was much less than that in the vegetated subsoil treatment. This result is in
473 accordance with substrate utilization hypothesis developed by Cheng and Kuzyakov (2005) and observed
474 in an incubation experiment by De Graaf et al. (2010). However, to our knowledge, this is the first time we
475 observed this mechanism in an *in vivo* experiment. According to this hypothesis, microorganisms prefer
476 labile C to stable C, thus resulting in a limited consumption of old C, especially that protected by fine soil
477 particles. This mechanism is observed in soils with high fertility and mineral nutrients, and when the input
478 of fresh new C is adequate, which is our case in topsoil. However, when mineral nutrients are low and
479 fresh C input is low and insufficient to switch substrate utilization preference, the low input of C increases
480 the activity of microbes, that augment the consumption of old C (Cheng and Kuzyakov 2005; De Graaf et
481 al. 2010), as in our subsoil treatment. In addition to the preexisting hypothesis, due to the use of
482 fractionation, we can argue that the entombing effect in the soil microbial pump hypothesis (Liang et al.,
483 2017) can expand the comprehension of the synergetic pattern. We observed that the synergetic pattern
484 between new C and old C changes was largely due to the same pattern existing in the SILT+CLAY C pool
485 that received more than 50% of total soil C. Due to the entombing effect, the maintenance of old C content
486 against old C loss in the SILT+CLAY pool may be a consequence of increased microbial biomass relying on
487 the old C resource that consume the old C in unprotected coarser fractions and transfer it in SILT+CLAY via
488 entombing of microbial exudates, exopolysaccharides and necromass (Cotrufo et al. 2013; Liang et al.
489 2017; Vidal et al., 2018). Accordingly, soil with a greater microbial biomass (in our study, topsoil) may have
490 more advantages to maintain the size of the stable C pool via entombing effect (necromass, microbial
491 exudates and exopolysaccharides). Such a kind of increase in C due to microbial necromass based old C
492 should not be considered a part of old C. However, to what extent the increase in microbial necromass
493

494 relies on old C and new C is unknown, hindering the validity of the speculation. Overall, understanding the
495 role of microbial necromass and its underlying mechanism is an important scientific lacuna in soil ecology
496 to explore in the future.

497

498 *3.4.3. Root traits influence new carbon gain and old carbon changes, and are strongly*
499 *mediated by soil variables (Hypothesis 3)*

500 The two plant species that we examined had contrasting functional root traits, i.e., SRL, diameter of
501 absorptive roots and C:N ratio that were negatively correlated with the gain of new C in the POM pool,
502 and with new C in the SILT+CLAY C pool, however not with high significance. While this finding may be
503 possibly due to the short-term experiment in which species impact is not yet fully exerted, it could also be
504 attributed to the nature of these traits. Functional traits such as C:N ratio, diameter of absorptive roots,
505 and SRL are classified as morpho-physio-phenological (MPP) traits according to Violle et al., (2007), and
506 the impact of these functional traits can be compensated by the effect of biomass, i.e., a performance trait
507 (Violle et al., 2007). In agreement with this hypothesis, we found root biomass a much better predictor of
508 new C gain in every soil C pool compared to the C:N ratio and SRL.

509 We found that the effect of species on new C gain is much less pronounced in subsoil than in topsoil,
510 although the disparity of trait values between the two species in subsoil was still very clear. This result
511 suggests that the effect of root traits on C sequestration is strongly mediated by soil characteristics. In the
512 previous, similar studies working, soil treatment was usually excluded (Roumet et al., 2016; Henneron et
513 al., 2019; Rossi et al. submitted, Chapter II, this thesis). In this study, we used two soil types that were
514 similar in granulometric texture, but greatly differed in physical, chemical and biological qualities. Topsoil
515 had greater initial C and N contents, aggregate stability and soil biodiversity than subsoil, suggesting that
516 better soil quality is a primordial factor in influencing plant performance in C sequestration.

517 Compared to new C, Δ old C were generally much less sensitive to plant traits, including MPP traits e.g.,
518 C:N ratio, SRL and mean diameter and biomass. This result suggests that Δ old C does not share the same
519 mechanism with Δ new C and was less dependent on ex vivo C flux from plants. Compared to the MPP
520 traits, root biomass was a slightly better predictor of Δ old C. This result can also be explained by the
521 preferential substrate utilization hypothesis (Cheng and Kuzyakov, 2005) and the boosting effect of root
522 biomass on microbial proliferation and activity (Fontaine and Barot 2005, De Deyn et al. 2008). Microbial
523 communities may prefer consuming new C to old C, resulting in better maintenance of old C.

524 *M. sativa*, as a N₂ fixing species increase microbial activity via symbiosis with *Rhizobium* bacteria (Poirer et
525 al. 2018), augmenting microbial exudation and input of exopolysaccharides in the SILT+CLAY protected
526 pool (Fehrmann and Weaver, 1978; Downie, 2010; Cotrufo et al., 2013). The increased biomass of *M. sativa*
527 and its lower C:N ratio (due to its N fixing ability) increase the labile C input in soil (Warembourg et al.
528 2003; Roumet et al. 2005; Hernández et al. 2017) again increasing mineralization and deposition in the
529 SILT+CLAY protected pool. Being able to differentiate fluxes of old C and new C in soil allowed to observe
530 the increased input of new C from N₂-fixing *M. sativa*. This higher input, when analyzing the total Δ C, was
531 hidden by the changes in old C that were soil dependent and not species dependent. This result helps to
532 explain why different studies have discrepant results regarding the C storage from N₂ fixing and non N₂
533 fixing species, where not always N₂ fixing species significantly increased Δ C compared to non N₂-fixing
534 species (e.g. Binkley, 2005; Fornara and Tilman, 2008; Chapter II, this thesis). The higher input of new C
535 that N₂ fixing specie provide thanks to the higher root biomass, lower C:N ratio and fastest growth, might
536 have been hidden by soil dependent old C changes.

537 Overall, our finding highlights the necessity of studying the effect of functional traits on C sequestration in
538 a more refined manner (i) differentiating soil C origins and pools for a given soil enables us to better
539 identify soil C flux pathways that are more susceptible to vegetation; (ii) including the effect of soil type

540 can allow us to determine the magnitude of influence of plant trait disparities so as to take into account
541 more complex effects of interaction between soil features and root traits in future experimental design.

542

543 *3.4.4. Microbiological activity can explain the disparity in new C and old C changes*
544 *between topsoil and subsoil*

545 The soil C saturation theory states that a soil with lower amount of C in the fine SILT+CLAY particle fraction
546 has a higher potential for organomineral interactions and the derived C storage in the fine SILT+CLAY C
547 pool (Six et al 2002). Given the lower initial C content in the SILT+CLAY pool at t0 and the significant slightly
548 higher fine fraction ratio in subsoil, we expected a faster increase in new C in the SILT+CLAY pool given
549 the same amount of C input from biomass. Our results support the C saturation theory to a certain extent,
550 as the increase in new C in the SILT+CLAY per unit root biomass or length in subsoil was slightly higher than
551 that in topsoil but not significant (Fig. S5). This difference was disproportionally less than the difference in
552 initial C content between two soil types. We argue that the fine fraction abundance and soil C saturation
553 can have a positive influence on C stored via organomineral interactions if other conditions, especially soil
554 microbiological conditions are previously met. Recent studies have highlighted the importance of
555 considering the robustness of soil microbial diversity as a soil quality indicator (Bouchez et al., 2016; Karimi
556 et al., 2017), thus challenging the conventional use of only physical and chemical soil quality indicators. In
557 this study, we have shown that microorganisms play a central role in the gain of new C and loss of old C in
558 the SILT+CLAY pool. The higher microbial activity and diversity in topsoil from t0 to t6 may compensate
559 the less favorable physical and chemical quality (lower fine fraction and higher initial C content) for C
560 sequestration. Therefore, if microbial communities are not considered in the prediction of soil C
561 sequestration, results will be flawed.

562 Along with greater microbial diversity, an increase in aggregate stability (MWD) and N content in topsoil
563 could also promote a synergetic effect that augments C sequestration. The physical protective role of
564 aggregates for C stock is widely documented (Hassink et al., 1992; Six et al., 2002; Chevallier et al. 2004;
565 Rasse et al. 2005; O'Brien et al., 2013; King et al., 2019). A high soil N content in topsoil will also improve
566 plant development and subsequent biomass, thus affecting C input and microbial diversity. We suggest
567 therefore, that a comprehensive indicator of soil health for plant performance and C sequestration should
568 incorporate physical, chemical and microbiological characteristics.

569

570 *3.4.5. Practical applications*

571 This study provides useful implications for future engineers to choose appropriate soil and species in road
572 embankment revegetation to favor C sequestration. First, with the dominant effect of soil over species
573 found in this study, choosing healthy and functional soil is of primary importance for C sequestration.
574 Topsoil has shown a clearly better performance in C sequestration than subsoil. However, implementing
575 topsoil over large scales is unrealistic when revegetating a site, because the amount of topsoil is relatively
576 limited and over-exploitation of topsoil may further provoke environmental issues for the location where
577 the topsoil is removed. Although subsoil has higher C sequestration potential due to its lower initial C
578 content, attention should be paid to the microbial diversity and functioning in subsoil. Inoculation of soil
579 with suitable microbial communities and fertilizer would therefore be necessary to favor both
580 revegetation and soil C sequestration (Dou et al. 2016, Guo et al., 2019).

581 Once soil quality is ensured, choosing appropriate species will be a bonus for boosting new C input and
582 protecting old C against priming. In our experiment that lasted 6 months, and so corresponds to the initial
583 planting stage in the field, *M. sativa* had a better performance than *L. perenne*, and also enhanced soil

584 aggregate stability, thus decreasing soil erodibility. However, the long-term effects of revegetation on
585 long-term soil C fates should also be investigated.

586

587 3.5. CONCLUSIONS

588 We designed an experiment with fully crossed treatments between vegetation and soil in microcosms and
589 used stable isotopic (^{13}C) labelling to assess new C input and old C changes in the soil system. We revealed
590 the distinct fates of new C and old C in soil, in both absolute values and relative values, among different
591 soil C pools related to soil fractions, highlighting:

- 592 • The major influence of soil, with topsoil having a higher C storage capacity compared to subsoil
593 due to higher soil quality that increase biomass development and C input, and higher microbial
594 biomass and activity that favors entombing of C in the stable SILT+CLAY pool.
- 595 • We evidence the necessity of considering both C fluxes in pools associated to soil fractions and
596 origin of C (new and old C) when studying C dynamics in soil. An example being old C decreasing
597 in the POM C pool and increasing in the SILT+CLAY C pool in topsoil. If only ΔC or $\Delta\text{old C}$ in bulk
598 soil was considered no changes would have been observed, and the old C would have been
599 considered inactive, masking the real mechanisms behind soil C sequestration in topsoil.
- 600 • New C increased not only in the POM C pool, but also in the more stable SILT+CLAY pool. Given
601 the short duration of the experiment, this flux is probably due to entombing of microbial
602 necromass and microbial exudates and exopolysaccharides more than degradation of POM.
- 603 • New C and old C covaried similarly in the SILT+CLAY C pool. A higher increase of new C resulted
604 in a lower decrease of old C due to microbiological switch of substrate preference.

- 605 • Changes in new and old C differed depending on plant and soil characteristics. N₂ fixing *M.*
606 *sativa* higher root biomass labile input in soil increased the amount of new C in soil. N₂ fixing
607 *M. sativa* also increased microbial biomass and activity that favor the mineralization of C and
608 transport into the SILT+CLAY protected fraction. Root biomass was the trait better correlated
609 with new C input in soil C pools.
- 610 • The lack of microbiological activity and the lower root biomass decreased the transfer of new
611 C in the SILT+CLAY pool in subsoil. For this reason, the lower C saturation did not increase the
612 total new C content in SILT+CLAY pool in subsoil as expected. When normalized for the root
613 biomass, however, the system showed the opposite behavior, and subsoil had a higher amount
614 of new C stored in SILT+CLAY for g of root. We argue that C saturation effect might be present
615 but is subdued to soil fertility and microbiological activity.

616 Such a fundamental understanding of plant-soil interactions may help us to better optimize soil and
617 vegetation management for road embankment revegetation. Long-term observations are now needed for
618 a better assessment of the roles of plant and soil characteristics in soil C cycling and long-term
619 sequestration.

620

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- 783

784 FIGURES AND TABLES

785 **Table 1:** Root, microbial and soil characteristics analyzed at time 0, after 183 days of experiment (time 6 months), and
 786 the difference between time 0 and time 6, for *M. sativa*, *L. perenne* sowed on topsoil and subsoil, plus bare soil controls.
 787 Root traits: Root biomass (g), specific root length (SRL, m g⁻¹ soil), diameter of absorptive roots (mm), adsorptive roots
 788 C:N ratio; Microbial characteristics: global metabolic activity (GMA, µgC-CO₂ g⁻¹ soil h⁻¹), Shannon metabolic diversity
 789 (H), concentration of DNA in solution as proxy for microbial biomass (DNA, ng µL⁻¹); Soil characteristics: mean weight
 790 diameter of aggregates (MWD, mm), soil nitrogen content (Soil N, mg N g⁻¹ soil), percentage of fine fraction <20µm
 791 in soil (FF, %).

		SUBSOIL			TOPSOIL			
		Bare soil	<i>L. perenne</i>	<i>M. sativa</i>	Bare soil	<i>L. perenne</i>	<i>M. sativa</i>	
Time 0	<i>Micorbial characteristics</i>	GMA (µgC-CO ₂ g ⁻¹ soil h ⁻¹)	7.56±1.1	-	-	8.68±1.94	-	-
		H (-)	0.85±0.17	-	-	1.14±0.05	-	-
		DNA concentration (ng µL ⁻¹)	1.00±0.00	-	-	13.67±3.06	-	-
	<i>Soil characteristics</i>	MWD (mm)	0.82±0.03	-	-	1.55±0.02	-	-
		Soil N (mg N g ⁻¹ soil)	0.75±0.04	-	-	1.25±0.06	-	-
		FF (%)	51.00±1.00	-	-	43.00±1.00	-	-
<i>Root traits</i>	Root biomass (g)	-	1.05±0.34	5.13±1.36	-	4.09±1.43	17.53±2.03	
	SRL absorptive roots (m g ⁻¹ soil)	-	34.92±4.72	14.77±7.87	-	23.25±2.98	10.25±0.57	
	Diameter absorptive roots (mm)	-	0.09±0.02	0.27±0.04	-	0.10±0.02	0.26±0.01	
	Absorptive roots C:N	-	90.78±15.53	20.12±2.42	-	51.48±10.73	20.23±1.31	
Time 6 months	<i>Micorbial characteristics</i>	GMA (µgC-CO ₂ g ⁻¹ soil h ⁻¹)	6.77±0.72	7.41±2.50	5.98±1.06	7.41±1.72	11.69±1.02	19.06±4.21
		H (-+)	0.78±0.07	0.93±0.08	0.90±0.04	1.16±0.01	1.17±0.00	1.17±0.00
		DNA concentration (ng µL ⁻¹)	4.00±2.22	3.00±1.41	5.20±1.64	18.40±5.90	18.40±5.59	19.00±4.32
	<i>Soil characteristics</i>	MWD (mm)	0.84±0.10	0.78±0.07	0.89±0.08	1.64±0.08	2.17±0.20	2.07±0.29
		Soil N (mg N g ⁻¹ soil)	0.63±0.05	0.65±0.08	0.65±0.05	1.19±0.14	1.12±0.10	1.23±0.10
		FF (%)	49.84±1.17	50.62±1.16	50.77±0.58	42.24±1.26	42.01±1.31	41.87±1.42
Δ t0-t6	<i>Micorbial characteristics</i>	GMA (µgC-CO ₂ g ⁻¹ soil h ⁻¹)	3.00±2.22	-0.16±2.73	-1.59±1.53	-1.27±2.59	3.01±2.19	10.39±4.63
		H (-)	-0.07±0.18	0.08±0.18	0.06±0.17	0.02±0.05	0.03±0.05	0.03±0.05
		DNA concentration (ng µL ⁻¹)	3.00±2.22	2.00±1.41	4.20±1.64	4.73±6.64	4.73±6.37	5.33±5.29
	<i>Soil characteristics</i>	MWD (mm)	0.02±0.1	-0.03±0.08	0.08±0.08	0.1±0.08	0.62±0.20	0.52±0.29
		Soil N (mg N g ⁻¹ soil)	-0.11±0.07	-0.1±0.09	-0.1±0.07	-0.06±0.15	-0.13±0.12	-0.02±0.12
		FF (%)	-0.01±0.02	-0.01±0.01	0.00±0.01	0.00±0.02	-0.01±0.02	-0.01±0.02

792 For Δt0-t6 the red values indicate a loss in 6 months, black value a gain

793

794 **Table 2:** Statistical analysis of the effect of treatments on on root, microbial and soil characteristics analyzed after 183
795 days of experiment for *M. sativa*, *L. perenne* sowed on topsoil and subsoil, plus bare soil controls. Root traits: Root
796 traits: Root biomass (g), specific root length (SRL, m g⁻¹ soil), diameter of absorptive roots (mm), adsorptive roots C:N
797 ratio; Microbial characteristics: global metabolic activity (GMA, µgC-CO₂ g⁻¹ soil h⁻¹), Shannon metabolic diversity (H),
798 concentration of DNA in solution as proxy for microbial biomass (DNA, ng µL⁻¹); Soil characteristics: mean weight
799 diameter of aggregates (MWD, mm), soil nitrogen content (Soil N, mg N g⁻¹ soil), percentage of fine fraction <20µm in
800 soil (FF, %).. Data where normal according to the Shapiro-Wilk test and the ANOVA test was utilized to asses statistical
801 differences.

	Soil	Variable	df	F	p
Effect of treatment	Topsoil	Root biomass	1,7	136.8	<0.001***
		SRL absorptive roots	1,6	73.33	<0.001***
		Diameter absorptive roots	1,6	224.6	<0.001***
		Absorptive roots C:N	1,7	32.61	<0.001***
		GMA	2,11	24.19	<0.001***
		H	2,11	5.925	0.0179 *
		DNA concentration	2,11	0.018	0.983
		MWD	2,11	9.953	0.00341 **
		Soil N	2,11	1.168	0.347
	FF	2,11	0.034	0.967	
	Subsoil	Root biomass	1,8	42.13	<0.001***
		SRL absorptive roots	1,7	22.98	0.00198 **
		Diameter absorptive roots	1,7	64.44	<0.001***
		Absorptive roots C:N	1,7	104	<0.001***
		GMA	2,11	0.914	0.429
		H	2,11	6.827	0.0118 *
		DNA concentration	2,11	2.181	0.159
		MWD	2,11	2.263	0.15
Soil N		2,11	0.079	0.925	
FF	2,11	0.662	0.535		
Effect of Soil	<i>M.sativa</i>	Root biomass	1,7	120.9	<0.001***
		SRL absorptive roots	1,6	1.314	0.295
		Diameter absorptive roots	1,6	0.098	0.765
		Absorptive roots C:N	1,7	0.006	0.939
		GMA	1,7	46.22	<0.001***
		H	1,7	150.6	<0.001***
		DNA concentration	1,7	44.35	<0.001***
		MWD	1,7	80.22	<0.001***
		Soil N	1,7	129.6	<0.001***
	FF	1,7	159.8	<0.001***	
	<i>L.perenne</i>	Root biomass	1,8	21.33	<0.001***
		SRL absorptive roots	1,7	18.3	0.00366 **
		Diameter absorptive roots	1,7	0.738	<0.001***
		Absorptive roots C:N	1,7	20.3	0.00278 **
		GMA	1,8	12.57	0.00757 **
		H	1,8	40.75	<0.001***
		DNA concentration	1,8	35.61	<0.001***
		MWD	1,8	220.4	<0.001***
		Soil N	1,8	72.42	<0.001***
	FF	1,8	132.1	<0.001***	
	Bare soil	GMA	1,7	0.398	0.548
		H	1,7	149	<0.001***
		DNA concentration	1,7	18.83	0.0034 **
		MWD	1,7	178.9	<0.001***
		Soil N	1,7	56.59	<0.001***
		FF	1,7	45.36	<0.001***

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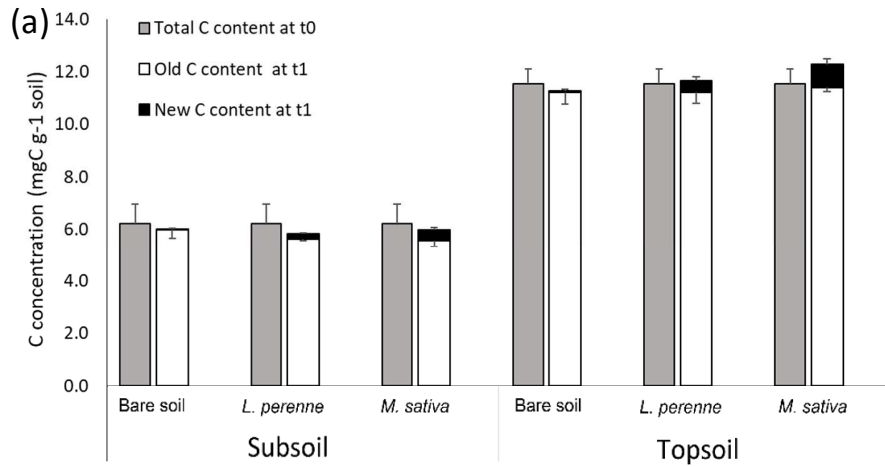
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804 **Table 3:** Pearson's correlation coefficients (r) showing relationships between microbial characteristics, root variables,
 805 soil structural characteristics and New, Old C, and ΔC in different soil C pools. Root traits: Root biomass (g), specific
 806 root length (SRL, m g⁻¹ soil), diameter of absorptive roots (mm), adsorptive roots C:N ratio; Microbial characteristics:
 807 global metabolic activity (GMA, $\mu\text{gC-CO}_2 \text{ g}^{-1} \text{ soil h}^{-1}$), Shannon metabolic diversity (H), concentration of DNA in solution
 808 as proxy for microbial biomass (DNA, ng μL^{-1}); Soil characteristics: mean weight diameter of aggregates (MWD, mm),
 809 soil nitrogen content (Soil N, mg N g⁻¹ soil), percentage of fine fraction <20 μm in soil (FF, %). Data where normal
 810 according to the Shapiro-Wilk test and the ANOVA test was utilized to asses statistical differences.

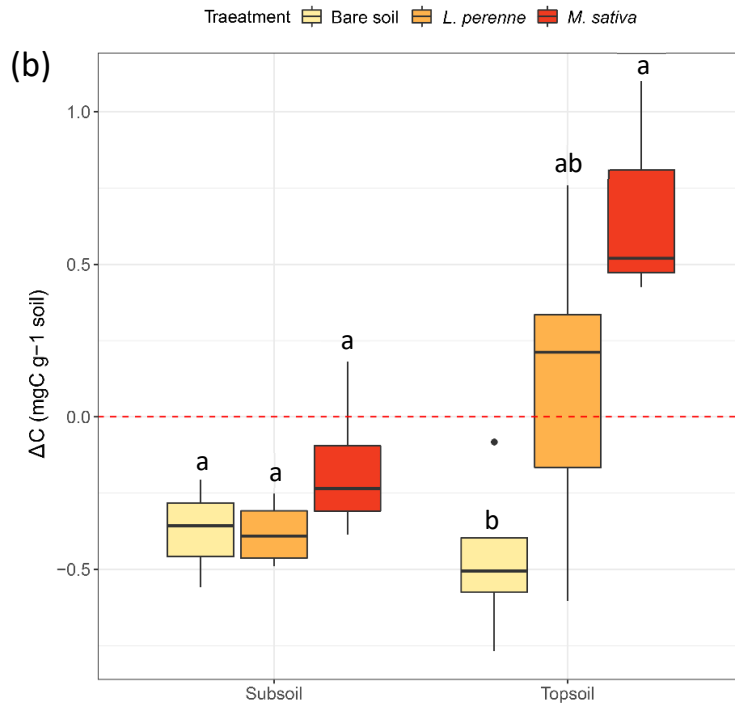
	Root biomass (R_bio)	Specific root length of absorptive roots (SRL_AD)	Diameter of absorptive roots (Diam_AD)	C:N ratio of absorptive roots (C:N_A D)	Global metabolic activity (GMA)	Shannon metabolic diversity index (H)	Microbial DNA concentration (DNA)	Aggregates mean weight diameter (MWD)	Soil nitrogen (Soil_N)	Fine fraction percentage (FF)
<i>NewC_{POM}</i>	0.77***	-0.69**	0.59**	-0.66**	0.57*	0.34	0.53*	0.36	0.43	-0.36
<i>NewC_{finePOM}</i>	0.57*	-0.4	0.06	-0.39	0.71***	0.90***	0.87***	0.97***	0.94***	-0.94***
<i>NewC_{silt}</i>	0.60**	-0.37	0.06	-0.35	0.73***	0.86***	0.85***	0.94***	0.94***	-0.90***
<i>NewC_{silt+clay}</i>	0.83***	-0.61**	0.45	-0.57*	0.84***	0.69**	0.71***	0.74***	0.82***	-0.76***
<i>NewC_{SUM}</i>	0.88***	-0.73***	0.56*	-0.69**	0.76***	0.57*	0.71***	0.61**	0.69**	-0.62**
<i>OldC_{POM}</i>	0.25	-0.08	-0.08	-0.04	0.38	0.53*	0.38	0.57*	0.59**	-0.57*
<i>OldC_{finePOM}</i>	0.58**	-0.3	0.14	-0.17	0.51*	0.36	0.33	0.38	0.43	-0.38
<i>OldC_{silt}</i>	0.06	-0.2	0.05	-0.17	-0.09	0.05	0.05	-0.02	-0.11	0.02
<i>OldC_{silt,clay}</i>	-0.55*	0.28	0	0.21	-0.71***	-0.83***	-0.70***	-0.87***	-0.87***	0.86***
<i>OldC_{SUM}</i>	-0.44	0.24	-0.07	0.23	-0.59**	-0.57*	-0.60**	-0.58**	-0.59**	0.56*
ΔC_{POM}	0.02	-0.13	0.24	-0.17	-0.16	-0.36	-0.16	-0.39	-0.38	0.39
$\Delta C_{finePOM}$	-0.47*	0.22	-0.13	0.09	-0.37	-0.17	-0.37	-0.17	-0.24	0.18
ΔC_{silt}	0.01	0.16	-0.04	0.13	0.17	0.05	0.17	0.12	0.21	-0.12
$\Delta C_{silt+clay}$	0.60**	-0.33	0.05	-0.26	0.74***	0.83***	0.74***	0.87***	0.89***	-0.87***
ΔC_{SUM}	0.72***	-0.52*	0.33	-0.50*	0.74***	0.63**	0.74***	0.66**	0.70***	-0.65**

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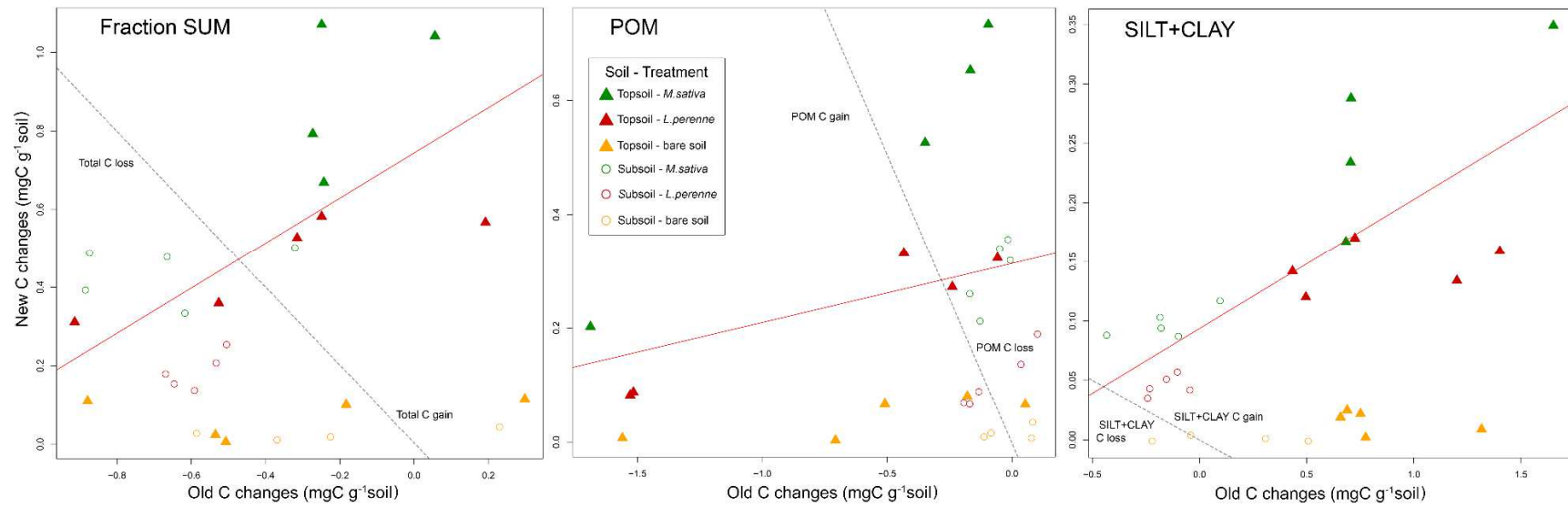


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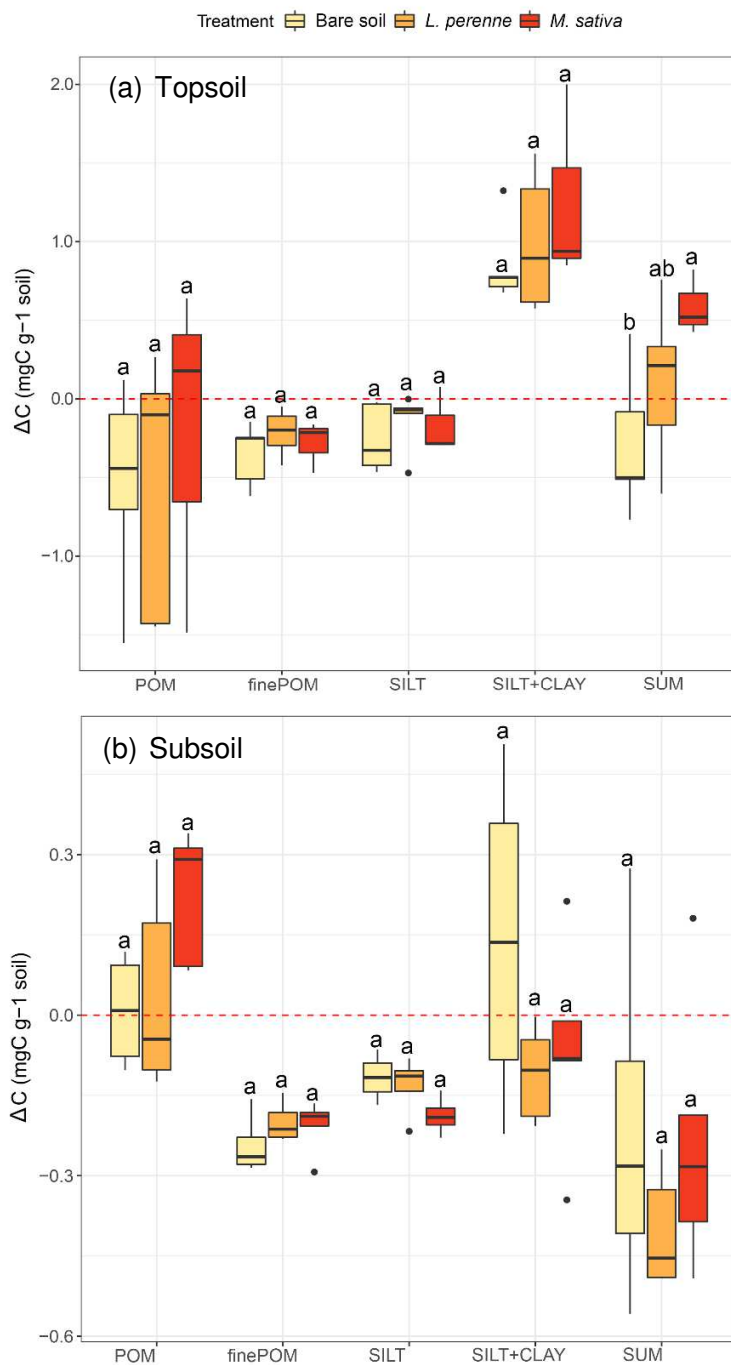
814

815 **Figure 1.** Total soil carbon (C) concentration (a) and concentration changes (b) among different soil types and
 816 vegetation treatments from *t*₀ (experiment set-up) to *t*₆ (harvest, i.e. 183 days after). In (a): total C concentration at *t*₀,
 817 corresponding to old C concentration at *t*₀, are all identical within each soil type. In (b), for each boxplot, the lower
 818 edge of the box corresponds to the 25th percentile data point, while the top edge of the box corresponds to the 75th
 819 percentile data point; the upper and lower vertical lines corresponds to the 90th and 10th percentile data points,
 820 respectively; the horizontal line within the box represents the median and black dots indicate outliers. Letters above the
 821 boxplots indicate statistically significant differences (p < 0.05*) between species and controls according to Tukey HSD
 822 test.



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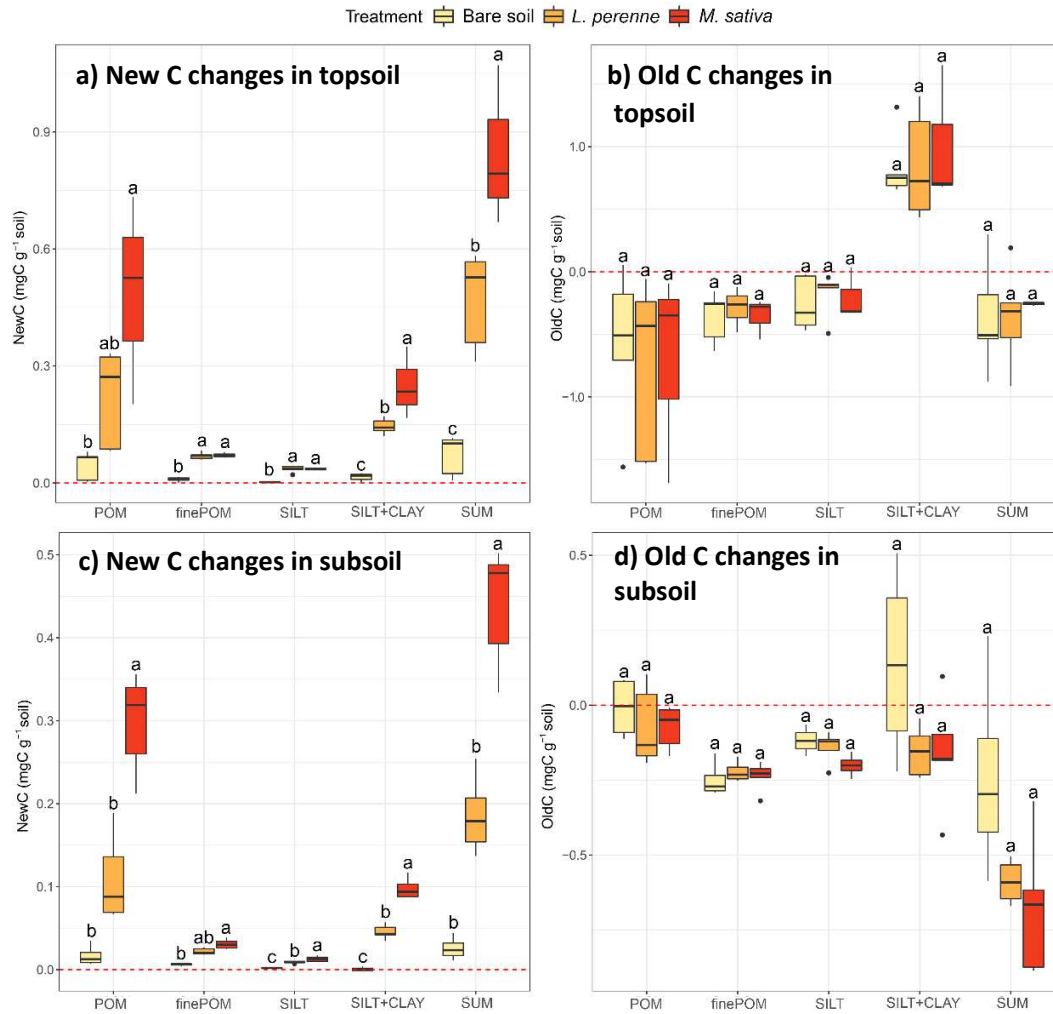
825 **Figure 2.** Relationships between the loss of OldC and gain in NewC in soil for (a) the sum of old and new C in every pool, (b) the POM pool and (c) the SILT+CLAY
 826 pool. The red solid line shows correlation between old c decrease and new c input for the vegetated treatment (without considering bare soil control). The grey dashed
 827 line shows the divide of the data between C gain (on the right) and loss (on the left) oin sum of fraction, POM and SILT+CLAY pools.



828

829 **Figure 3:** Comparison of the difference in carbon (ΔC) after 6 months in different soil C pools and for each treatment in
 830 a) subsoil and b) topsoil. In each boxplot, the lower edge of the box corresponds to the 25th percentile data point, while
 831 the top edge of the box corresponds to the 75th percentile data point. The line within the box represents the median
 832 and black dots indicate outliers. Different letters above the boxplots indicate statistically significant differences ($p < 0.05$)
 833 between families and controls according to Tukey HSD test.

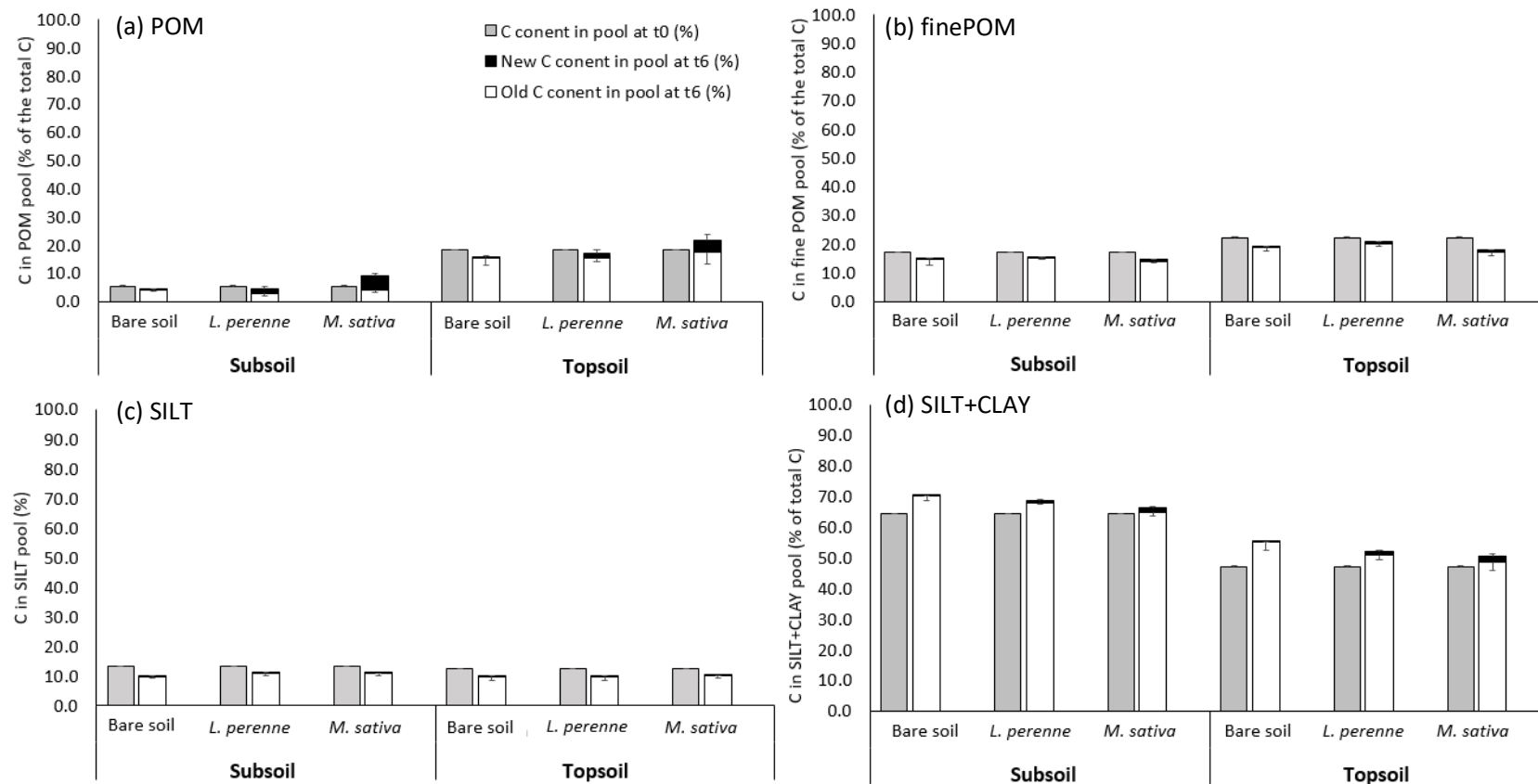
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 836 **Figure 4:** Gain of new C and changes in old C in 4 different C pools (POM, finePOM, SILT, SILT+CLAY) and in the
 837 total bulk soil (as sum of the different pools, Sum) for bare soil control (light yellow), *L. perenne* (orange), and *M. sativa*
 838 (red). a) shows the fluxes of new C in topsoil, b) the fluxes of old C in topsoil, c) the fluxes of new C in subsoil, and d)
 839 the fluxes of old C in subsoil. In each boxplot, the lower edge of the box corresponds to the 25th percentile data point,
 840 while the top edge of the box corresponds to the 75th percentile data point. The line within the box represents the
 841 median and black dots indicate outliers. Different letters above the boxplots indicate statistically significant differences
 842 ($p < 0.05$) between species treatments according to Tukey HSD test.

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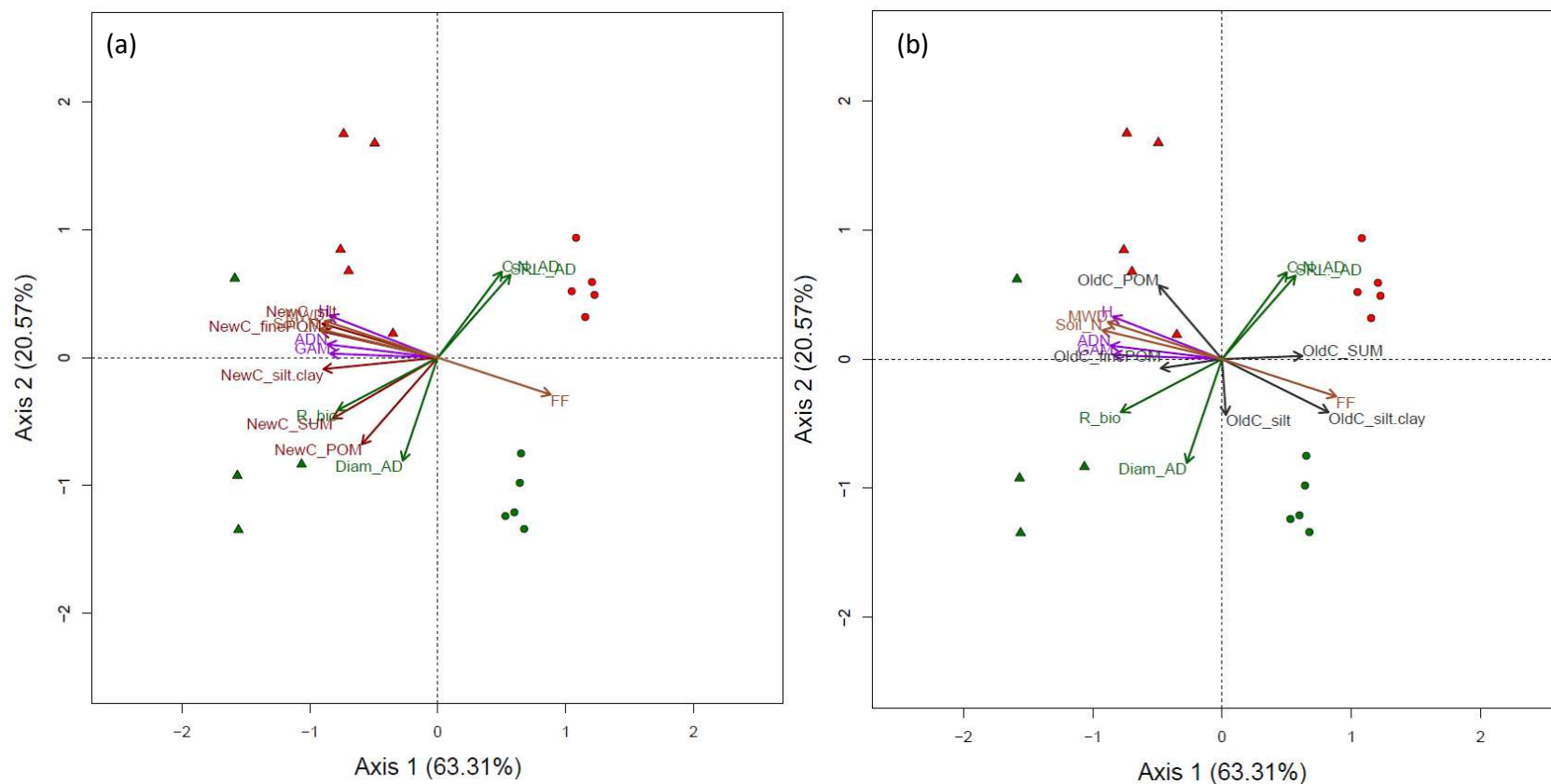
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846 **Figure 5:** C concentration in % of total C in the different soil C pools at t0 and t6 (after 183 days of experiment). At t6 the C is divided in % of new C in soil (black)
 847 and old C in soil (white). (a) shows the C concentration in % in POM pool, (b) in fine POM pool, (c) in SILT pool, and (d) in SILT+CLAY pool.

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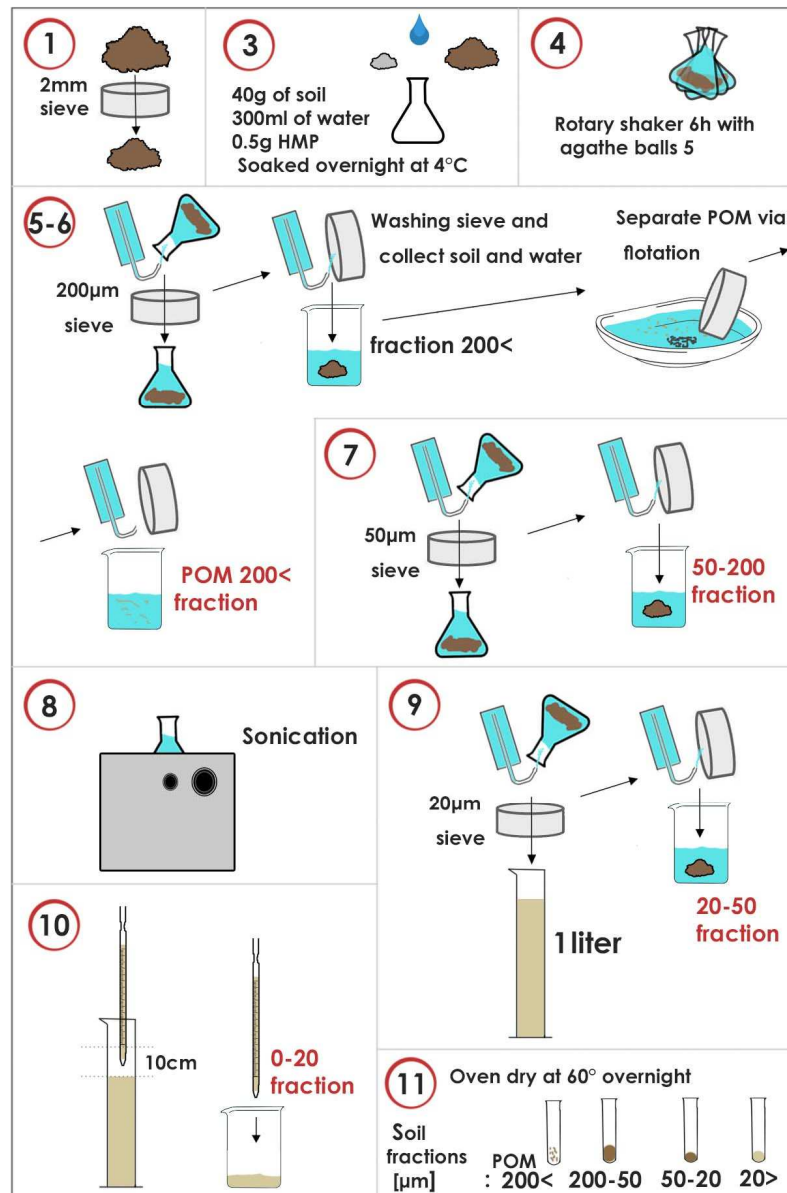
851 **Figure 6:** Principal Component Analysis of (a) new C input and (b) old C fluxes in different soil C pools and root traits (SRL_AD: specific root length of absorptive
 852 roots, C:N_AD: C:N ratio of absorptive roots, R_bio: root biomass and Diam_AD: diameter of absorptive roots), microbial processes indicators (ADN: concentration
 853 of DNA as proxy for microbial biomass, GMA: global metabolic activity, H: Shannon metabolic diversity), and soil structure indicators (MWD: mean weight diameters
 854 of aggregates, Soil_N: nitrogen content in soil, and FF: percentage of fine fraction <20 μm) in soil. Triangles are topsoil and dots subsoil. Green is *M. sativa* and red
 855 is *L. perenne*. (a) and (b) shows the same PCA analysis, but in (a) only the arrows of new C are shown, while in (b) only the arrows for old C, for a better comprehension
 856 of the graph.

857 **SUPPLEMENTARY MATERIALS**

858 **Method S1:** Soil fractionation according to Gavinelli et al. (1995) methodology

859 Gavinelli et al. (1995) methodology:

- 860 1. Soil is sieved at 2mm and 40g are collected for fractionation
- 861 2. Soil presoaked overnight at 4 °C in 300 mL of deionized water with 0.5g HMP (sodium metaphosphate)
- 862 3. Shaken with 5 agate balls (d 10 mm) in a rotary shaker, maximum frequency for 2h in case of sandy soils,
863 6h for other soils.
- 864 4. Soil suspension wet sieved through a 200 µm
- 865 5. Fraction remaining on sieves (2000-200 µm) washed with water in a bowl for POM separation via flotation,
866 while the remaining >200 µm fraction is collected in a baker for further fractioning
- 867 6. The POM is separated from the sand fractions by submerging the 2000-200 µm fraction in deionized water.
868 The POM will float while the sand will drown. Carefully collect the POM using a sieve and separate it from
869 the sand (**coarse POM fraction**). The sand fraction is collected in a glass beaker after separation from POM
870 (sand fraction)
- 871 7. >200 µm suspension is sieved at 50 µm and the 200-50 µm is gently washed with deionized water from
872 the sieve and collected in a glass baker (**finePOM fraction**)
- 873 8. >50 µm suspension sonicated for 10 minutes
- 874 9. >50 µm suspension sieved with 20 µm screen and 50-20 µm is gently washed with deionized water from
875 the sieve and collected in a glass baker (**coarse SILT fraction**)
- 876 10. Transfer of >20 µm suspension in 1 L glass cylinder and add water to bring the volume to 1 L
- 877 11. >20 µm suspension shaken by hand (30 tumbling) and collection of 100 ml immediately after (aliquot for
878 the **fine SILT+CLAY fraction**)
- 879 12. The resulting beakers containing the soil suspension of the different fractions are collected and oven dried
880 at 60 °C until all the water evaporates

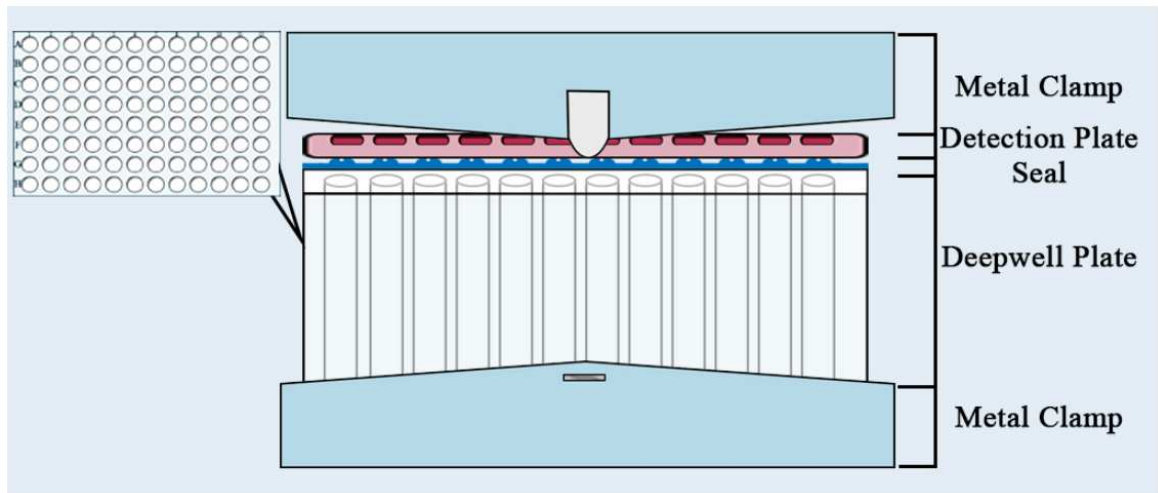


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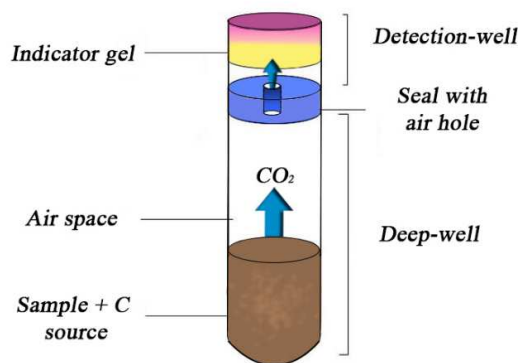
883

Figure S1: Soil fractionation according to the Gavinelli et al. (1995) methodology



884

885 **Figure S2** (ONICA et al.2018): scheme of a MicroResp system



886

887 **Fig. S3** (ONICA et al.2018): scheme of the functioning of a deep well detection system. Soil is placed in a deepwell and sealed,
 888 the CO₂ respired and accumulating in the detection well change the colour of the detection gel according to the equation [4.5]

889 **Method S2**

890 The principle of the MicroResp is to utilize a colored detection gel (Cresol red) that change color when changing
 891 pH or increasing the CO₂ concentration according to the equation:



893 When the pH diminishes, the Cresol red turns from pink to yellow. (Fig. S3)

894 The first step in the process is to prepare the detection gel in the MicroResp plaques. The preparation of detection
 895 gel is a fundamental step to assess the calibration curve of each detection plaque and calibrate the results based
 896 on the gel preparation, where $\% \text{CO}_2 = A + B / (1 + D * \text{DO}_{\text{norm}})$. First an Agar solution is prepared, mixing 3g of agar

897 with 100ml of deionized water, mixed in an autoclave for 20 minutes at 120°C. After the indicator solution is
898 prepared mixing 0.315 g NaHCO₃ (final: 2.5 mM), 16.77 g KCl (final: 150 mM) and 18.75 mg cresol red (final: 12.5
899 µg ml⁻¹ or 20 ppm), in 900 ml of deionized water and brought at 65°C to dissolve. The 1000ml solution is
900 transferred to an open bottle and stored at 4°C for 6 months maximum. To prepare the gel 100ml of the 3% agar
901 solution are melted and kept at 60°C. Separately, other 200ml of the indicator solution are melted at 60°C while
902 stirring, and after mixed with the agar solution. The mixture is then distributed in the detection plaques, 150µl of
903 solution for each one of the 96 detection well of each plaque. To avoid the formation of bubbles the gel is kept
904 throughout the process in a baker of boiling water and the tip of the pipette used to distribute the solution is
905 preheat in boiling water. When the gel is solidified (2-3h) the plaques are stored in a dessicator at room
906 temperature in a dark room with a baker of soda and a cup of water for 2-3 days to allow each microplate to reach
907 CO₂ equilibrium. After they are covered in parafilm.

908 After the detection plaque are prepared a calibration process is needed for every new detection plaque. 12 strips
909 (8 wells each) made for calibration containing the indicator gel are scanned used a Victor 1420 Multilabel Counter
910 (PerkinElmer, Massachusetts, USA) to assess the initial DO₅₉₀ (DO_{initial}). Twenty-four 150ml serum vials are
911 prepared, each containing ½ a strip of detection gel (4 wells), and injected with known CO₂ concentration with a
912 syringe in order to have a CO₂ concentration range (from 0% CO₂ concentration to 4% CO₂ concentration, 6040
913 vol CO₂ (µl), increasing the concentration in each vial of 0.1%). The strips are incubated for 2h at 25°C to achieve
914 balance with the CO₂ in each serum vial. After 2h the concentration of CO₂ is assessed in the vials using the GC-
915 microcatha measurement. The detection strips are then retrieve and immediately read at 590nm to assess the
916 DO_{final}. Finally the calibration is finalized as follow:

- 917
- Normalize DO data: $DO_{norm} = DO_{final} / DO_{initial} * average(DO_{initial})$
 - Draw the DO_{norm} vs [CO₂] calibration curve in %.
 - Fit the curve (rectangular hyperbola regression $\%CO_2 = A + B / (1 + D * DO_{norm})$)
- 918
- 919

920

921 The next step in the MicroResp protocol is to prepare the different substrate solutions. The idea is to give 1.5 mg
922 of C for each g of dry soil (substrate saturation) and reach a humidity level of 80% of field capacity. For each soil
923 type is therefore necessary to determine 1) the field capacity in g of water per g of soil and 2) the soil mass
924 distributed in each well by the MicroResp filling device. Field capacity for the soil was calculated at 28.3%. Three
925 plaques for each soil were filled with the MicroResp filling device and weighted to estimate the average soil
926 content in each MicroResp well, set at 0.5±0.04g well⁻¹. 1.2ml of solution have been added to each MicroResp well.

927 The solution have been prepared using milliQ water and sterilized using 0.45 µm paper filters, stored in sterilized
928 falcon tubes at 4°C.

929 The substrates used for the MicroResp analyses were chosen based on their biological properties and are reported
930 in Table S1. In every detection plate an extra substrate with pure MillQ water were added as control. In each
931 plaque (96 wells) we tested 2 samples, one in each half of the plaque, for a total of 48 wells per samples organized
932 as follow: 3 sub replicates per substrate (15 substrate) plus 3 sub replicate per the millQ water control (Fig.S4).

933 Each sample was prepared as follow:

- 934 • Identify the deepwell plate and the sample used
- 935 • Tare the deepwell plate
- 936 • Hide half of the filling device, place it above a deepwell plate, covered with plastic sheeting, fill half of
937 the filling device with the ground and remove the excess with a spatula.
- 938 • Pull the plastic sheet to drop the soil into the wells, then weigh and record the mass of soil used for the
939 half plate (48 wells). Tare again before filling the second half of the plate with the other soil sample.

940 The samples are analyzed as follow:

- 941 • At time 0 the substrate are added using a multichannel pipette to each wells of the deepwell plate, cover
942 with the parafilm and incubated at 25°C in the dark for 2h
- 943 • Before the end of the 2h incubation, the DO_{590t0} of each detection placed is taken with a Victor 1420
944 Multilabel Counter (PerkinElmer, Massachusetts, USA). After two hours the detection plate is placed
945 above the analysis plate with and sealed with a clamp. Resume incubation at 25°C for an additional 4
946 hours.
- 947 • At time 6h: the detection plate are detached from the deepweell and immediately read using a Victor
948 1420 Multilabel Counter (PerkinElmer, Massachusetts, USA) to determine the DO_{590t6}

949 For the data analysis the following steps were taken:

- 950 • Retrieve DO data at t0 and t6 for each plate. Check that the coefficient of variation of the DO_{590} of each
951 detection half plate at t0 does not exceed 5% (otherwise, remove the outliers DO values).
- 952 • Normalize the data: $DO_{norm} = DO_{t6}/DO_{t2} * \text{average}(DO_{t2})$
- 953 • Convert DO_{norm} to % CO2 from calibration data: $\%CO_2 = A + B / (1 + D * DO_{norm})$. The data for the
954 calibration curve were A = -0.29, B = -0.87, D = -7.72

955 • Convert these %CO₂ to SIR ($\mu\text{g C-substrate g}^{-1} \text{ soil h}^{-1}$) according to the incubation time and soil mass in
956 each well.

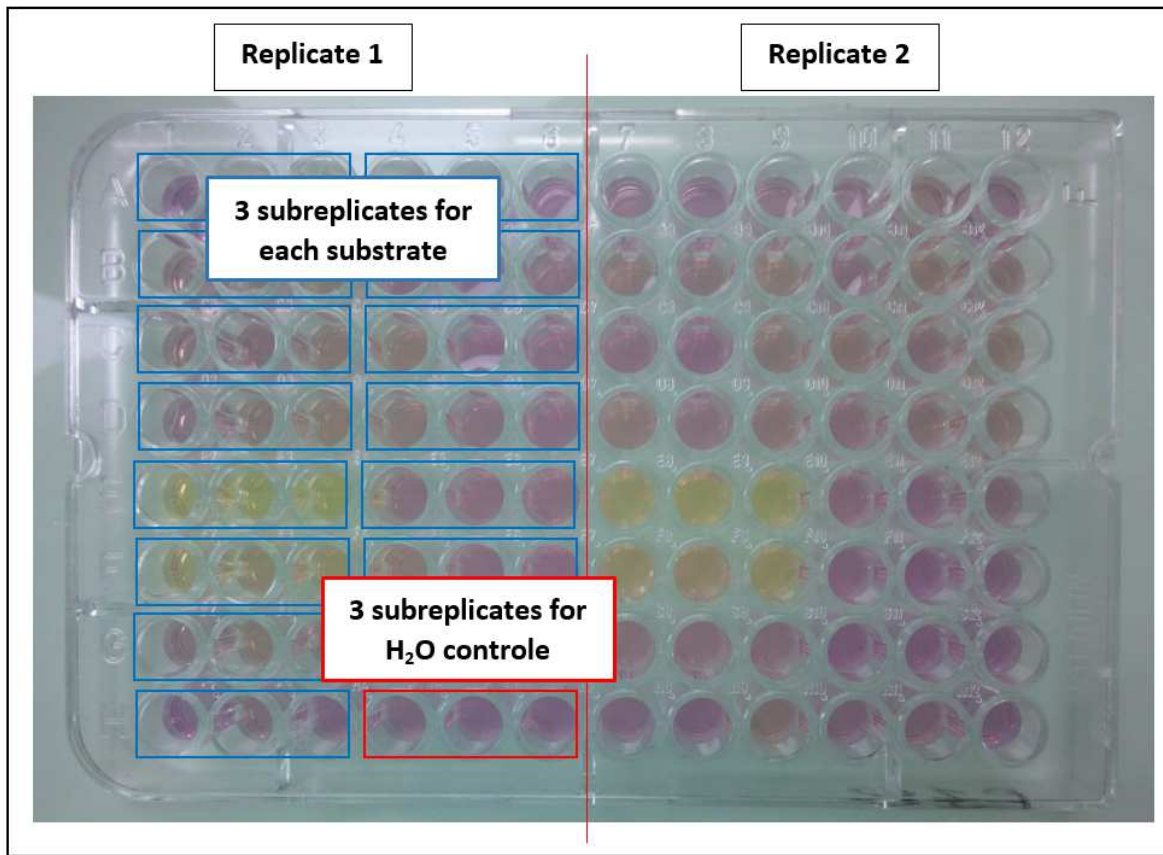
957 The aim of the MicroResp is to characterize the differences in functional activity of microbiological communities.

958 The activity was assessed on soil sampled and t₀ and t₆.

959

Table S1: Substrates used for the MicroResp

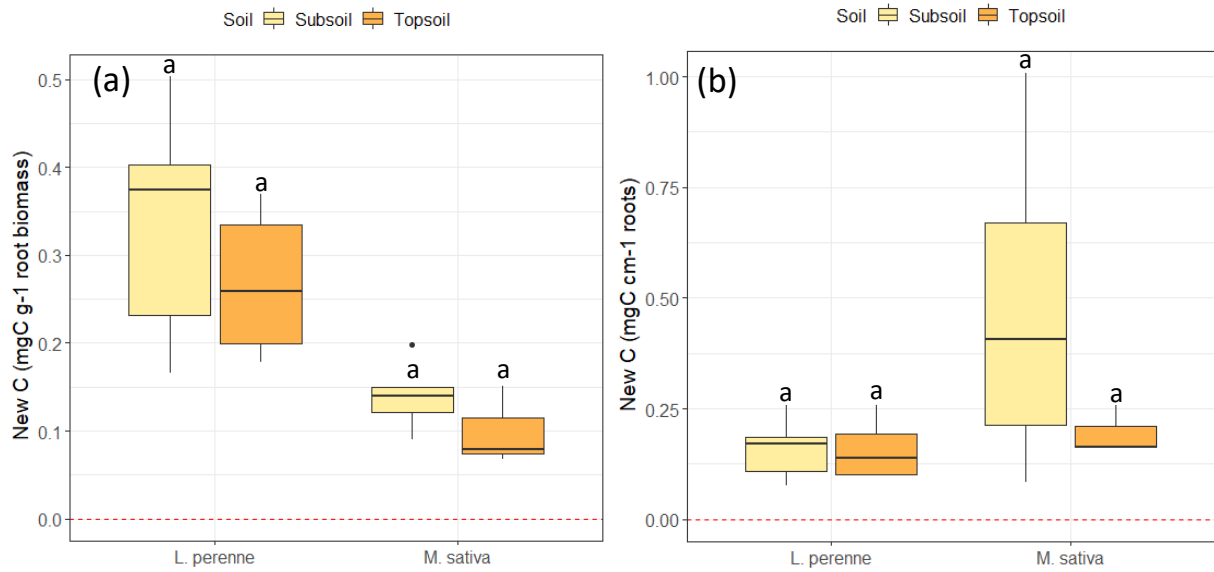
Code	Substrate	Interest and biological properties	Reference	Formula
Carbohydrates :				
GLU	D-glucose	Potential comparison with SIR and cellulose compound	Reactive to different soils	C ₆ H ₁₂ O ₆
XYL	Xylan	Compost of hemicellulose, changes strongly according to the seasons	Reactive to different soils	(C ₅ H ₈ O ₅) _n
CEL	Cellulose	plus dur à dégrader et comparaison potentielle avec résultats DCP	Bérard	(C ₆ H ₁₀ O ₅) _n
Amino acid				
ASP	L-Asparagine	Reactive to different soils		C ₄ H ₈ N ₂ O ₃
SER	L-Serine	Reactive to different soils		C ₃ H ₇ N ₁ O ₃
LYS	L-Lysine	Reactive to different soils		C ₆ H ₁₄ N ₂ O ₂
GLY	Glycine	responds well when decomposed results Berard, precursor ac uric	Bérard, article	C ₂ H ₅ NO ₂
GLUT	L-Glutamine	Linked to the metabolism of nitrogen and ammonia fixation on glutamic acid	données terrain	C ₅ H ₁₀ N ₂ O ₃
Nac	N-acetyl glucosamine	Chitin monomer, found in insects exoskeleton and fungi	Dalmonech	C ₈ H ₁₅ NO ₆
Carboxylic acid (more recalcitrant)				
OX	Oxalic acid	Root and exudates component linked with Malic acid	From field trial, Dalmonech, Bérard	C ₂ H ₂ O ₄
UR	Ureic acid	extruded by isopods (and diplopods)	From field trial, Dalmonech, Bérard	C ₅ H ₄ N ₄ O ₃
MAL	Malique acid	Root and exudates component, useful in fermentation processes		C ₆ H ₆ O ₅
Phenolic acid (strongly recalcitrant)				
CAF	Cafeic acid	Close to rosemalic acid (extruded by Lamiaceae)	Bérard	C ₉ H ₈ O ₄
SYR	Syringic acid	produit de la dégradation de pigments végétaux la malvidine	Dalmonech	C ₉ H ₁₀ O ₅
VAN	Vanillic acid	produit de dégradation de la lignine par les champignons	Oren	C ₈ H ₈ O ₄



962

963 **Figure S4:** Scheme representing the organization of the MicroResp detection plate, showing the three replicates for each
 964 substrate used, the three for the H₂O control and the separation of 2 replicates analyzed in the same MicroResp system.

965



966

967 **Figure S5:** (a) New C moved in the SILT+CLAY fraction in subsoil (light orange) and topsoil (orange) for g of root biomass for the
 968 two different treatments (*L. perenne*, *M. sativa*) in 183 days of experiment. (b) New C moved in the SILT+CLAY fraction in subsoil
 969 (light orange) and topsoil (orange) for cm of root for the two different treatments (*L. perenne*, *M. sativa*) in 183 days of experiment.
 970 , for each boxplot, the lower edge of the box corresponds to the 25th percentile data point, while the top edge of the box
 971 corresponds to the 75th percentile data point; the upper and lower vertical lines corresponds to the 90th and 10th percentile data
 972 points, respectively; the horizontal line within the box represents the median and black dots indicate outliers. Letters above the
 973 boxplots indicate statistically significant differences ($p < 0.05^*$) between species and controls according to Tukey HSD test.

974

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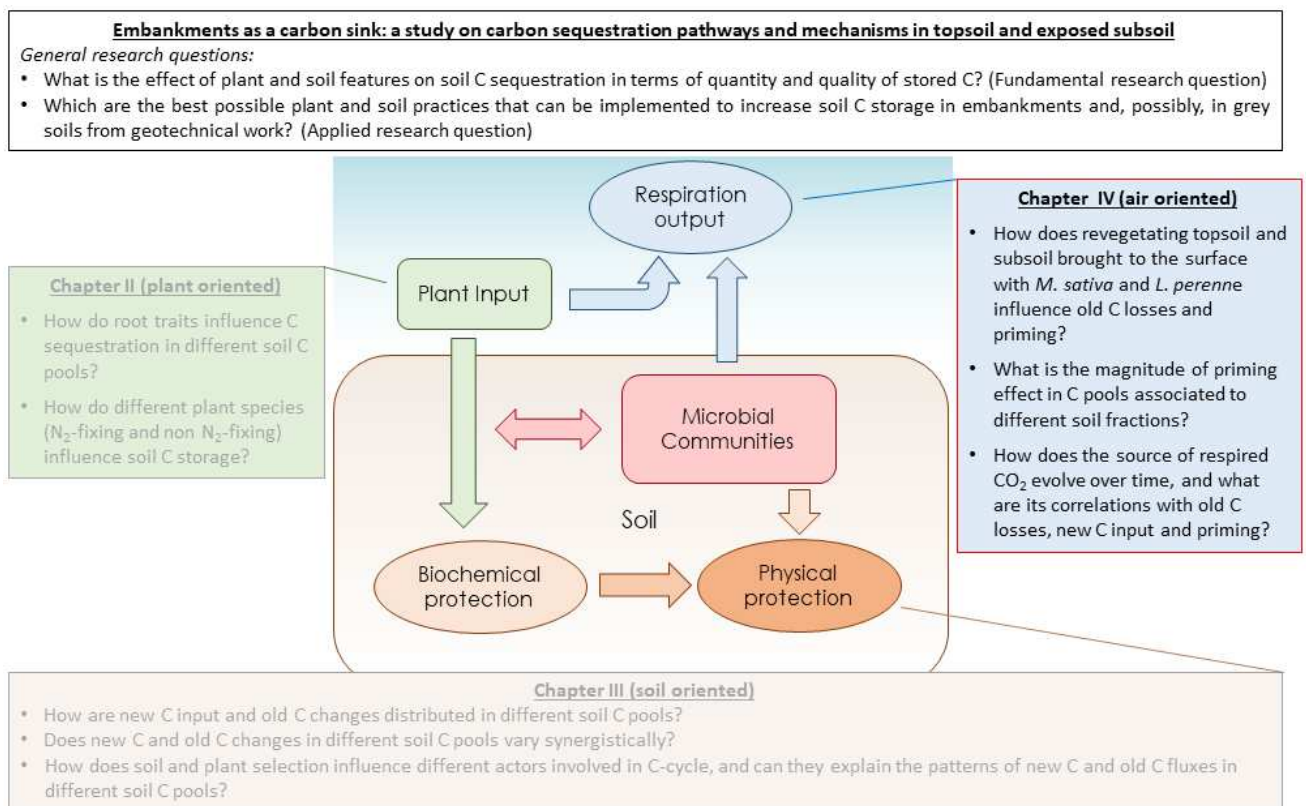
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Chapter IV: Soil quality drives the priming effect and plant species refine it



In Chapters II and III we investigated the C storage in different C pools and its relationship with soil, root and microbiological characteristics. However, vegetation can also have a negative effect on soil C storage increasing the preexistent old C mineralization and loss compared to bare soil - the so called 'priming effect'. In Chapter IV we aim to tackle this aspect, and study the how plant species and soil showing contrasting characteristics influence the priming effect, analyzing soil respiration origin and changes in soil old C in bulk soil and different soil C pools, and describing the main factors that influence our plant-soil system: microbiological substrate preference and N competition.

4.1. INTRODUCTION

Using vegetation to increase soil C sequestration in soil is recognized as an efficient method to mitigate atmospheric CO₂ content. Accordingly, national and supranational organizations designed international programs to promote C storage in soil e.g., the 4p1000 initiative (www.4p1000.org, Minasny et al. 2017). The net input of C in an ecosystem is jointly determined by fresh biomass input of new C via plant biomass turnover and labile rhizodeposition (new C) and losses of old preexistent C in soil (old C) (Kuzyakov and Domanski, 2000; De Deyn et al., 2008; Lal 1994; Smith et al., 2000; De Deyn et al., 2008). Both processes are not independent, but can have strong interactions. Especially, concerns about the fate of the preexisting old C loss due to such a fresh new C input from plants have been raised increasingly. Such a phenomenon, called the priming effect (PE), refers to input of labile C from plants that can accelerate (positive PE) or decelerate (negative PE) mineralization and losses of preexistent old C from soil (Kuzyakov et al., 2000, Fig. 1).

The PE depends on the nature of the substrate consumed by the soil microorganisms, which could be altered by plant and soil conditions (Hamer and Marschner, 2005; Shahzad et al., 2015; Huo et al., 2017). Therefore, understanding the impact of plant and soil characteristics on the amount of primed C has become a key research objective. Among the diverse factors affecting C sequestration, soil particle size and associated C pools is considered as a major factor affecting PE. The commonly used classification of soil C pools associated to soil particle size fractions in literature refers to four C pools. First C in particulate organic matter (POM) (C_{POM} , 2000-200 μ m), and C in fine POM ($C_{finePOM}$, 200-50 μ m) originating from plant litter debris at different levels of degradation, and less protected from degradation (Kögel-Knabner, 2002). After the pools more stable due to their organomineral binding with fine soil particles: C in coarse SILT (C_{SILT} , 50-20 μ m) and C in fine SILT + CLAY ($C_{SILT+CLAY}$, <20 μ m), deriving from exudation and processed C from microbiological communities in form of exudates, exopolysaccharides and microbial necromass (Sollins et al., 1996; von Lützow et al., 2006; Cotrufo et al. 2013, Liang et al., 2017). For example, Huo et al. (2017), found that rhizospheric PE was significantly greater in finely grained

24 soil. Surprisingly, the effect of soil particle size fraction on PE has seldom been investigated (but see Ohm et al.
25 2007), nor the interactions with soil type and plant species. In a framework of soil C sequestration, both soils that
26 differ in initial old C quantity in different soil C pools, and species of contrasted root growing strategies, have been
27 shown to have significant effects on the sequestration of C in soil and relative size of C pools. We argue that, to
28 better understand the effects of revegetation on C-cycle and storage, even the influence of soil and species on
29 priming effect needs to be examined, considering the changes of old C in different C pools related to soil particle
30 size fractions.

31 To quantify the PE, knowing the fate of old C in soil is essential and usually performed using a stable isotopic
32 labelling approach (Kuzyakov, 2006). Monitoring respired CO₂ reflecting the source of the consumed substrate and
33 the remaining old C changes in soil are two major means to assess the PE. When plants are growing in an
34 atmosphere constantly enriched with ¹³C atmosphere, the input of plant derived C has a higher abundance of ¹³C.
35 Consequently, the abundance of ¹³C in respired CO₂ (A¹³C, %) depends on the mineralized C source: a higher A¹³C
36 if the main respiration source is the consumed plant new C input, and a lower A¹³C if the source is the preexistent
37 old C in soil (e.g. Fontaine et al. 2004, 2007; de Graaf et al. 2010). Another effective way to study priming effect is
38 assessing the losses of preexistent old C in soil with and without vegetation. When analyzing the C in bulk soil, the
39 ¹³C signal also allows the differentiation between preexistent old C, and fresh new C derived from plant input.
40 Comparing losses of old C in a vegetated soil allows for the quantification of the priming effect and whether it is
41 positive or negative PE.

42 On newly constructed road embankments, subsoil is increasingly used to replace topsoil that is stripped off during
43 the construction process. Compared to topsoil, subsoil contains less C, but the old C present is more stable than
44 in topsoil for several reasons. C in subsoil is associated with the finest soil particles and stabilized via organomineral
45 interactions (i.e., SILT and SILT+CLAY) (Eusterhues et al., 2005; Chabbi et al., 2009). Subsoil has less microbial
46 biomass (Taylor et al. 2002; Andersen and Domsche 1989; Ekklund et al. 2001), and activity due to oxygen

47 limitation (Rumple and Kögel-Knabner, 2010), and reduced plant inputs (Fontaine et al. 2007) increasing C
48 residence time. Finally, physical separation of microbes and C decrease the possibility for C mineralization (Von
49 Lützow et al. 2006; Holden and Fierer 2005). Subsoil excavation, mixing and revegetation alter all of these
50 protection mechanisms. How the revegetation influences PE and the fate of old C in subsoil, especially in the
51 SILT+CLAY pool remains unclear. To the best of our knowledge, no studies on the priming effect of subsoil brought
52 to the surface have been performed.

53 We aim at comparing the priming processes in two soils with same origin but contrasting characteristics (topsoil
54 with typical fertility, high microbial biomass and nitrogen (N) content versus subsoil with low fertility, microbial
55 biomass and N content). The soils were vegetated with two herbaceous species: the di-nitrogen (N₂) fixing species
56 *Medicago sativa* L., and the non N₂-fixing grass *Lolium perenne* L.. Soil respiration, changes in new C, old C and the
57 priming effect for total C and that in each C pool were quantified. We hypothesize that (i) topsoil will have higher
58 losses of old C due to greater microbial biomass and activity; however, (ii) subsoil will have a greater positive
59 priming effect because it is very highly disturbed compared to the initial conditions, and (iii) the C priming effect
60 will differ among soil fractions.

61

4.2. METHODOLOGY

4.2.1. *Experimental setup*

Soil used for growing plants was excavated from Pisciotta (Italy, 40°07'N 15°14'E/40.116667°N) at two depths of the same soil profile: topsoil (0-30cm depth) and subsoil (110-140cm depth). The soil is a clay loam soil (USDA) with a slightly different granulometry between topsoil and subsoil (topsoil: 27.3% clay, 31.1% silt, 41.6% sand; subsoil: 34.8% clay, 36.8% silt, 28.4% sand). The pH in topsoil was 7.0 and in subsoil was 8.4.

Air dried soil was crushed and sieved to 5mm to homogenize it. We mixed and divided the soil in four sections (quartiles) and 36 different pots were prepared collecting one scoop of soil from each section until the desired weight in each pot has been reached (Fig. S1). We added 6.9 kg of soil into each pot. During the preparation, three soil samples were removed and put aside for chemical analyses. These samples represent the initial soil, or time zero (t₀). Inside each pot, a 60 mm deep plastic ring with a diameter of 80 mm was fitted that could be closed with an airtight dome for subsequent measurements of soil respiration (Fig. S2).

N₂-fixing *Medicago sativa* L. and non N₂-fixing *Lolium perenne* L. were sowed as monocultures with exactly the same pattern. In each pot, three seeds were put at six equidistant spots. After germination, one seedling was removed with scissors at ground level, at each spot. For each soil type (i.e. top- and subsoil) and species, six replicate containers were prepared and six bare containers per soil type were used as controls (n = 36 in total)

Containers were placed into three identical microcosms at the Ecotron growth facility at Montpellier, France (<http://www.ecotron.cnrs.fr/>) (Fig. S3). In each microcosm, two replicates of all treatments, i.e., 12 pots, were placed randomly to avoid any effect of microcosm on plant growth and soil processes. Plants were grown at a constant air temperature of 21°C and at 80% humidity (to reduce the soil water loss by evapotranspiration). Artificial light was provided by three lamps (Gavita PRO 300 LEP 02, Netherlands) in each microcosm with a 12h day/night cycle, shifted to allow air sampling at the same moment of the plant's circadian rhythm (data not shown)

85 in this study, Fig. S4). A shade was placed on the lamps and the distance of the lamps from the plants was adjusted
86 to achieve the most possible homogenous light intensity on the foliage ($300 \mu\text{mol m}^{-2} \text{s}^{-1}$). Soil moisture was kept
87 at $45 \pm 10\%$ of the soil water holding capacity for the entire duration of the experiment. A system of plastic pipes
88 was installed in the chamber to allow irrigation without having to open the chamber and disturb the ^{13}C
89 concentration (Fig.S5). Pots were irrigated every 2-3 days, according to their evaporation rate. However, with the
90 growth of plants, the increase in biomass and in transpiration had to be considered to calculate the amount of
91 water needed to keep the soil at the desired moisture content. For this reason, every 2 weeks, pots (in
92 correspondence with the air sampling) were removed from the chamber, weighted and randomized inside the
93 chambers. Knowing the amount of water added in the previous 2 weeks, the initial soil moisture content, and the
94 final soil moisture content, we were able to calculate the increase in evapotranspiration every 2 weeks and adjust
95 the amount of water needed (data not shown).

96 After the germination of seedlings, the atmosphere was enriched with ^{13}C , reaching a concentration of 2%
97 (approximately two times higher than the natural ^{13}C abundance of 1.1%, in other words $\delta^{13}\text{C}$ of CO_2 in the chamber
98 was roughly +760, as compared to the ambient -8). The air enrichment with ^{13}C was infused during the photoperiod
99 and the ^{13}C infusion stopped during the night period. The experiment was carried out for 183 days, starting the 29
100 September 2017 and ending the 31 March 2018.

101

102 4.2.2. *Air sampling*

103 Air sampling rings were built with two openings in their belowground section to allow root growth in their
104 perimeter, and a double ring structure (one inside the other) that could be filled with water. Inserting the plastic
105 dome inside the double ring structure filled with water ensures an airtight sealing, allowing soil respiration to
106 accumulate inside the chamber (the plastic ring cover had an area of 8.5cm^2 and a height of 6cm, for a volume of
107 340cm^3) (Fig. S6). Every two weeks, we assessed the percentage of ^{13}C in the respired CO_2 .

108 To conduct the air sampling, ¹³C enrichment was stopped 24h before the sampling procedure to allow the ¹³C
109 accumulated in soil to leak out of macropores and cracks that could pollute the results.

110 The day of the sampling, each chamber was open and the air sampling of the time 0 (t0) was performed as soon
111 as the photoperiod stopped. The protocol for the air sampling consisted of:

- 112 1. Pour water in the ring (Fig. S7a)
- 113 2. Close the ring with the plastic dome (Fig. S7b)
- 114 3. Insert the syringe in the rubber cap of the plastic dome and collect 5ml of air to set the reference time 0 (t0)
115 (Fig. S7c)
- 116 4. Immediately transfer the collected 5ml sample from the syringe to an Exitainer under vacuum to store gas (Fig.
117 S7d)
- 118 5. After 2 hours of incubation, without moving the dome, insert the syringe in the rubber cap of the dome, collect
119 5ml of gas enriched with the soil respiration, and transfer it in the exitainer following steps 3 and 4. This sample
120 will represent the Time 1 (t1) air sample, as the amount of CO₂ in the chamber atmosphere after a 2h
121 incubation period.
- 122 6. Samples analyzed with an elemental analyzer Isoprime100 coupled with an Elementar Varo Isotope Cube

123 Results from the isotope analyzer provided the CO₂ concentration in ppmV at time 0 (CO_{2t0}) and time 2 hours
124 (CO_{2t1}). The abundance of ¹³C in respired CO₂ was given in δ¹³C, according to the equation:

$$125 \quad \delta^{13}C = \frac{R_{sample}}{R_{VPDB}} - 1 \quad [1]$$

126 Where R_{sample} is the carbon isotope ratio of the sample (¹²C/¹³C) and R_{VPDB} the ratio of the international standard
127 reference Vienna Pee Dee Belemnite (R_{VPDB} = 0.0111802, Werner and Brand, 2001). The δ¹³C was adjusted
128 according to the CO₂ concentration of the analyzed samples as δ¹³C_{SR}:

$$129 \quad \delta^{13}C_{SR} = \frac{(CO_{2t1} * \delta^{13}C_{t1}) - (CO_{2t0} * \delta^{13}C_{t0})}{(CO_{2t1} - CO_{2t0})} \quad [2]$$

130 Where $\delta^{13}C_{t1}$ is the isotopic composition of CO_2 at after 2 hours of soil respiration and $\delta^{13}C_{t0}$ the isotopic
131 composition at time 0.

132 To calculate the fractional abundance of ^{13}C in the respired CO_2 ($A^{13}C$), first the carbon isotope ratio was derived
133 from [1] as follows:

$$134 \quad R_{sample} = \left(1 + \frac{\delta^{13}C_{SR}}{10^3}\right) * R_{VPDB} \quad [3]$$

135 Finally, to calculate the isotope abundance $A^{13}C$ (%):

$$136 \quad A^{13}C = R_{sample} / (1 + R_{sample}) \quad [4]$$

137 To calculate the percentage of CO_2 derived from fresh plant input mineralization (fPlant) first the soil derived CO_2
138 concentration (CO_{2c}) in μmol have been calculated as:

$$139 \quad CO_{2c} = \Delta CO_{2P} \times \frac{P \times Vc}{R \times T} \quad [5]$$

140 Where ΔCO_{2P} is the CO_2 concentration in (ppmV) is the difference of CO_2 concentration in the sampling chamber
141 (in ppmV) at time 0 and after 2h of incubation time; P the atmospheric pressure in Pa; Vc the volume of the
142 chamber in m^3 ; R the ideal gas constant $8.314 J K^{-1}$; and T the temperature in K.

143 After the amount we calculated the quantity C(C_Q in μg) in the respired CO_2 as:

$$144 \quad C_Q = (CO_{2c} * 12 * (1 - A^{13}C)) + (CO_{2c} * 13 * A^{13}C) \quad [6]$$

145 Where 12 and 13 are the atomic weight of ^{12}C and ^{13}C . The ^{13}C amount ($^{13}C_Q$ in μg) in the respired CO_2 have been
146 determined as:

$$147 \quad ^{13}C_Q = C_Q \times A^{13}C \quad [7]$$

148 Where the $E^{13}C$ is the excess of ^{13}C (in %) compared to the bare soil control at the beginning of the experiment t_0
149 ($E^{13}C = A^{13}C$ at time $x - A^{13}C$ of bare soil control at time 0, equal to 1.076 in topsoil and 1.082 in subsoil). After the
150 plant derived C (pC in μg) was calculated as:

$$151 \quad pC = \frac{{}^{13}C_{\mu g}}{E^{13}C_{atm}} \quad [8]$$

152 Where $E^{13}C_{atm}$ is the excess of ^{13}C in the chamber atmosphere (average of +0.8%). Finally, to calculate the
153 percentage of C in CO_2 deriving from plants C input (f_{plant} , in %):

$$154 \quad f_{plant} = \frac{pC}{C_{\mu g}} \quad [9]$$

155

156 **4.2.3. Soil and biomass sampling**

157 The volume of soil in each pot (20 x 20 x 10 cm³) was divided in two halves vertically with a saw. One half was air
158 dried and used for the soil analysis and microbial measurements, and the other half was used for the measurement
159 of plant traits. Plants were cut at the root collar to divide aboveground and belowground biomass. The resulting
160 mixed sample of soil and roots were placed on a 2 mm sieve and carefully washed to disperse the soil, and the
161 plant individuals were divided (if possible). Above- and belowground biomass was collected, oven dried at 60°C,
162 and weighed to determine dry weight.). Following McCormack *et al.* (2015), we visually separated and sampled
163 transporting (long, thick, high-order roots (>3) and absorptive roots (short, thin, low-order roots 1 – 3),, finely
164 ground and analyzed with an Elementar Varo Isotope Cube to determine their C and $\delta^{13}C$ signal.

165 The soil half used for soil sampling was subsequently divided into shallow soil (0-3.5 cm) and deep soil (3.5-10 cm).
166 Deep soil was air dried, crushed, mixed, and divided into four sections. One 5 ml scoop from every section was
167 collected to form a composite subsample, then sieved at 2 mm. Three subsamples were collected for each replicate
168 pot, and analyzed with an elemental analyzer Isoprime100 coupled with an Elementar Varo Isotope Cube, to
169 determine C content, nitrogen (N) content and $\delta^{13}C$ signal. Samples collected at time 0 and at the end of the

170 experiment (after 183 days, t_6) were analyzed. The difference between t_0 and t_6 gave the changes in C after 6
171 months (ΔC).

172 Simultaneously, 40g of soil from the same deep layer of the pot of the bulk soil samples were collected and
173 fractioned after Gavinelli et al. (1995) (See Chapter III: Method S1, Figure S1). The resulting five fractions (POM:
174 $>200\mu\text{m}$, finePOM: $200-50\mu\text{m}$, SILT: $50-20\mu\text{m}$, SILT+CLAY: $<20\mu\text{m}$) were analyzed for C and $\delta^{13}\text{C}$ with an elemental
175 analyzer Isoprime100 coupled with an Elementar Varo Isotope Cube. The sum of C in different fractions represents
176 the total C in the fraction sample (SUM). A subsample of 0.1 g was taken from each 40 g sample and analyzed
177 without fractioning to determine the total C in the bulk sample. The difference between total C in bulk soil and the
178 sum of C in the different soil fractions was used to assess the correctness of the fractionation and was equal to
179 93.3%.

180 The increased $\delta^{13}\text{C}$ signature of the atmosphere in the microcosm chamber, allowed the calculation of the
181 proportion of C stored in soil directly derived from the input of *M. sativa* and *L. perenne* (new C), to differentiate
182 it from the carbon already present in the soil at the beginning of the experiment (old C). To calculate the
183 proportion, an isotope mixing model (after Balesdent and Mariotti 1996) was used:

$$184 \quad \%C_{new} = \frac{\delta(t_1) - \delta(t_0)}{\delta_B - \delta(t_0)} \quad [10]$$

185 Where %Cnew is the percentage of new carbon in the measured SOC of a specific fraction, $\delta(t_1)$ is the $\delta^{13}\text{C}$
186 signature of the SOC measured in a specific fraction at the end of the experiment (t_1), $\delta(t_0)$ is the $\delta^{13}\text{C}$ signature
187 of the SOC of a specific soil fraction before the experiment (t_0), δ_B is the $\delta^{13}\text{C}$ signature of the new C input in the
188 system, in our case, the signature of the root biomass (i.e., the mean signature of absorptive and transport roots).
189 The choice of root biomass as the $\delta^{13}\text{C}$ reference for C input was made since root material was considered the
190 main source of fresh C, given that litter was removed every two weeks from the pots. Multiplying the total soil C
191 by %Cnew provides the amount of new C in mgC g^{-1} soil.

192 Since the changes in carbon in the system (ΔC in mgC g^{-1} soil) are composed of the two fluxes: input of new C
193 (ΔC_{NEW}) and changes in oldC, the changes of preexistent carbon in soil (ΔC_{OLD}) were calculated as:

$$194 \quad \Delta C_{OLD} = \Delta C - \Delta C_{NEW} \quad [11]$$

195 The effect of vegetation regarding the losses of old C (priming, PE in mgC g^{-1} soil) was calculated as:

$$196 \quad PE = \Delta C_{OLDV} - \Delta C_{OLDBS} \quad [12]$$

197 Where ΔC_{OLDV} is the change in old C in the vegetated soil fraction (in mgC g^{-1} soil), while ΔC_{OLDBS} is the mean of old
198 C changes in the bare soil controls (in mgC g^{-1} soil). If priming has positive values (positive PE) – it means that
199 vegetation increases old C mineralization, with the value corresponding to an increased loss of old C in vegetated
200 soil compared to bare soil. Likewise, if priming has negative values (negative PE), it means that vegetation
201 decreases old C mineralization and losses, with the value corresponding to a decrease in old C loss in vegetated
202 soil compared to bare soil.

203

204 **4.2.4. Statistical analysis**

205 The normal distribution of residues was verified using a Shapiro-Wilk test ($\alpha p = 0.05$). If data were normally
206 distributed, a one way analysis of variance (ANOVA) was performed to test the effects of soil type (topsoil versus
207 subsoil) and plant species (*M. sativa*, *L. perenne*, bare soil) on C priming and $A^{13}C$ in respired CO_2 .

208 If data were not normal a Kruskal – Wallis test was used. Linear regressions ΔC , new C, old C and priming with $A^{13}C$
209 in respired CO_2 were analyzed and R^2 and p values calculated. All the statistical analyses were performed using the
210 open-source statistical environment 'R', version 3.4.3 (R Development Core Team, 2017) using the packages *vegan*
211 and *Hmisc*. (Oksanen et al. 2019, Harrel 2007).

212

213 4.3.RESULTS

214

215 4.3.1. Soil characteristics and changes in carbon content

216 A positive increase in C in bulk soil after six months was found in topsoil planted with *M. sativa* only (ΔC , Table 1).

217 However, on subsoil, a net loss of total ΔC over 6 months was found. In bare soil and soil planted with *L. perenne*,

218 losses of total ΔC were higher in topsoil than subsoil, with the greatest loss in bare topsoil (Tables 1, 2). In bulk

219 soil, new C was significantly greater in topsoil than in subsoil (Tables 1, 2) and regardless of soil type, new C input

220 was always greater in soils planted with *M. sativa* compared to *L. perenne* (Table 2).

221 Based on bulk soil data, the losses of old C in topsoil are the highest in bare soil compared to *L. perenne* and *M.*

222 *sativa*, which are not statistically different (Table 1, Table 2). In subsoil, old C losses were significantly lower than

223 in topsoil, with the most losses in soil planted with *M. sativa*, compared to bare soil and *L. perenne* (Table 1).

224 The losses of old C in SUM of fractions were lower in topsoil and not significantly different among fractions or

225 between species and bare soil (Table 1). In subsoil, SUM was comparable in soils planted with *L. perenne* and *M.*

226 *sativa*, however the losses in bare soil are lower in the SUM compared to bulk soil data.

227 After 6 months we observed that vegetation significantly influenced N content in both subsoil and topsoil when

228 sowed with *M. sativa*, compared to bare soil. Soil N content at t6 for *L. perenne* did not show any significant

229 change compared to that at t0 (Fig. S8).

230

231 4.3.2. Priming effect

232 In bulk soil that had vegetation present, PE was negative in topsoil and there was no significant effect of species

233 (Fig. 2a). In bulk soil subsoil planted with *M. sativa*, PE was positive and old C loss was significantly greater than

234 bare soil, but in subsoil planted with *L. perenne*, old C loss was not significantly different from bare soil and there

235 was no PE (Fig. 2b). In topsoil SUM, PE was negative but was not significant different between plant species,

236 although it had a lower intensity compared to bulk soil (Fig. 2a). In subsoil, the PE in SUM was positive, although
237 there were no differences among species (Fig. 2b). In topsoil fractions, PE was negative except in SILT+CLAY, and
238 no significant differences occurred in fractions between *L. perenne* and *M. sativa*, except in fine POM (Fig. 2a). In
239 subsoil, priming was positive in each fraction, with the highest effect in SILT and SILT+CLAY fractions, and the only
240 difference between *L. perenne* and *M. sativa* occurred in the POM fraction (Fig. 2b).

241

242 4.3.3. Evolution of ^{13}C abundance in respired CO_2 ($A^{13}\text{C}$) over 6 months

243 There was a significant effect of both soil type (Kruskal-Wallis, $p < 0.001$) and vegetation treatment (Kruskal-Wallis,
244 $p < 0.001$) on the abundance of ^{13}C in respired CO_2 ($A^{13}\text{C}$). In topsoil, the abundance of ^{13}C in the respired CO_2
245 increased over 6 months, with the highest $A^{13}\text{C}$ from *L. perenne* (2.02%) and *M. sativa* on 20/02/18 (1.99%), while
246 in bare soil, $A^{13}\text{C}$ was greatest on 09/02/18 (1.96%). In topsoil there was a significant effect of treatment (Kruskal-
247 Wallis, $p < 0.001$) (Fig. 3a,b,c). Over six months, in topsoil planted with *M. sativa*, the abundance of ^{13}C in CO_2
248 increased from $1.55 \pm 0.05\%$ to $1.73 \pm 0.03\%$ (with an increment of +11.2%). In the same period, the abundance of
249 ^{13}C in CO_2 from *L. perenne* increased from $1.51 \pm 0.07\%$ to $1.7 \pm 0.02\%$ (with an increment of +13%), and bare soil
250 increased from $1.43 \pm 0.13\%$ to $1.61 \pm 0.07\%$ (+13%).

251 In subsoil, the highest peak of $A^{13}\text{C}$ was found under *M. sativa* the 20/02/18 ($1.49 \pm 0.07\%$), followed by bare soil
252 on 06/03/18 ($1.41 \pm 0.07\%$) and *L. perenne* on 23/01/18 ($1.37 \pm 0.04\%$) (Fig. 3d,e,f). The effect of treatment was
253 significant (Kruskal-Wallis, $p < 0.001$), with the highest $A^{13}\text{C}$ under *M. sativa*. No significant differences between
254 bare soil and that planted with *L. perenne* were found. In subsoil, $A^{13}\text{C}$ of soil respiration under *M. sativa* increased
255 from $1.31 \pm 0.01\%$ to $1.41 \pm 0.05\%$ (with an increment of +9%) and bare soil from $1.26 \pm 0.01\%$ to $1.32 \pm 0.03\%$ (with
256 an increment of +5%), while *L. perenne* decreased from $1.32 \pm 0.04\%$ to $1.31 \pm 0.02\%$ (with a decrement of -0.2%),
257 however the decrease was not significant (Fig. 3e).

258 **4.3.4. Evolution of ratio of CO₂ derived from fresh plant new C input (f_{Plant})**

259 There is a significant increase of respired CO₂ derived from fresh plant inputs (f_{Plant}) in the vegetated treatments
260 over the 6 month period. Topsoil had a +152% increase of f_{Plant} , and subsoil had a +84% increase. When observing
261 the trend over time, topsoil had a greater increase than subsoil in f_{Plant} (Fig. 4; $p < 0.001$)

262

263 **4.3.5. Correlations between OldC loss, NewC input, priming and $A^{13}C$**

264 In bulk soil, both new C and old changes in soil and the $A^{13}C$ were positively and significantly correlated with the
265 $A^{13}C$ of respired CO₂ (all data grouped together). Data points from different species were clearly segregated, with
266 more changes in new C and old C and $A^{13}C$ in respired CO₂ for *M. sativa* (Fig. 6a,b). In the respired CO₂, $A^{13}C$ was
267 negatively correlated with priming effect (Fig. 6c; Kruskal - Wallis, $p < 0.01$).

268

269 4.4. DISCUSSION

270 We observed a strong influence of soil type on old C stability and soil priming for both total C and that in the
271 SILT+CLAY pool, which masked the effect of plant species. Revegetating topsoil results in a negative priming effect,
272 with a lower mineralization of old C compared to the bare soil control. However, in subsoil, the priming effect
273 depends on the species, with no priming effect under *L. perenne*, and a positive priming effect under *M. sativa*,
274 that increased old C losses compared to the bare soil control. We will tackle these effects separately for the sake
275 of clarity, but they are nonetheless closely linked and the C-cycle depends on a feedback mechanism between soil,
276 plants and microbial communities.

277

278 4.4.1. *Subsoil and topsoil revegetation: identifying the substrate preference of microbial* 279 *communities*

280 The priming effect was highly negative in topsoil, but was marginally positive in subsoil. On topsoil a high input of
281 fresh new C allowed microbiological communities to change the substrate preference for energy and nutrients
282 requirements from old C in soil to fresh new C inputted by plants. This phenomenon seems to support the
283 Preferential Substrate Utilization hypothesis (Cheng, 1996; Cheng and Kuzyakov, 2005). Subsoil had less N than
284 topsoil, resulting in reduced plant development (Chapter III, this thesis), and a consequent lower input of labile C
285 into soil from rhizodeposition. As subsoil is subjected to long-term limitation of nutrients, microbial functioning is
286 decreased, thus promoting the development of oligotrophic communities with high metabolic diversity
287 (Blagodatskaya and Kuzyakov, 2008). This set of conditions makes subsoil a perfect candidate for positive priming
288 effect. Specific dormant microbial groups can be activated by the input of low available substrates, such as oxalic
289 acid, that have a high impact on community shift, and synthesize a broad variety of enzymes that promote old C
290 decomposition and a positive priming (Falchini et al., 2003; Landi et al., 2006; Blagodatskaya and Kuzyakov, 2008).
291 Finally, fungi have been shown to play an important role in C degradation and priming (Panikov 1995; Bell et al.
292 2003; Blagodatskaya and Kuzyakov, 2008). The input of fresh C might activate spore or cysts dormant in subsoil,

293 which can penetrate previously inaccessible micro zones with hyphae development (Blagodatskaya and Kuzyakov,
294 2008). The low amount of labile C input, together with the low N availability, will not be enough to shift the
295 increased microbial metabolism towards labile C input in a significant way, and microbial communities will increase
296 old C mineralization to acquire energy and nutrients (De Graaf et al., 2010). In topsoil, instead, the higher biomass
297 development and the consequent high input of labile fresh C from plants will enhance fungal: bacterial gene copy
298 ratios (Griffiths et al., 1998; Broeckling et al., 2008; Chiginevaa et al., 2009). Our data showed higher microbial
299 abundance and diversity, as well as a higher level of activity (Chapter III, this thesis), corresponding well to such a
300 phenomenon. The higher inputs of labile C in the system may, nevertheless, shift the microbial preferential
301 consumption from preexistent old C to fresh C input (as shown in Fig. 5), and result in a negative priming effect.
302 This phenomenon is well supported by the $A^{13}C$ and percentage of plant derived C in respired CO_2 , that was always
303 higher in topsoil compared to subsoil.

304 Experiments investigating substrate preference or competition mechanisms effect on PE have usually been
305 performed in controlled incubation conditions (e.g., Fontaine et al., 2007; De Graaf et al., 2010). Our study is novel
306 in that plants were grown in different types of non-sterilised soil. Also, we demonstrate that the SILT and
307 SILT+CLAY C pools played a pivotal role in determining the amount and trajectory of PE. Particularly in subsoil,
308 where PE was positive in SILT and SILT+CLAY C pools, thus questioning the point of view that these C pools are
309 highly stable (Chapters II and III, this thesis). However, SILT+CLAY C pools is very reactive to input of new C (Chapter
310 III, this thesis) and Fontaine et al. (2007) showed how input of fresh C in subsoil can increase the mineralization of
311 stable C and lead to positive PE. Our results suggest that SILT+CLAY pool is stable when conditions are not abruptly
312 changed: in topsoil positive PE is present only in POM. An abrupt change of conditions, however, can bring to
313 instability of C associated with fine soil fractions and the resulting PE, as observed in subsoil SILT and SILT+CLAY.

314

315 4.4.2. The impact of plants on the two soil types: competition for nitrogen

316 The effect of vegetation on PE was largely influenced by soil type. We did not observe any significant difference in
317 PE between topsoil planted with *M. sativa* or *L. perenne*. Plant species had more influence on PE in subsoil, despite
318 the disparity between the results from bulk soil and from fractionation. *L. perenne* better mitigated the undesired
319 positive PE than *M. sativa*, especially in bulk soil, where no PE occurred in subsoil sown with *L. perenne*. This result
320 on of plant species effect on PE in subsoil, however limited, is in line with the competition hypothesis (Cheng and
321 Kuzyakov, 2005). When plants are grown on an N poor soil, mineralization of old SOC from microbial communities
322 can be reduced due to more efficient N removal by plant roots, that hinders microbial activity, resulting in a
323 negative priming effect (Cheng & Kuzyakov, 2005). This phenomenon can explain the effect of *L. perenne* on
324 positive PE mitigation in subsoil. Increased rhizodeposition will increase old C consumption and can result in a
325 positive PE. This mechanism is exactly what we observe when planting soil with *M. sativa* which has a higher
326 biomass development than *L. perenne* on subsoil (Chapter III, this thesis), and therefore lowers rhizodeposition
327 (Fu and Cheng, 2002; Cheng et al., 2003; Dijkstra et al., 2006). Moreover, *M. sativa* is associated with *Rhizobium*
328 bacteria that allow fixation of N₂ directly from the atmosphere, and rely less on N mining for growth. In this case,
329 the 'competition effect' is avoided because *M. sativa* can acquire N from a different source. To test this hypothesis,
330 we analyzed the difference in N content in soil between time 0 and time 6 months under the three different
331 treatments and two soils (Fig. S8). We observed that bare soil and *L. perenne* do not significantly differ from t0
332 and between each other. However, *M. sativa* increased the amount of soil N, strengthening our hypothesis that
333 competition for N was decreased. Also, substrates from fresh plants' new C input become available, and it
334 increases microbial activity but do not permit a substrate preference switch, resulting in higher mineralization of
335 old C and a positive priming effect (De Graaf et al., 2010). These results are supported in several studies, where N-
336 rich rhizodeposition is believed to be linked with higher PE (Fu and Cheng, 2002; Cheng et al., 2003; Cheng and
337 Kuzyakov, 2005). In topsoil, such a phenomenon was not observed. We speculate that higher fertility levels and
338 rhizodeposition level (Chapter III, this thesis) mask the competitive effect between roots and microorganisms,

339 providing enough nutrients via plants fresh C deposition to the microorganisms to allow them not to have to rely
340 on mining soil C and compete with plants (Cheng and Kuzyakov, 2005; De Graaf et al., 2010).

341 Such a framework considering both soil and vegetation features incorporate the two hypotheses: the Substrate
342 Preferential Utilization hypothesis and the Competition hypothesis (Fig.6), supporting the reconciliation
343 reconciliation between the two proposed by Cheng and Kuzyakov (2005): in fertile soils, the substrate preference
344 will drive the PE, while in poor soil competition will shape the trajectory and magnitude of PE.

345

346 *4.4.3. The priming effect and its implication in practice*

347 From an applied point of view, when revegetating soil in geotechnical constructions, especially subsoil, it is
348 necessary to consider soil fertility, as it will affect i) biomass development, ii) nutrient competition in soil and fresh
349 substrate availability and, consequently, iii) the priming effect. In nutrient poor subsoils, the use of non N₂-fixing
350 species (e.g. *L. perenne*) will result in a low priming effect. A possible solution to avoid the priming effect when
351 revegetating subsoil (or nutrient poor soils in general) could be to couple inoculation of microbial strains that
352 consume labile C with N fertilization, to increase fertility and nutrient availability, and try to switch the microbial
353 consumption from preexistent oldC to new C.

354

355 **4.5. CONCLUSIONS**

356 We examined the priming effect in a crossed experimental design with two soil types and two plant species. We
357 highlighted the complex interactions between the two sources of factors and demonstrated the importance of soil
358 quality (in terms of N content and microbiological activity and biomass) in determining the trajectory and
359 magnitude of PE over that of plant species. When soil quality is high, such as topsoil, positive PE can be mitigated
360 and negative PE can occur thanks to high fresh new C input. However, in N-poor subsoils, old soil C, especially the
361 stable old C in the SILT+CLAY pool, can be susceptible to the PE, depending on the competition between plants

362 and soil microorganisms. Therefore, plant species could play a non-negligible role in influencing the tendency and
363 magnitude of PE.

364 Our results suggest that topsoil, with higher rhizodepositions, allows microbial communities to switch from
365 consuming old C to new C mineralization, resulting in a negative priming effect. In subsoil, microbes will mine old
366 C for nutrients due to low new C input. Competition for N is fundamental to shape the priming effect, and in poor
367 subsoil, *L. perenne* had no priming effect due to N competition. Therefore, when a soil is severely limited in
368 nutrients the competition effect will be predominant; while when conditions are not so limiting the substrate
369 preference will dominate. These findings are in line with the reconciliation of hypothesis from Cheng and Kuzyakov
370 (2005). The $\delta^{13}\text{C}$ and its correlations with old C losses and priming helps to understand the processes in different
371 soils, but alone this is not enough to investigate the effects of priming. Old C losses of vegetated and bare soil
372 control need to be taken into account, together with stable ^{13}C labeling methods confirming its potential for
373 priming studies.

374

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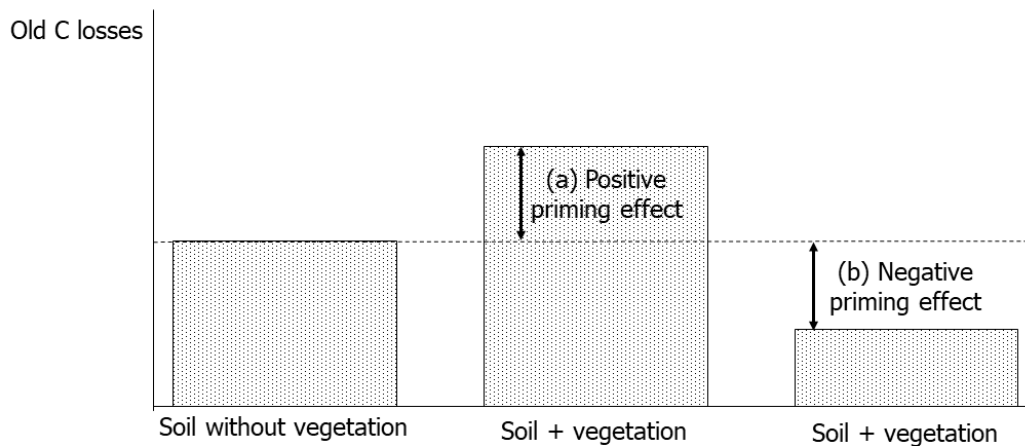
471 FIGURES AND TABLES

472 **Table 1:** ΔC is the difference of C content in bulk soil between time 0 and time 6 months in mgC g^{-1} soil. New C is the input of C
 473 in bulk soil deriving from the vegetation growth during the 6 months of the experiment in mgC g^{-1} soil. Old C is the losses of old
 474 C in bulk soil. The last column shows the losses of old C in the sum of fractions data. ΔC new C, and old C have been calculated
 475 for each treatment (*M.sativa*, *L. perenne* and bare soil) and each soil (topsoil: 0-30cm depth, subsoil: 110-140cm depth). Different
 476 letters next to the average value indicate statistically significant differences ($p < 0.05$) between species or families according to
 477 Tukey HSD tests.

Soil	Data set:		Bulk soil data			SUM of fractions
	Treatment	ΔC (mgC g^{-1} soil)	New C (mgC g^{-1} soil)	Old C (mgC g^{-1} soil)	Old C (mgC g^{-1} soil)	
Topsoil	bare soil	(c) -1.60 ± 0.20	(c) 0.10 ± 0.05	(b) -1.70 ± 0.15	(a) -0.36 ± 0.44	
	<i>L. perenne</i>	(b) -0.48 ± 0.33	(b) 0.68 ± 0.09	(a) -1.16 ± 0.27	(a) -0.36 ± 0.40	
	<i>M. sativa</i>	(a) 0.01 ± 0.41	(a) 1.22 ± 0.16	(a) -1.21 ± 0.29	(a) -0.26 ± 0.02	
Subsoil	bare soil	(b) -0.50 ± 0.07	(c) 0.04 ± 0.03	(ab) -0.54 ± 0.08	(a) -0.24 ± 0.35	
	<i>L. perenne</i>	(a) -0.26 ± 0.08	(b) 0.22 ± 0.03	(b) -0.49 ± 0.06	(ab) -0.59 ± 0.07	
	<i>M. sativa</i>	(a) -0.17 ± 0.18	(a) 0.45 ± 0.11	(a) -0.61 ± 0.11	(b) -0.67 ± 0.23	

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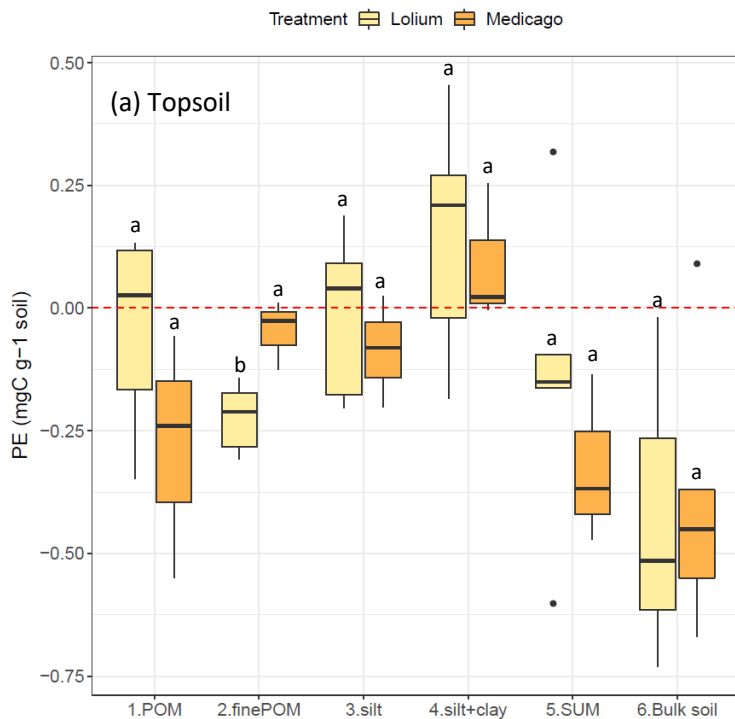
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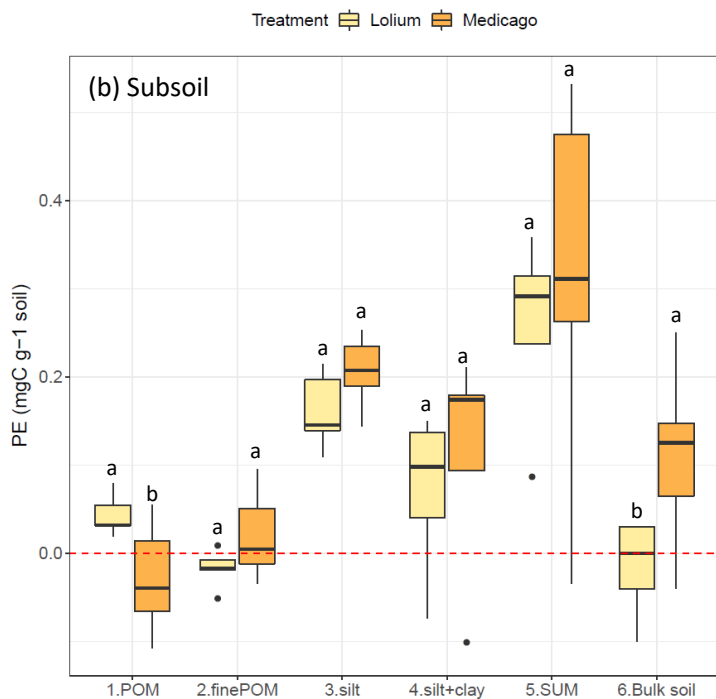
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481 **Figure 1:** Graphic explanation of positive priming effect (a) and negative priming effect (b).

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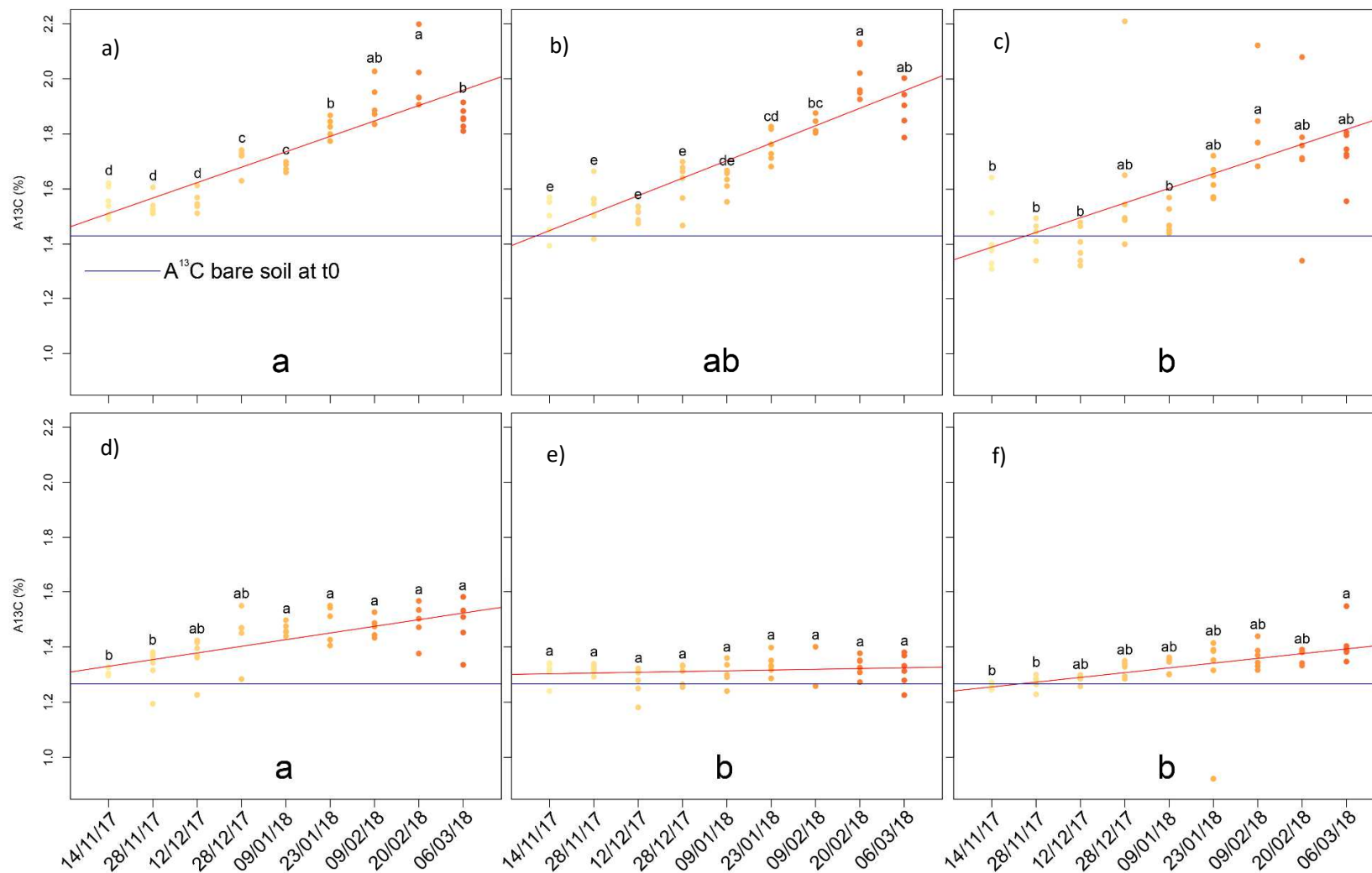
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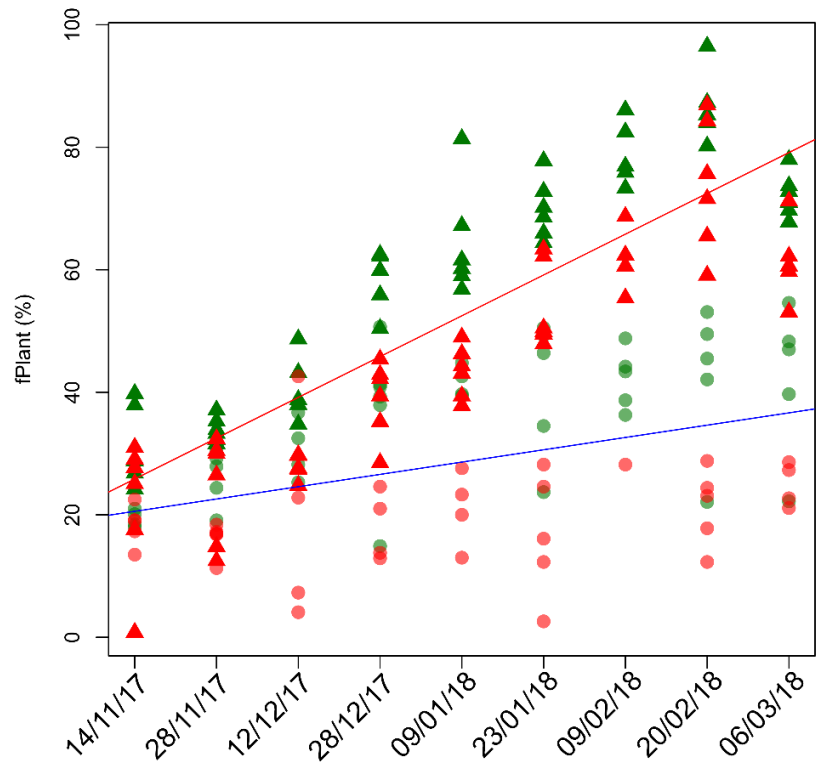
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Figure 2: Comparison of priming effect after 6 months of revegetation between different soils ((a) topsoil and (b) subsoil) for each vegetated treatment (*L. perenne* light yellow, *M. sativa* orange). The data presented shows the priming in the different soil fractions (1.POM, 2.finePOM, 3.silt, 4.silt+clay) the sum of the soil fractions (5.SUM) and the bulk soil data (6.Bulk soil). Negative values means a reduced loss of old C, positive values an increased loss of old C. In each boxplot, the lower edge of the box corresponds to the 25th percentile data point, while the top edge of the box corresponds to the 75th percentile data point. The line within the box represents the median. Different letters above the boxplots indicate statistically significant differences ($p < 0.05$) among species according to a ANOVA test

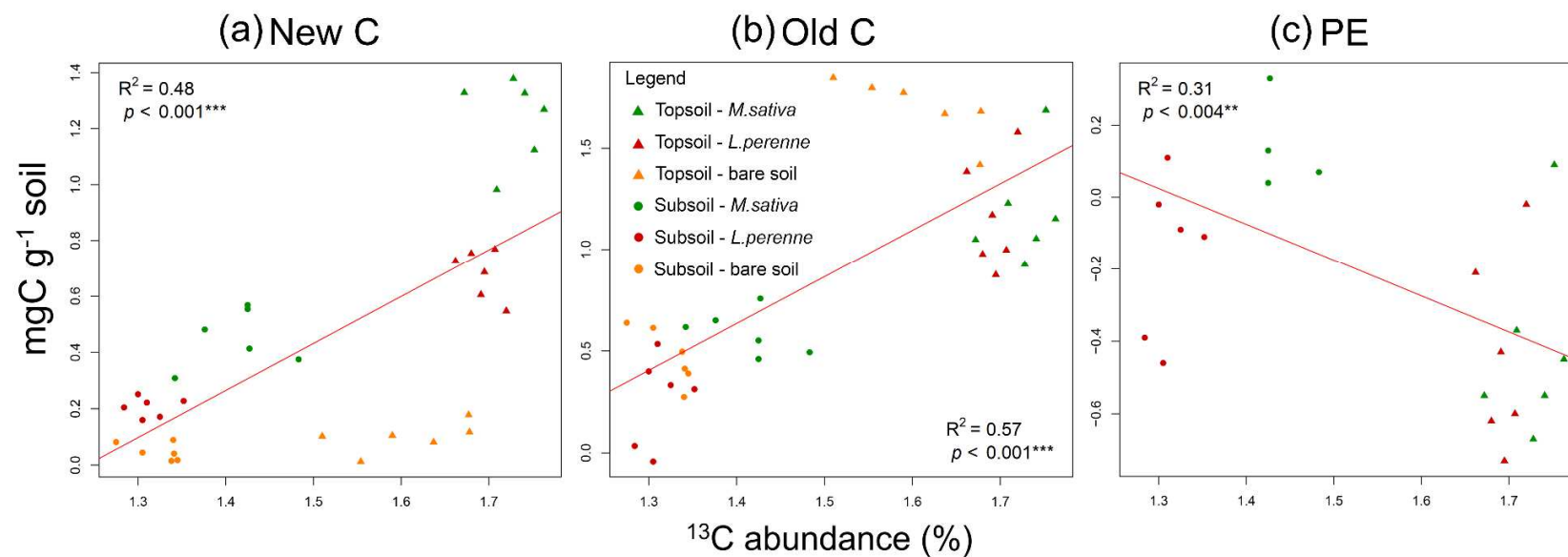


486 **Figure 3:** Evolution over time of A¹³C, as the percentage of ¹³C in the respired CO₂. The results shows the trend for a) topsoil vegetated with *M.sativa*, b) topsoil with *L.perenne* and c) topsoil bare soil control. The second row shows the results for d) subsoil sowed with *M.sativa*, e) subsoil with *L.perenne* and f) subsoil bare soil control. The different dots shows the A13C result for the single subsample of the treatment in shade of yellow/red according to the sampling time. Different letters above the dots indicate statistically significant differences ($p < 0.05$) according to a Tukey HSD test among different sampling dates. Different letters at the bottom of the graph indicate statistically significant differences ($p < 0.05$) according to a Tukey HSD test among different treatments.



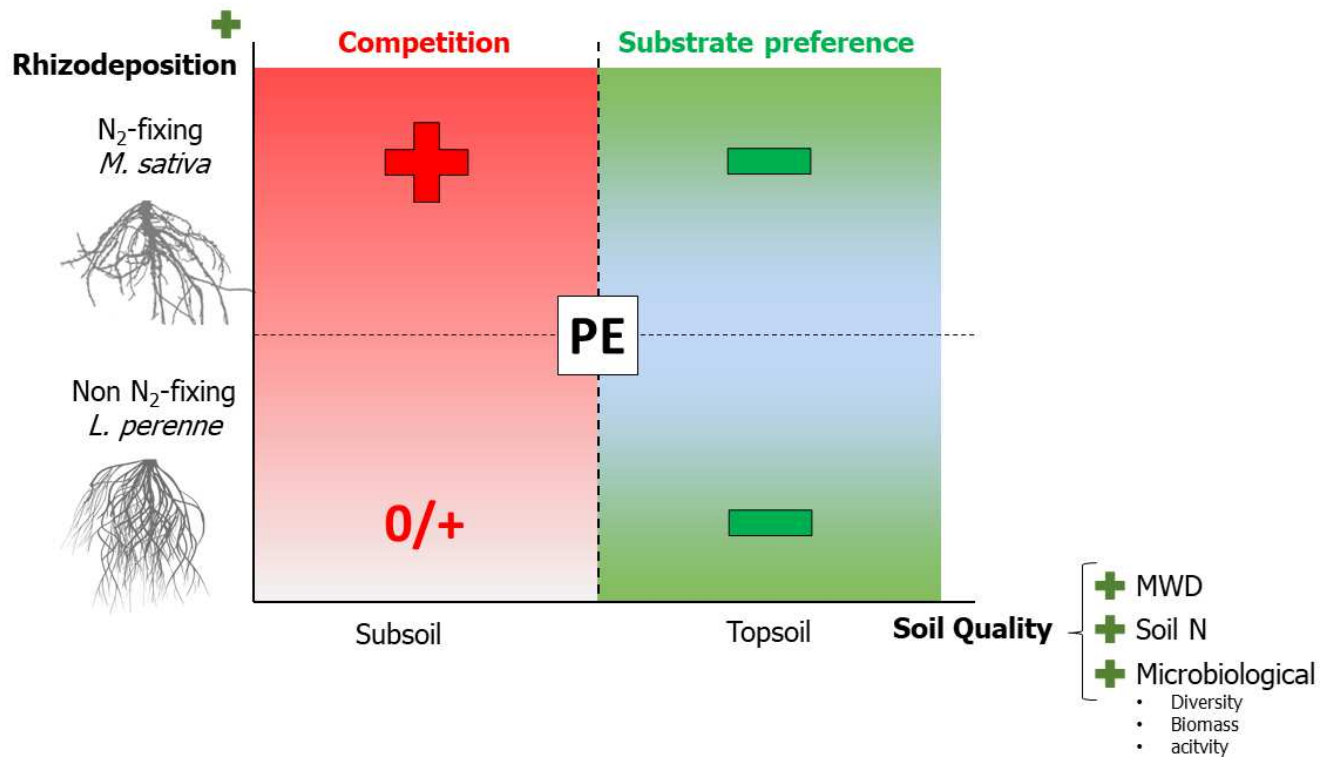
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488 **Figure 4:** Percentage of C derived from mineralization of plant inputs in soil respired CO₂ (f_{Plant}) over 4 months in the two
 489 vegetated soils. Triangles represent topsoil and dots subsoil. Green represent *M. sativa* and red *L. perenne*, different saturations
 490 have been given to better differentiate the points. The red solid line represent the increase according to a linear model of the
 491 f_{Plant} in topsoil over the 4 months, the blue solid line in subsoil. The slopes of the linear models are significantly different, $p <$
 492 0.001.



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Figure 5: Correlations between a) New carbon inputs in soil, b) old SOC C losses (OldC) and c) priming effect (Priming), with abundance of ¹³C in respired CO₂ (A¹³C). Topsoil is represented by triangles, subsoil by dots, *M.sativa* is green, *L.perenne* red and bare soil control orange. The red line shows the correlation according to a linear model, and the significance (R^2 , p) is shown in a corner box in every graph.

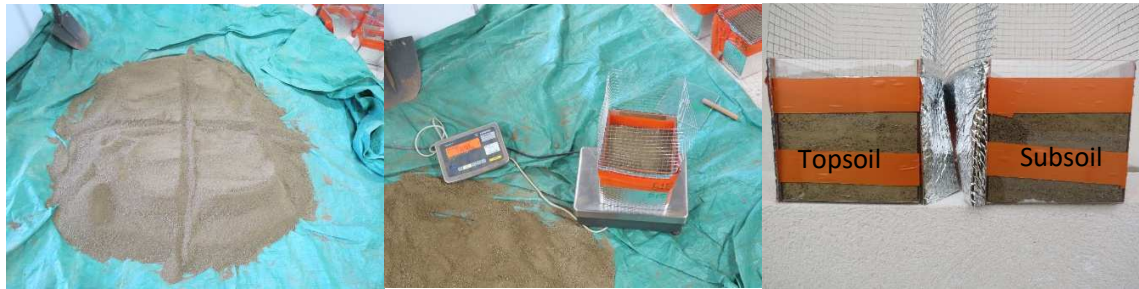


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499 **Figure 6:** Reconciliation of Preferential Substrate Utilization hypothesis and competition hypothesis. a) shows the effect
 500 of soil, where the higher input in the fertile soil allow microbial communities to switch preference of substrate and
 501 decrease old C mineralization, while in subsoil, with low fertility and low input of fresh new C, competition drive the
 502 priming effect that is generally higher than topsoil. b) shows the species effect in the two soil conditions. In subsoil low
 503 rhizodeposition from *L. perenne* stimulate and competition for N hinder old C mineralization and result in a slightly
 504 negative priming effect. In topsoil, the N rich rhizodeposition from *M. sativa* increase the soil N content and decrease
 505 competition, allowing microbial communities to mine more efficiently old C and resulting in positive priming effect. In
 506 topsoil, contrary to what expected, we did not find any difference between *L. perenne* and *M. Sativa* in priming effect,
 507 suggesting a lower influence of rhizodeposition when the system is rich and efficiently colonized by roots.

508

509 SUPPLEMENTARY MATERIALS



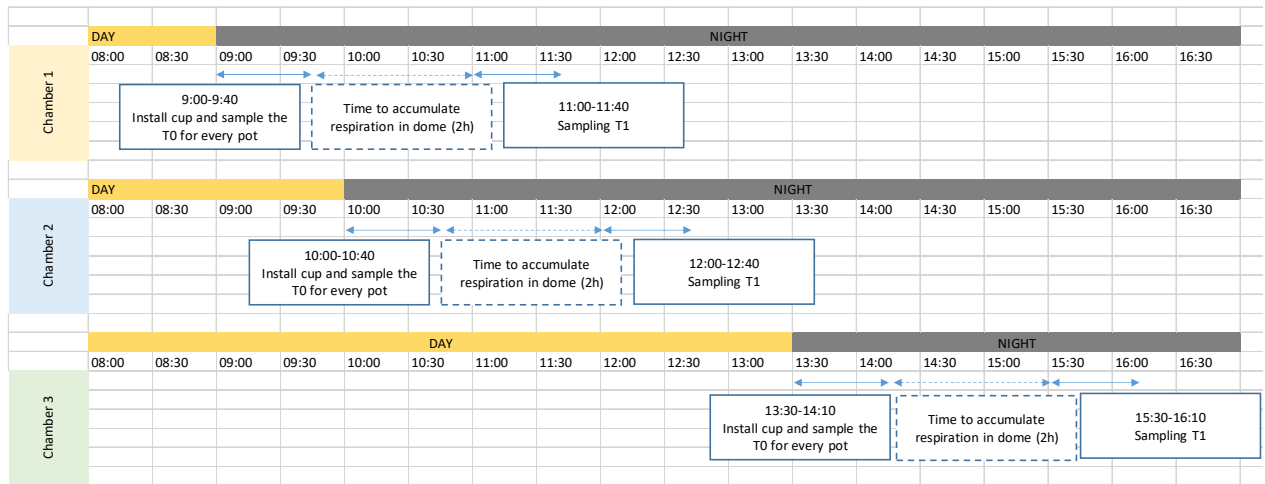
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511 **Figure S1** : Pot preparation. Starting from left to right : Quartile of soil prior to filling the pots, weighting of the pots while
512 filling and examples of two pots filled with the two different types of soil The soil has been added collecting one scoop
513 of soil from each quartile and keep moving to the next quartile, in the same order, until the desired weight was reached.



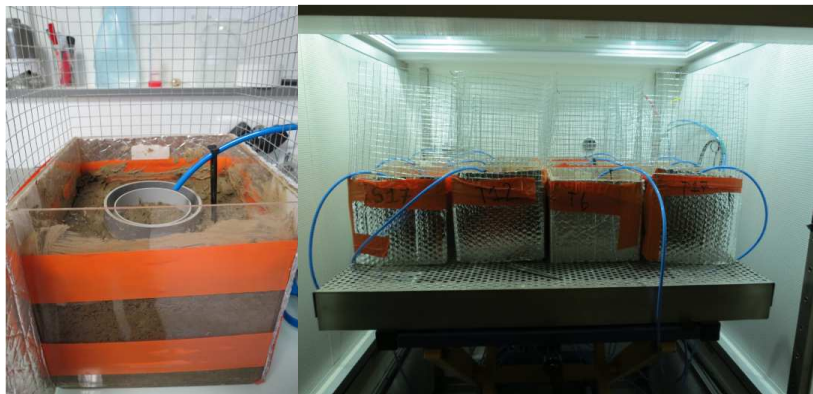
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515 **Figure S2** : From left to right: example of the ring used for soil respiration analysis, ring with the plastic dome in place
516 and ring inside the soil, the red crosses mark the spots were the seeds were planted



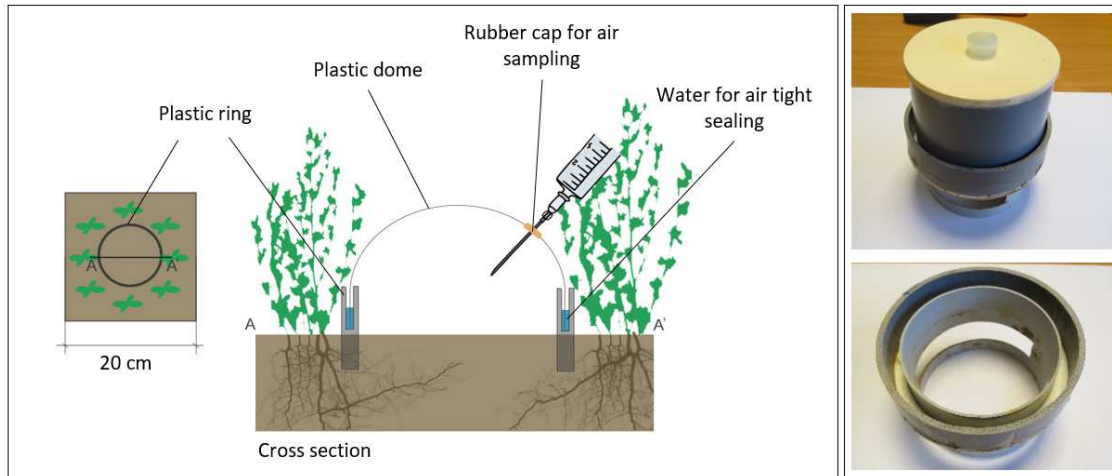
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518 **Figure S3** : Pots positioned in the growing chamber at a fix distance from lights in the Ecotron facilities



519
 520 **Figure S4** : Day and night cycles in the three different grow chambers with timing for air samples collection
 521



522
 523 **Figure S5** : plastic tubes used for irrigation, fix on the single pot on the left and positioned in the chamber on the right
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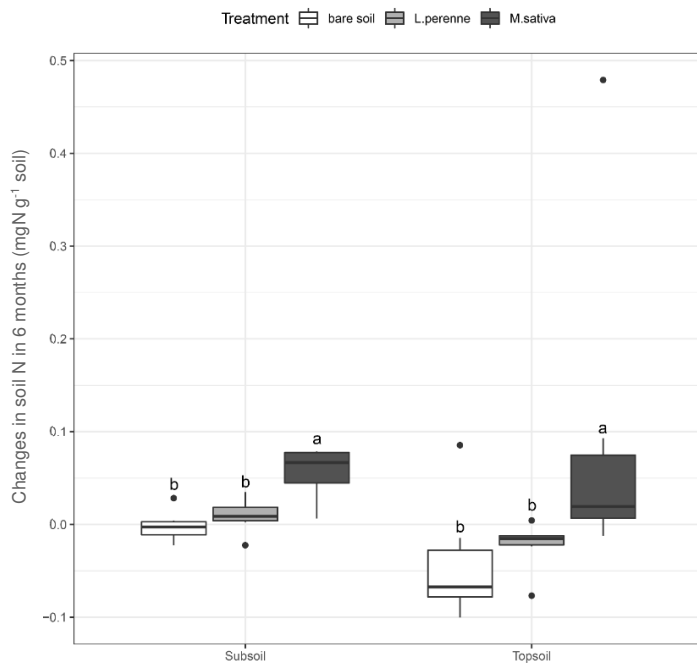
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526 **Figure S6** : Scheme of the plastic dome used for soil respiration analysis



527

528 **Figure S7** : Air sampling protocol : a) fill ring with water, b) place the plastic cap on top, c) take a 5ml air samples using
 529 a syringe on the rubber cap and d) transfer the air taken in the airtight exainers



530

531 **Figure S8:** Comparison of N content in soil between the beginning of the experiment (t0) and the end (t6) in the two
 532 soils (subsoil and topsoil) and the three treatments (*M.satva*, *L.perenne*, and bare soil). No significant differences have
 533 been found between bare soil and *L.perenne*, which show no changes from the initial C content. *M.satva*, instead, is
 534 significantly different compared to the other treatments, showing an increase of soil N. In each boxplot, the lower edge
 535 of the box corresponds to the 25th percentile data point, while the top edge of the box corresponds to the 75th percentile
 536 data point. The line within the box represents the median. Different letters above the boxplots indicate statistically
 537 significant differences ($p < 0.05$) among species and control according to a Tukey HSD test.

538

Chapter V: General discussion

1 5.1. Carbon quality matters: coarse particle pool versus fine particle pool

2 Assessment of C stock of an ecosystem usually considers total soil carbon, not the C in individual fractions.
3 As a result, soil C is presented as a simple number at the plot, catchment, regional or national scale, e.g.,
4 the 4P1000 goal, that considers only total C (Minasny et al., 2017). Yet, the soil C pool is a chemically and
5 physically complex system in which C compounds associated with different soil particle fraction sizes may
6 greatly differ in stability and mean residence time. As a result, increasing attention has been paid to the
7 understanding and characterization of soil organic C quality, here defined as the relative amount of fast-
8 turnover particulate organic matter C fractions (C_{POM} , C_{finePOM}) and stable clay and silt associated C fractions
9 (C_{SILT} , $C_{\text{SILT+CLAY}}$) (Balesdent et al., 1998; Cotrufo *et al.*, 2013; Cardinael et al., 2015). A soil with good C
10 quality should be targeted in C sequestration practices, aiming to have a high proportion of stable C in the
11 SILT+CLAY pool.

12 In both experiments of my thesis, soil particle size fraction related C pools were characterized to assess C
13 quality besides C quantity, and proved to be fundamental in the understanding of the plant-soil system. In
14 Chapter II and III, there was no species effect on the total amount of C stored in soil. However, the quality
15 of C, i.e. its accumulation in different soil fractions, was influenced by the root traits of the different plant
16 species analyzed (Ch. II/III) as well as the soil type chosen (Ch. III), and their effects on microbiological
17 communities. In Chapter IV, applying the concept of 'priming effect' on fraction associated C pools enabled
18 us to highlight that the positive priming in subsoil was mainly due to an increased mineralization of C in
19 the SILT+CLAY pool, while topsoil showed a homogeneous negative priming among pools, allowing us to
20 better disentangle the priming mechanisms in different soil types. All of these findings not only highlight
21 the great importance of looking at C sequestration at the fraction scale, but also challenge the supposed
22 high stability of the SILT+CLAY pool.

23 5.2. Carbon origin matters: new carbon versus old carbon

24 The labelling approach to distinguish new C input in soil has been widely applied and is a relatively new
25 frontier in plant-soil studies examining soil C storage (i.e. Dijkstra and Cheng, 2007; Paush et al. 2013;
26 Haddix et al., 2016). In my thesis, the stable labelling approach during 183 days of plants growth allowed
27 to disentangle plant-soil processes connected with soil C storage, and was an efficient way to study soil C
28 storage. We observed how changes in soil C were mainly attributable to the input of new C. Also, in Chapter
29 III we observed a positive synergy with new C input and old C losses, with higher new C input connected
30 with lower old C losses. This result was in accordance with results from De Graaf et al. (2010), and
31 supported the preference substrate utilization theory (Cheng and Kuzyakov, 2005).

32 Considering new C gains and old C changes in different soil fractions allowed us to further unveil
33 mechanisms of soil C storage that would have been hidden without this double approach of isotope
34 labelling and soil fractionation. In Chapter III there was a high response of POM and SILT+CLAY pools to
35 the input of fresh new C from plants, highlighting the double pathway of new C accumulation in soil, from
36 turnover in POM and exudation or microbial mineralization and deposition in the SILT+CLAY pool (Cotrufo
37 et al., 2013). Moreover, the fractionation allowed us to underline how old C is active in tospoil, being
38 mineralized and accumulating from the coarser fraction to the fine SILT+CLAY pool. These changes in old
39 C among pools would have been hidden by analysis of total C in bulk soil, and old C would have been
40 wrongfully considered inactive.

41 New C and old C in fractions were only studied in Chapters III and IV, but the lesson learnt from these
42 chapters could make us rethink results in Chapter II, in which new C and old C were not distinguished. If
43 not considering old/new C, in both Chapter II and III, we observed that there was no significant effect of
44 species in C storage in different fractions (ΔC ; Chapter III, Fig. 2, 3). However, considering the new C and
45 old C changes in different fractions, we were able to identify the effect of species on the input of new C

46 (Chapter III, Fig. 4), that in ΔC was masked from the changes of old C and its accumulation in the SILT+CLAY
47 pool. Not only distinguishing between old and new C dynamics helped us to understand the effect of
48 species and the different behaviour of soil pools, but even to shed light on the relationships between
49 factors involved in C-cycling and C storage in different pools. Comparing the correlations in Chapter III
50 Table 3 between microbial/soil and root characteristics and new C, old C and ΔC , underlined how new C
51 accumulation was better predicted compared to ΔC (see also Henneron et al. 2019).

52 We, therefore underline the power of coupling the study of C sources with soil fractionation and related
53 C pools, which helped us greatly to analyse the mechanisms behind soil C storage, as explained in the next
54 sections.

55

56 5.3. Microbial community matters: priming and entombing

57 The microbial community is the factor shaping all the processes involved in C cycling. Our main goal was
58 to investigate the influence of species and soil selection on soil C storage in topsoils and exposed subsoils.
59 However, our findings highlight that the role of these two factors always indirectly pass through microbial
60 community biomass and activity via two mechanisms:

61 i) the priming effect (Chapter IV) was the response of microbial communities to revegetation, shaping the
62 losses of C in the system. The productivity of the system (based on soil fertility and plant species
63 performance) will determine the substrate preference of microbial communities and the direction and
64 magnitude of the priming effect (Cheng and Kuzyakov, 2005; De Graaf et al. 2010), as shown in Chapter
65 IV. Moreover, when soil N was low, we found an effect of competition for nitrogen influencing the priming
66 direction and intensity (Cheng and Kuzyakov, 2005). These results support the reconciliation of the

67 Preferential Substrate Utilization Hypothesis with the Competition Hypothesis (Cheng and Kuzyakov 2005),
68 and underline the pivotal role of microbial communities in the priming effect.

69 ii) the quality of C stored due to microbial transformations. The common view that microbial communities
70 reduce C sequestration due to increased soil respiration is becoming increasingly redundant. In chapter II
71 and III we found strong links of microbial activity and biomass with C sequestration in the fine SILT and
72 SILT+CLAY fractions. In this regard, two main theoretical framework have been developed in the last years:

- 73 • Microbial Efficiency-Matrix Stabilization (MEMS) from Cotrufo et al. (2013). The main hypothesis
74 behind this framework states that labile C from plant inputs is the main source for microbial
75 exudates and exopolysaccharides, which are the precursors of stable SOM in aggregates and
76 organomineral compounds.
- 77 • Soil microbial carbon pump (MCP) from Liang et al. (2017). Microbial communities are the main
78 factors shaping the sequestration of C by 1) degrading via *ex-vivo* modifications the stored C,
79 leaving high recalcitrant and persistent SOM in soil, 2) increasing via *in-vivo* turnover the stability
80 of stored C, as microbial necromass and metabolites, i.e., the 'entombing effect'.

81 Microbial necromass was not measured in this thesis. However, in Chapter IV, I provide evidence of the
82 entombing effect. The increase of soil C is mainly due to accumulation of new C in POM and in SILT+CLAY
83 and mineralization of old C in the SILT+CLAY pool. These C increases were correlated with microbial activity
84 and biomass. As predicted by the MEMS model, the quality of input influences the destination of C: with
85 microbial activity enhanced by labile inputs, while recalcitrant input was stabilized via *ex-vivo*
86 transformations. The transformed labile C by microorganisms as exopolysaccharides was then stabilized
87 in the SILT and SILT +CLAY pools – via *in vivo* turnover that promoted 'matrix stabilization' via
88 organomineral interactions, as part of the 'entombing effect'. We, therefore, argue that the MEMS and

89 MPC theories are complementary (Fig. 1), and the use of fractionation enables us to further expand their
90 understanding by quantifying the effects of the different pathways of stabilization (Fig. 2).

91 This thesis highlights the necessity to jointly consider priming and entombing effects as the two faces of
92 the same 'microbial coin'. The balance between these two processes will affect the final C sequestration
93 efficiency, as stated by Liang et al. (2017). We observed, however, how a higher microbial activity have a
94 positive effect on both i) C entombing via necromass and exopolysaccharides deposition and ii) reducing
95 the priming in fertile soil with high microbial biomass, suggesting an overall positive effect of microbial
96 communities on C sequestration.

97

98 5.4. Root traits matter: N₂ fixing species (Fabaceae) vs conservative non N₂ fixing 99 species (Poaceae)

100 Plants are widely recognized as the main factor influencing C input in soil via litter fall, root mortality and
101 exudation (Six et al., 2002; Derrien et al., 2016; Sokol et al., 2019). This thesis focused partly on the
102 comparison between N₂-fixing species (*Fabaceae*) with non N₂-fixing species (*Poaceae*) commonly used for
103 revegetating embankments in the South of France. Coupling isotopes labelling with fractionation
104 techniques, we demonstrate the beneficial effect of N₂-fixing species and the specific root traits that
105 species possessed in this study (low C:N, high hemicellulose and low lignin content, high root elongation
106 rate, low absorptive root diameter, low SRL, and high biomass) on total C sequestration and its
107 accumulation in stable C pools. The once common view that stable C storage is driven by selective
108 preservation of recalcitrant compounds is once again challenged by these results. We found root biomass
109 to be a better predictor of new C gain in every soil pool, more than any other root trait. The reason might
110 be that morpho-physio-phenological (as SRL and C:N, and diameter) can be compensated by the effect of
111 biomass, as performance traits (Violle et al., 2007). In soil planted with Poaceae species, C storage in the

112 POM C pool was greater due tissue recalcitrance inhibiting microbiological activity, microbiological
113 biomass and overall mineralization. However, this effect was masked in N₂-fixing species with their higher
114 root biomass and related C input. It is necessary to underline that N₂-fixing species are associated with
115 *Rhizobium* bacteria. This association increases the N content in the root biomass, decreasing the C:N ratio
116 and their recalcitrance. In addition, the symbiosis with *Rhizobium* bacteria increases microbial activity and
117 the deposition of microbiological exopolysaccharides (Garcia et al., 2001). These effects overall increase
118 the new C input in the system, especially in the SILT and SILT+CLAY (Cotrufo et al., 2013). For this reason,
119 is important to consider that the root traits related to high labile C input in this thesis are characterized by
120 an intrinsic high microbiological activity due to symbiosis of N₂ fixing species with *Rhizobium*. When
121 studying Leguminous species and C sequestration, the effect of the symbiosis and the root traits connected
122 with high labile input are therefore synergic in increasing new labile C input, and difficult to disentangle. In
123 general, root biomass and chemical traits (C:N ratio, and lignin/cellulose/hemicellulose ratio) were a better
124 predictor for C storage compared to architectural traits. Recent studies have highlighted correlations
125 between root economic spectrum (RES) and C storage (De Deyn et al. 2008; Roumet et al. 2016; Prieto et
126 al. 2016; Poirier et al. 2018; Henneron et al 2019). The main characterization of RES - as coordinate
127 variation of root respiration rate, decomposability, and morphological and chemical traits related to C
128 economy (Roumet et al., 2016) - is given by the distinction between fast growing, acquisitive species (e.g.
129 Fabaceae) and slow growing, conservative species (e.g. Poaceae) (Chapter II, this thesis). Therefore, our
130 findings are consistent with the bulk of literature that find N₂-fixing Fabaceae in the spectrum of acquisitive
131 species characterized by high input of labile C in the soil, and N₂-fixing species Poaceae representing
132 conservative species with low input of recalcitrant old C (Prieto et al., 2016; Henneron et al., 2019). These
133 results support the correlations between RES and soil C sequestration potential, mediated by root growth
134 strategies and different C economies (Roumet et al. 2016; Poirier et al. 2018; Henneron et al 2019).

135 However, when studying RES correlations with C storage is important to differentiate between N₂ fixing
136 and non N₂ fixing species.

137

138 5.5. Soil matters: a major factor in carbon-cycle regulation, but due to indirect 139 effects

140 Soil fractions and the related C pools have shown to be fundamental and understand the C-cycle in soil.
141 The results of this thesis show that soil type has the highest impact on both C storage and priming effect.
142 Reduced root biomass in subsoil due to low fertility decreases the input of new C in every soil C pool and
143 the transfer of C in the SIL+CLAY pool via microbial metabolic transformations (Cotrufo et al., 2013; Vidal
144 et al., 2018). It also has a negative effect on priming, since labile C input are not high enough to allow
145 substrate preferential switch of microbial communities (Cheng and Kuzyakov, 2005; De Graaf et al. 2010).

146 However, when observing the direct effect of soil characteristics on the soil C storage, it is surprising to
147 observe that C saturation has no effect on the increase of protected C in the SILT+CLAY fraction. In our
148 experiment, subsoil had a higher clay percentage and lower initial C content compared to topsoil,
149 decreasing soil C saturation that should positively influence C storage in the SILT+CLAY C pool (Six et al.,
150 2002; Eyles et al., 2015; Shahbaz et al., 2017). Under these premises, we expected a higher rate and
151 amount, or at least relative amount, of C stored in fine SILT+CALY pool. However, topsoil had a higher
152 increase in the SILT+CLAY C pool compared to subsoil in absolute terms, while in relative terms they were
153 comparable. The reasons behind this behavior is attributable to plant biomass and microbial communities
154 in different soils. A lower fertility of subsoil decreases the input of C via plant biomass. The decreased
155 microbial activity and abundance decreased the input of processed C in the SILT+CLAY C pool. Without C
156 input in the SILT+CLAY C pool, increased potential for organomineral interactions did not influence the
157 amount of stored C. With this diagnosis, we could not claim that clay content and C saturation had no

158 effect on potential C storage. However, we can affirm that they were less important to C input by plants
159 and C metabolic transformation by microbial community in respect to soil C storage.

160 Soil N content had a high impact on soil C-cycle, increasing fertility and biomass production and,
161 consequently, new C input. Regarding the priming effect in soil, we confirm that in a poor soil the
162 competition for N reduced the consumption of C from microorganisms and had a positive effect on soil C
163 storage (Cheng and Kuzyakov, 2005). We confirmed that N rich soils positively influenced soil C storage
164 (Dou et al. 2016, Guo et al., 2019).

165 Aggregate stability had a significant effect on C storage due to a double feedback mechanism: new C input
166 in the system participated in creating more stable aggregates that, in turn, protected the encompassed C
167 (Tisdall and Oades 1982; Caesar-Tonthat 2002; Nichols and Wright 2005). The decrease of old C
168 mineralization in SILT+CLAY C pools were the direct result of the higher physical protection of C in stable
169 aggregates (Chevallier et al., 2004). Soil structure, and especially aggregates, seems to be the main direct
170 soil effect that influences C stability, and more studies need to be developed on the subject.

171

172 5.6. Ecological engineering toward a carbon sequestration goal

173 I suggest that we do not only consider soil C storage potential from a point of view of mineralogy or clay
174 content (Hassink 1992, Hassink al. 1997) or C saturation (Six et al. 2002), but we should pay more attention
175 to soil health. More specifically, we need to assess its fertility levels, such as N content, aggregate stability
176 and microbial community development (biomass and/or activity). Microbial diversity could also be an
177 important indicator. These indicators are connected with higher input of C in soil via increased biomass
178 production, transfer to the protected SILT+CLAY pool and negative priming due to the switch of substrate
179 utilization. An overview of the effect of soil and plants on C sequestration can be found in Figure 2.

180 The use of fertile topsoil increases carbon accumulation when compared with poor subsoil and it is
181 therefore desirable for revegetation of geotechnical soils. When revegetating fertile topsoil, fast growing
182 N₂ fixing species with high input of labile C are more efficient to store C in the protected SILT+CLAY pool
183 via higher root input and microbial turnover. Revegetating topsoil also induces a negative priming effect,
184 increasing preexistent C stability.

185 However, the use of topsoil is not always possible. Some particular conditions, eg in quarries and minor
186 road embankments, might require revegetation of subsoil. In this case, I advise to:

187 1) Fertilize soil to increase biomass production and C storage in soil. Fertilization is recognized to
188 increase soil C storage in both unprotected and protected C pools (Dou et al. 2016, Guo et al.,
189 2019). However, to my knowledge, no studies exist on the effect of fertilization on subsoil brought
190 to the surface. Moreover, the C impact of different fertilizer production and transport needs to be
191 compared with the benefits for C storage, or the final result might be detrimental for global C
192 storage.

193 2) Inoculate with microbial communities: we argue that inoculation of bacteria and mycorrhiza
194 (especially *Rhizobium* associated with Leguminous species to increase nodulation, and arbuscular
195 mycorrhiza fungi) would increase the C input in the SILT+CLAY C pool. Li et al. (2016) found a
196 decrease in C loss due to an increase of soil microbial biomass. However they did not consider the
197 different fluxes of C (New C and Old C), so it is not possible to assess if the increase of respiration
198 was detrimental for C balance. Kuimei et al. (2012) observed an increased soil C sequestration with
199 arbuscular mycorrhiza fungi inoculation in a reclaimed mine soil treated with coal gangue, fly ash
200 and sludge.

201 The lower C saturation did not increase protected C storage in subsoil in our experiment, but is still
202 promising for potential C storage if fertility and microbial requirements are met.

203 If neither fertilization nor microbial inoculation are possible, we suggest avoiding the use of N₂ fixing
204 species, since the increase in microbial biomass connected with those species will result in a higher
205 mineralization of old SOC.

206 Figure 3 shows a simple flowchart that provides suggestions for revegetating geotechnical soils and
207 optimize C sequestration. However, it is to be noted that this flowchart is based on results that present a
208 major shortcoming: the short timeframe of experiments. Long-term experiments are now much needed,
209 to explore how the results from this thesis are influenced over time.

210

211 5.7. What research remains to be performed?

212 Countless opportunities for research are possible in the C storage domain. However, this work on C storage
213 in revegetated geotechnical soil sparked some specific questions that I feel should to be tackled to have a
214 more comprehensive view of the system, from both a mechanistic and an applied point of view.

- 215 • It would be vital to extend these studies on fractions and C sources on long term experiments.
216 How subsoil evolves and 'become topsoil' is a fundamental aspect to be considered in studies of
217 plant development, microbiological characteristics and soil aggregation etc. However, the cost and
218 experimental setup makes it difficult to implement long-term constant isotope labeling
219 experiment. A solution would be to use C₃ plants grown on soil planted only with C₄ plants (or
220 vice-versa) as they have different isotopic signatures (Hobbie and Werner, 2004; Kuzyakov 2006).
- 221 • While in topsoil, research on aggregate protection and formation are numerous, they have not
222 reached a complete consensus on the processes involved. In subsoil, instead, the role of
223 aggregates in C protection and their formation processes remains still obscure. In Annex I, I present

224 a preliminary work to investigate in depth the influence of soil structure on C protection in topsoil
225 and exposed subsoil.

226 • In this thesis, I investigated the influence of microbiological communities on C-cycling and
227 sequestration based on microbiological activity and biomass. However, refined identification of
228 microbiological community structure and diversity would help unveiling key processes and factors
229 in C-cycling. Studying the evolution of microbiological communities on subsoil brought to the
230 surface would be fundamental to better understand its C sequestration potential, the processes
231 behind it, and to have an insight into soil evolution.

232 • More studies have to be carried out on inoculation with different strains of fungi and bacteria to
233 understand the mechanisms and influence of inoculation on C-cycling and its role as a 'pump' for
234 complexed C in the protected silt and silt+clay C pools.

235 • Microbial communities influence most of the major processes involved in soil C storage, directly
236 or indirectly. More precisely: i) total C input, as microbes area commonly used indicator for soil
237 fertility and health (Waksman, 1922; Waksman and Starkey, 1924; Mader et al., 2002; Suzuki et
238 al., 2005; Schloter et al., 2018), ii) quality of the stored C, determining if the C will accumulate in
239 the unprotected C pool (when microbial abundance and activity is low) or if it will be metabolized
240 and transferred in the protected C pool (when microbial abundance and activity is high) (De Deyn
241 et al, 2008; Cotrufo et al., 2013; Liang et al. 2017), iii) total respired C (increased with increasing
242 microbial activity) and priming effect (decreased when microbial activity is inhibited due to
243 competition or a switch of substrate preference from old C to new C) (Cheng and Kuzyakov., 2005;
244 De Graaf et al., 2010), and iv) increased aggregate formation and stability (Tisdall and Oades 1982;
245 Caesar-Tonthat 2002; Nichols and Wright 2005). Relying on clay abundance or C saturation levels
246 to determine the quantity of protected C might be inaccurate, since they are not an indicator for
247 soil health, and since microbial communities will determine the C input in the silt+clay C pool. For

248 these reasons we state that, given the correlations between microbial activity and many of the
249 key processes involved in C storage, further research should be carried out regarding the use of
250 microbial characteristics as an indicator for potential soil C storage.

251

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FIGURES AND TABLES

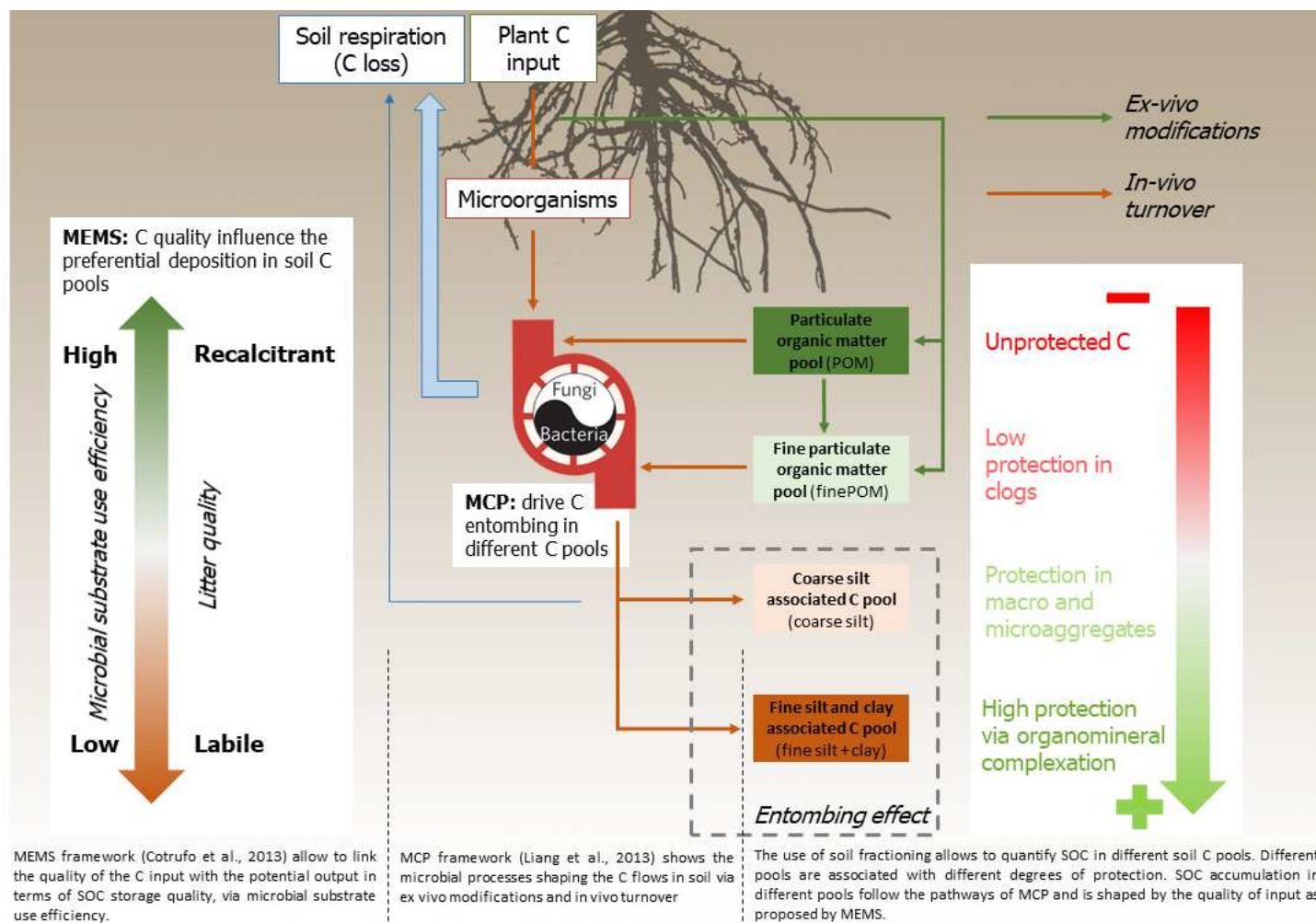


Figure 2: Conceptual framework illustrating complementarity between MEMS and MCP models, enriched by including the soil C pools. Primary plant's carbon inputs quality influence the final stabilization of stored C (MEMS, Cotrufo et al. 2013). The microbial carbon pump determine the entombing of C in the soil system, via ex-vivo modifications (green arrows) and in-vivo transformation (brown arrows) (MCP, Liang et al. 2017). Finally, the quality of C input will determine the C distribution in different soil C pools associated to soil fractions, through the microbial carbon pump. Labile C input will favor in-vivo turnover, increasing C in the silt and silt+clay fractions, protected in microaggregates and via organomineral interactions with fine silt and clay minerals. Recalcitrant C decrease in vivo turnover, and C accumulate mainly in unprotected POM and finePOM fractions via ex-vivo modifications.

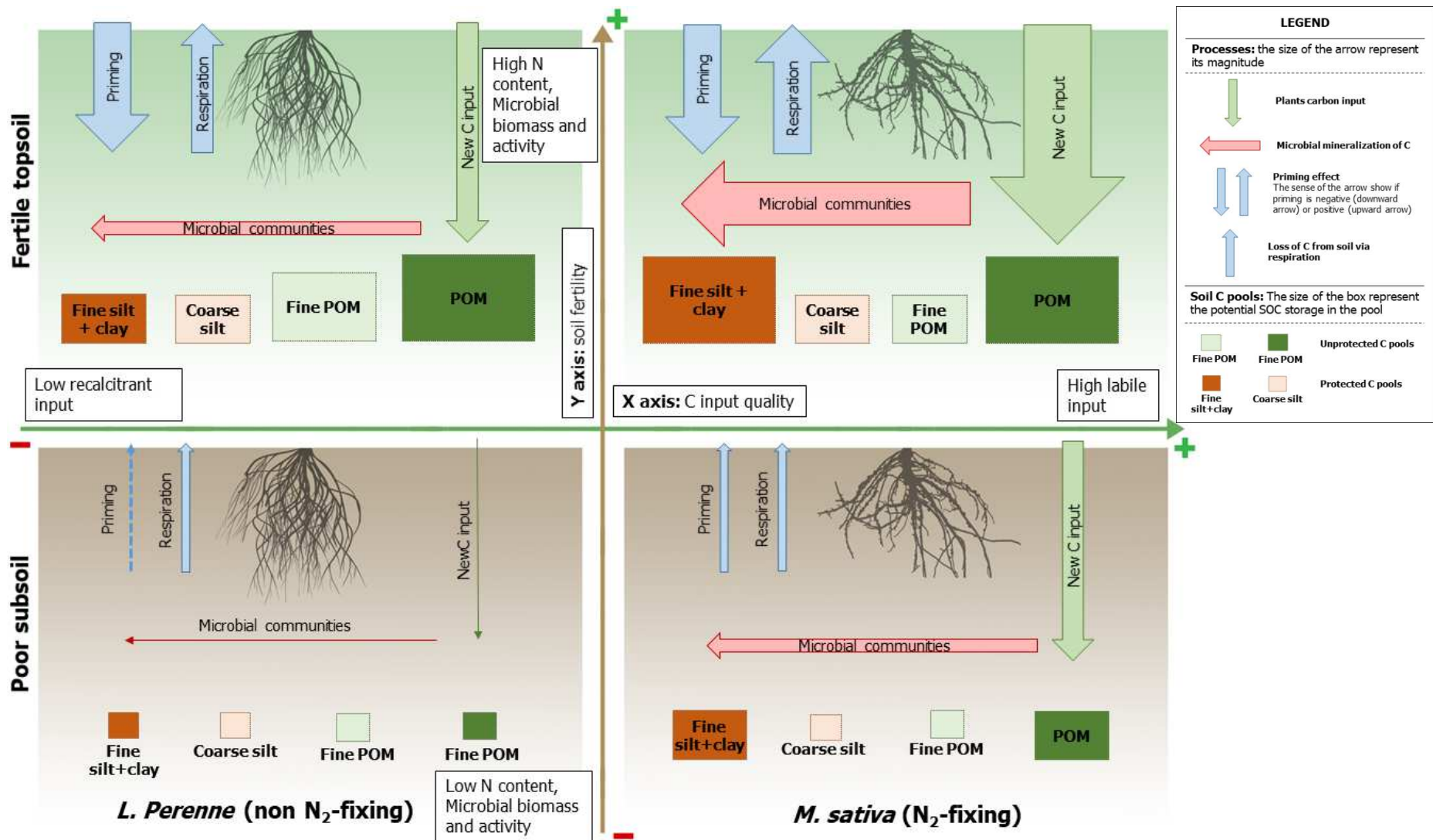


Figure 3: Scheme illustrating the changes in the C storage mechanisms along two main axes: x – Soil fertility and y – C input quality. Top right has the higher C storage potential, with high fertility soil revegetated with N₂-fixing species, that have high labile C input, positively influencing POM and, most importantly, fine silt+clay fraction accumulation through high microbial activity. Top left corner shows the potential soil C storage of fertile soil revegetated with non N₂-fixing species. The lower input of recalcitrant C decrease microbial biomass and activity, and increase POM and finePOM accumulation via decreased mineralization. In fertile topsoil priming effect is high and negative. Bottom right corner shows the effect of revegetating poor subsoil with N₂-fixing species: decreased C input due to fertility decreases the accumulation in the POM fraction, and decreased microbial biomass/activity its complexation in the protected silt+clay fraction. Priming effect is lower in intensity but positive; increasing the loss of preexistent SOC. Bottom left corner shows the effects of revegetation of poor subsoil with non N₂-fixing species. Decreased input and microbial activity decrease the C accumulation in every soil fraction, however priming effect almost absent due to plant-microbes competition for nitrogen.

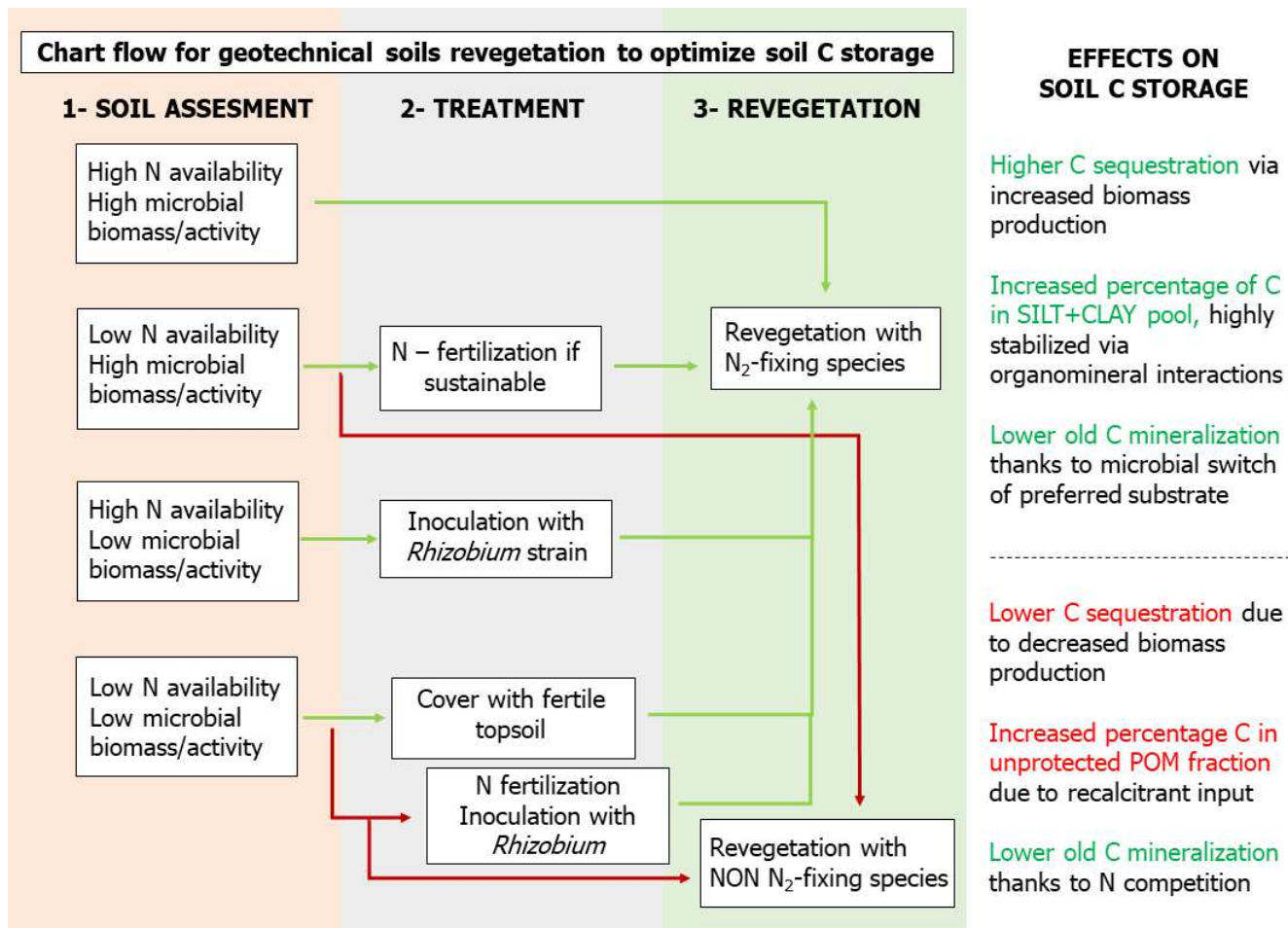


Figure 4: flowchart for soil revegetation to increase C sequestration in geotechnical embankments. First step is soil assessment in regard to N content and microbial biomass and activity. The second step shows the treatments to implement if N content is low (fertilization) or microbial communities are underdeveloped (inoculation). The fourth step shows the suitable species for revegetation given the soil conditions. The last on the effects on C sequestration given the soil, the treatments and the plant species selected. Green arrow shows the suggested pathway to take, red arrow the alternative unadvised pathway if the first is not possible. This flowchart, however, is based on a short term experiment. Long term experiment should be implemented to improve it.

Annex I: Perspectives: the influence of vegetation on soil microstructure and its implications on soil carbon sequestration: a geotechnical approach

1 INTRODUCTION

2 A particular interest in sustainability has been voiced in both environmental and geotechnical disciplines,
3 given the global climate change challenge that requires immediate action in multiple sectors. However,
4 the research on sustainability remains largely confined in the peculiarity of each discipline, which has its
5 specific assumptions and methodology characterizing the sustainability. We argue that a higher
6 multidisciplinary approach is needed to hybridize research and find transdisciplinary methodologies and
7 points of view on sustainability in every discipline. In this part of the research, we aim to bridge C storage
8 research from a plant/soil science point with geotechnical engineering research. Soil microstructure have
9 proven to be a good common ground between the two fields, since it is largely studied both with regards
10 to C sequestration and in the geotechnical engineering field for characterization of the soil structural
11 properties.

12 Soil structure has been proven to be central with regards to C sequestration, especially regarding the role
13 of aggregates. Aggregates forms through binding of soil particles by fine roots and fungal hyphae (Tisdall
14 and Oades 1982). Glycoproteins, polysaccharides, and mucilage, from plants cement their structure and
15 influence their stability (Tisdall and Oades 1982; Caesar-Tonthat 2002; Nichols and Wright 2005).
16 Aggregates occlude C in their structure, physically impairing the accessibility of microbes (O'Brien &
17 Jastrow, 2013). The efficiency of aggregates C protection will depend on their stability and the amount of
18 C stored inside their structure. Aggregates is a dynamics process, and the higher is their stability the higher
19 they will resist to disaggregation (Eyles et al., 2015). In soil sciences most of studies refers to three main
20 classes: microaggreagets (0.02-0.2mm), macroaggregates (0.2-3mm) and clots (3-5mm). Aggregation in
21 soil will deeply influence the void ratio, a common indicator used in geotechnical research to define soil
22 structural characteristics. Aggregates structure gives a double porosity behavior to soil, with micropores
23 characteristic of intra-aggregates structure, and macropores formed by the interaction between different

24 aggregates (Koliji et al., 2008). In geotechnical studies, a common method used to investigate soil porosity
25 and deriving void ratio is mercury intrusion porosimetry (MIP). MIP allow characterizing the cumulative
26 and relative abundance of voids of different pore classes (Russo et al., 2016). The aggregation process and
27 characteristics are expected to be correlated with void ratio in different pore classes. Soil porosity and
28 connectivity also influence the possibility of microbes to be in contact with substrates and their
29 consumption (McCarthy et al., 2008; Lugato et al., 2009). Vegetation can influence soil porosity in different
30 classes due to i) root channels, ii) hyphae development, iii) wet and dry cycles, iv) cementation and clogging
31 of micropores due to rhizodeposition and v) aggregation and disaggregation processes favored by plants
32 influence (McCarthy et al., 2008; Lugato et al., 2009). Extensive studies have been implemented on
33 aggregate formation and C protection. However, most of the studies see aggregates as ‘building blocks of
34 soil’ (Malamoud et al., 2009) and overlook the more complicated structure deriving by their interactions.
35 Moreover, studies on aggregate formation and C protection have seldom been implemented on subsoil,
36 where the aggregate formation processes are still debated. Soil microstructure has great potential to
37 shape C sequestration in soil, and we aim to assess evolution of aggregate characteristics due to
38 revegetation in topsoil and subsoil brought to the surface. Moreover, we aim to assess pore ratio in soil
39 and the influence of vegetation using MIP, and correlate it to aggregate characteristics, to better
40 understand how aggregates shape soil structure. The use of MIP together with aggregate stability and C
41 analysis will allow comparing methods characteristics of different disciplines and exploring possible
42 exchanges and overlapping between these fields.

43

44 OBJECTIVES AND HYPOTHESIS

45 Our first objective is to investigate the influence of vegetation on microstructure using the MIP
46 methodology. For this, we assessed bulk void ratio at time 0 and after 6 months of two soils showing

47 contrasting characteristics (topsoil 0-30 cm depth and subsoil 110-140 cm depth) vegetated with *M. sativa*
48 and *L. perenne*, plus bare soil controls. Comparing void ratio in bare soil control after 6 months of
49 experiment (time 6) with the initial soil (time 0) will allow us to assess the effect of wet and dry cycles on
50 microstructure (since soil was kept at 45% of water holding capacity with irrigation to compensate
51 evaporation) and compare it with the effect of revegetation. Void ratio will be cumulated in different
52 classes relative to different aggregates and processes as a proxy for: microaggregates porosity,
53 macroaggregates porosity deriving from microaggregates interaction and clots porosity deriving from
54 macroaggregates interactions.

55 Our second objective is to characterize aggregates and their characteristics for bare soil control and
56 vegetated treatment with *M. sativa* and *L. perenne*. For this, we measured aggregate stability, quantity of
57 C protected inside of aggregates, quantity of fresh new C inputted in different aggregates classes in 6
58 months, thanks to the constant CO₂ enrichment with ¹³C.

59 Our final objective is to investigate the relationship between aggregate stability and void ratio in different
60 pore classes, ii) between C protected in different aggregate classes and void ratio in different pore classes
61 and iii) if new C input in different aggregate size influence macro- and microporosity.

62 We hypothesize that plants will increase macroaggregates and clots porosity due to channeling of roots
63 and aggregates formation. However, vegetation might clots micropores due to rhizodeposition (McCarthy
64 et al., 2008). In this case, aggregate stability will increase with increased void in macroaggregates and clots
65 (due to aggregate formation and inter-aggregate porosity) while it will be negatively correlated with
66 porosity in microaggregates, due to bioclogging from microbial activity that cement and increase
67 aggregate stability (Ivanov and Chu, 2008). However, wet and dry cycle will probably drive the formation
68 of soil structure.

69 Another hypothesis is that the protection of C will increase when decreasing the porosity (and void ratio)
70 in macro and microaggregates, since the microbes will not have access to the occluded C. Finally, we expect
71 that new C input is positively correlated with porosity in macroaggregates and clots (due to the role that
72 fresh C input plays in aggregate formation, and root channeling connected with new C deposition) but
73 negatively correlated with porosity in microaggregates, due to the clogging from rhizodeposition and
74 microbiological exudation and exopolysaccharides. The analysis will be conducted using Pearson's
75 correlations between C protection and new C input in microaggregates, macroaggregates and clots, and
76 the void ratio (as proxy for porosity) in three different pore classes.

77

78 STATE OF THE WORK

79 Analyses on aggregate stability, C protection in aggregates and new C input in different aggregates classes
80 have been performed. A first MIP analysis campaign has been carried out to design the work. A second
81 campaign to acquire MIP replicates is in progress and expected to finish by the end of October 2019. After,
82 correlations with aggregate properties will be investigated to study the relations between vegetation, soil
83 structure in terms of aggregation and porosity, and C protection. Methodology and preliminary results are
84 presented in the following sections.

85

86 **METHODOLOGY**

87

88 *Mercury intrusion porosimetry (MIP) curves and cumulative porosity for different pore classes*

89 MIP test allow to characterize the porosity of the sample in an entrance pore diameter that ranges
90 between 0.001 to 300 μm . Abundance of pores of different diameters define the microstructure of the soil.
91 Once grouped in different pore size classes, we believe to find correlations between pores and aggregate
92 characteristics in soil.

93 **Methodology:**

- 94 1. 1-2 g of sample are dehydrated according to the freeze-drying method (Delage et al. 1984).
 - 95 2. MIP test is performed in a double chamber Micromeritics Autopore III apparatus.
 - 96 3. Place the sample in the filling low-pressure apparatus (dilatometer).
 - 97 4. The samples is outgassed and under vacuum, and after filled by mercury. The chamber is at
98 ambient absolute pressure.
 - 99 5. Pressure is then rise up to 200 kPa using of compressed air
 - 100 6. Chamber is depressurized and the samples were transferred to the high-pressure unit,
 - 101 7. The pressure is then raised to 205 MPa following a previously set intrusion program. At any
102 intrusion step a time sufficient to observe a quasi-static penetration of mercury was allowed.
 - 103 8. A blank test is performed to corrections the results and prevent errors deriving from the
104 compressibility of the intrusion system.
 - 105 9. Finally, SEM analyses were performed on dehydrated samples in order to highlight their fabric.
- 106 Output: Intruded void ratio and pore size density function for different pore classes rangion from 0.001 to
107 300 μm

108 ***Aggregate stability***

109 Aggregates provide physical protection to carbon. However, the degree of protection of carbon depends
110 on their stability. The more stable are the aggregates, the more resilient will be to changes in
111 environmental condition, decreasing their disaggregation and ensuring stable physical protection to the
112 carbon occluded. Mean weight diameter is a standard indicator for aggregate stability, as the mean
113 diameter of aggregates that, starting from a 5-3mm diameter, have undergone a disaggregation process
114 via wet and drying. The higher the MWD, the lower the disaggregation.

115 **Methodology according to le Bissonais et al. (2006):**

- 116 1. 20 g of soil collected and air dried
- 117 2. The sample is sieved first at 5mm and after at 3mm, to isolate the 3-5mm fraction
- 118 3. Aggregates are put in the oven for 24h so they are at the same matrix potential
- 119 4. 5g of 3-5mm fraction are weighted and gently immerse in a 250 cm³ beaker filled with 50 cm³ of
120 ethanol for 10 minutes
- 121 5. Ethanol is sucked off with a pipette
- 122 6. Sample transferred in a 250cm³ Erlenmeyer flask containing 50cm³ of deionized water and
123 brought to 200cm³
- 124 7. Flask is agitated 20 times and left 30 minutes for sedimentation of coarse particles
- 125 8. Water sucked off with a pipette
- 126 9. Mixture of soil and water transferred to a 50µm sieve previously immersed in ethanol
- 127 10. Sieve gently moved 5 times to separate <50 µm from those >50 µm, use of ethanol for the wet
128 sieving to reduce additional breakdown
- 129 11. >50 µm fraction is collected from the 50-µmsieve, oven-dried and gently dry-sieved by hand on a
130 column of six sieves: 2000, 1000, 500, 200, 100 and 50 µm

131 12. Mass percentage of the different fraction is calculated, and for subtraction even the <50µm
132 fraction

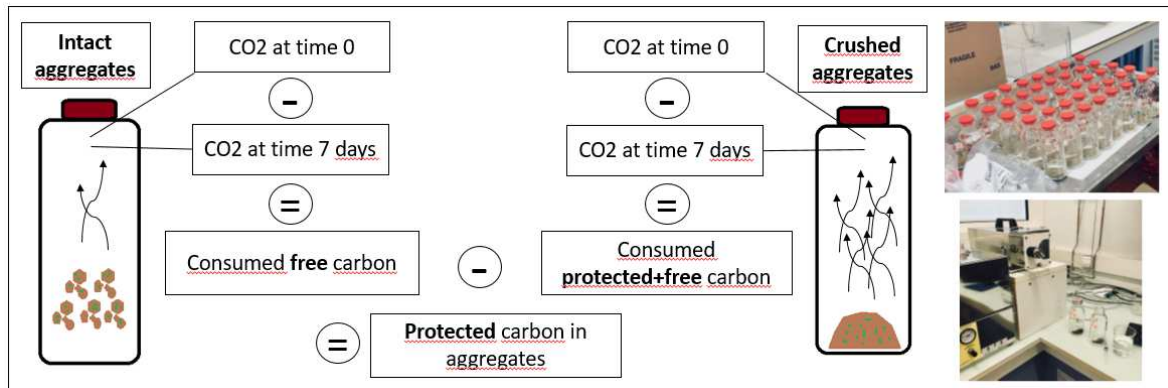
133 13. MWD is calculated as the sum of the mass fraction of soil remaining on each sieve after sieving
134 multiplied by the mean aperture of the adjacent mesh

135 Output: MWD for different soils and treatments

136

137 ***Protected carbon: aggregate mineralization***

138 Aggregates provide physical protection to carbon. However, not all the carbon is automatically protected
139 inside the aggregates. The protection will depend on the porosity of the aggregates and the amount of
140 microbial biomass enclosed in the aggregates itself. This will ultimately influence the capacity of microbes
141 to get in contact with the C substrate and mineralize it. We aim to assess the degree of C protection in
142 different aggregates sizes of the two different soils (topsoil vs subsoil) and three different treatments (*M.*
143 *sativa*, *L. perenne*, bare soil). The soil nature will influence pore size, microorganisms' abundance and
144 aggregate stability. First, to assess the unprotected C in aggregates we measure the amount of CO₂
145 released when incubating undisturbed aggregates of different classes. After, to assess the amount of total
146 C (unprotected and protected) we finely grind aggregates the aggregates (to remove their physical
147 protection on carbon) and assess the CO₂ respired during incubation. The difference between these two
148 values (CO₂ deriving from protected C and CO₂ deriving from total C) will allow us to assess the amount of
149 soil derived CO₂ that is protected in different aggregates classes (3-5 mm, 0.2-3 mm, 0.02-0.2 mm) for
150 different soils (topsoil and subsoil) and species (*M.sativa*, *L.perenne* and bare soil) (Figure A1).



151

152 **Figure A1:** : scheme of the microcatalometer methodology

153 **Methodology:**

- 154 1. Manually crush the soil and push it through a 5000 μm sieve (aggregates 3000-5000 μm fraction)
- 155 2. Sieve at 3mm and 0.2mm (aggregates < 200 μm fraction and 3 mm to 200 μm).
- 156 3. Collect the different fractions and separate them in half
- 157 4. Crush half one half of each fraction to obtain two subsamples: uncrushed aggregates and
- 158 crushed aggregates (20g of aggregates for each sample)
- 159 5. Bring them to 75% of water holding capacity
- 160 6. Samples placed in 125 ml jars with parafilm allowing the interchange of gases (but not water)
- 161 and incubated at 28 $^{\circ}\text{C}$ for 7 days.
- 162 7. Each sample was adjusted for soil moisture and, just after, the bottles were air tightly closed
- 163 and measurements of respiration made. After 6 hours of incubation (without any gas
- 164 interchange) measurements were made.
- 165 8. The differences of CO₂ between these two measurements gave the amount of respired CO₂ in
- 166 6h per treatment, soil and aggregate class.

167 Output: amount of respired CO₂ for crushed and uncrushed aggregates. The difference between these
168 measurements represent the aggregates protected carbon in potential respiration. These results were
169 available for 3-5mm, 0.2-3mm and 0.02-0.2mm aggregate classes.

170

171 *Plant derived fresh carbon (new C) stored inside aggregate structures*

172 It is well known how aggregates provides protection for C, however the aggregate formation processes
173 are still debated. Especially in subsoil were little is known about aggregate structures and formation. C
174 deriving from plants, often processed by microbes, is recognized as one of the main actors in aggregate
175 formation. The input of C as plants' exudates and microbiological exudates and exopolysaccharides
176 cement the mineral structure of the aggregate that will provide protection from microbial mineralization.
177 We aim to investigate the pathways of C input in different aggregate classes to acquire information on
178 aggregate formation and C protection in the two different soils.

179 **Methodology:**

- 180 1. A subsample from the aggregate abundance samples was taken, representative of the following
181 aggregate classes:
 - 182 1) aggregates 2000-5000 µm (2-5 mm)
 - 183 2) aggregates 200-3000 µm (0.2-3 mm)
 - 184 3) micro-aggregates 20-200 µm (<0.2mm)
- 185 2. The subsample is finely ground with an agate mortar and stored
- 186 3. The sample were analyzed to asses SOC and ¹³C with an elemental analyzer Isoprime100 coupled
187 with an Elementar Varo Isotope Cube at INRA Nancy.

188 4. To calculate the proportion of NewC in aggregates, an isotope mixing model according to the work
189 of Balesdent and Mariotti (1996) was used:

$$190 \quad \%C_{new} = \frac{\delta(t1) - \delta(t0)}{\delta B - \delta(t0)} \quad [1]$$

191 Where %C_{new} is the percentage of new carbon in the measured SOC of a specific aggregate
192 fraction, $\delta(t1)$ is the $\delta^{13}C$ signature of the SOC measured in a specific aggregate fraction at the
193 end of the experiment (t1), $\delta(t0)$ is the $\delta^{13}C$ signature of the SOC before the experiment (t0), δB is
194 the $\delta^{13}C$ signature of the new C input in the system, in our case the signature of the root biomass
195 (as the average of adsorptive and transport roots signature).

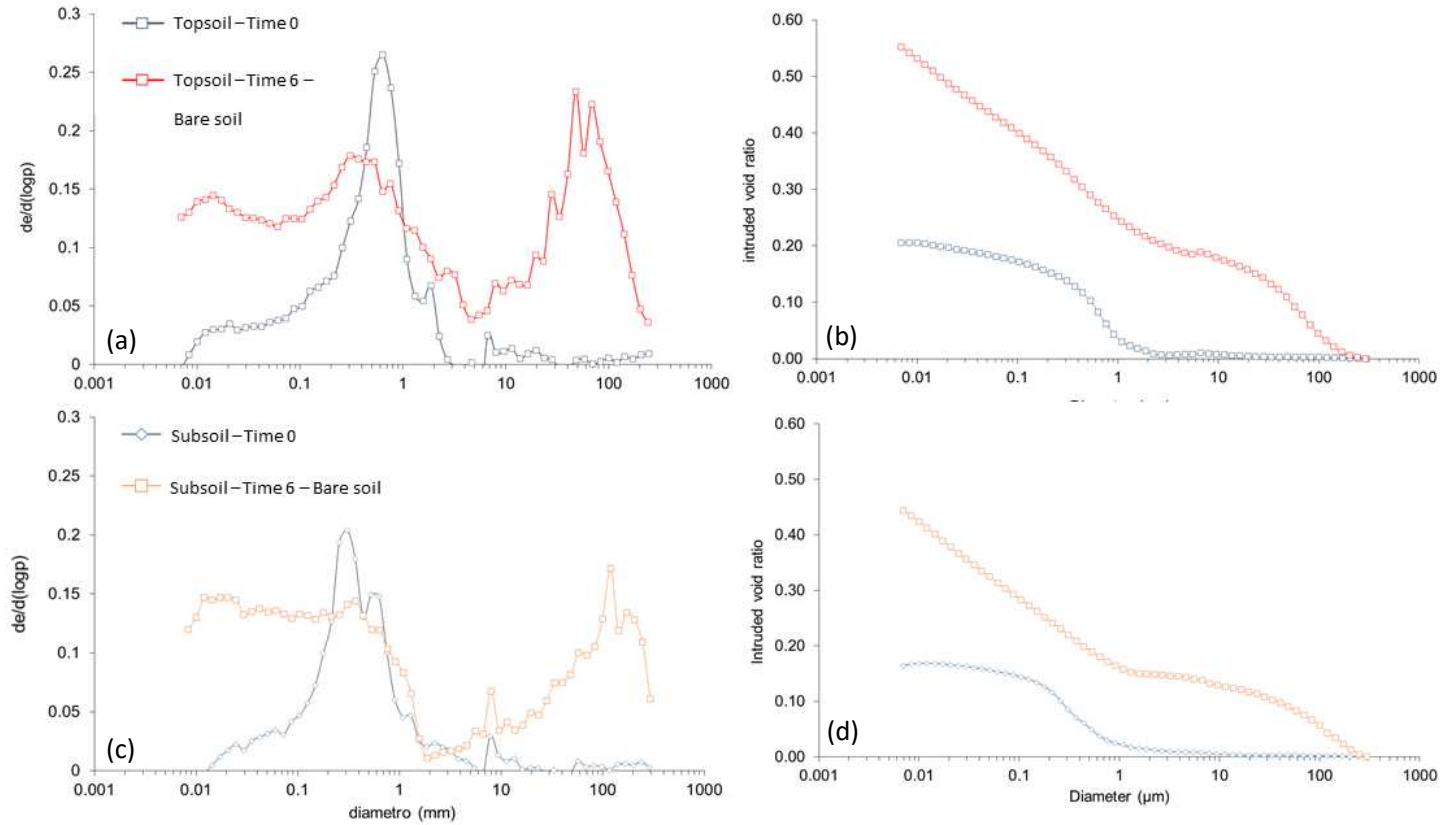
196 5. Multiply the total SOC for the %C_{new} provides the amount of NewC in mgC g⁻¹ soil.

197 Output: Amount of total SOC and NewC in the different classes (3-5 mm, 0.2-3 mm, 0.02-0.2 mm) for
198 different soils (topsoil and subsoil) and species (*M.sativa*, *L.perenne* and bare soil).

199 PRELIMINARY RESULTS AND MAIN DISCUSSION POINTS

200 *Mercury intrusion porosimetry (MIP) curves and cumulative porosity for different pore classes*

201 **Bare soil condition: Effect of wet and dry cycles**

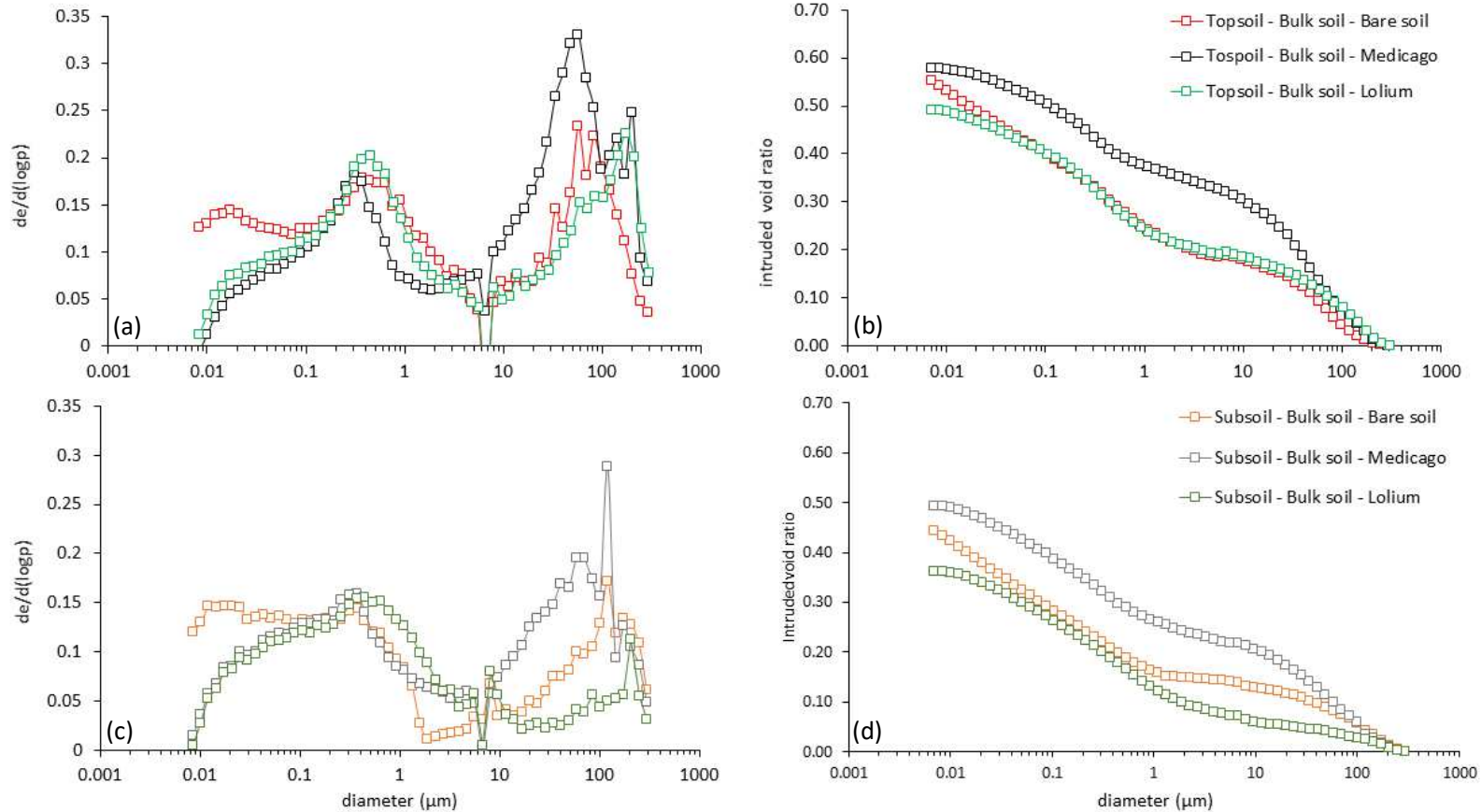


202

203 **Figure A2:** Evolution of microstructural voids in 6 months of wet and dry cycles, with no vegetation sowed. Figure (a) shows the pore size density function of topsoil
204 at time 0 (blue boxes) and time 6 (red boxes) without vegetation. (b) shows the total intruded void ratio for different diameter classes in topsoil. (c) shows the pore size
205 density function of subsoil at time 0 (blue boxes) and time 6 (orange boxes) without vegetation (d) the total intruded void ratio for different diameter classes in
206 subsoil.

- 207 • Wet and dry cycles have a high effect on soil structures: soil structure pass from a mono-modal curve structure to a bi-modal curve
208 structure in both soils.
- 209 • Void ratio highly increase in the micropores (<0.1 μm) and macropores (>10 μm) classes due to wet and dry cycles. Mesopores (0.1-10 μm)
210 decrease during the 6 months experiment.
- 211 • The wet and dry cycles proved to influence both aggregates formation and stabilization processes (Shiel et al. 1988; Deneff et al. 2001). We
212 hypothesize that, after the soil preparation (that included crushing and sieving) the soil lost its primary microstructure. Wet and dry cycles
213 increase aggregate formation and, consequently, the microporosity deriving from intra-aggregates structure and the macroporosity
214 deriving from inter-aggregates interactions.
- 215

216 **Vegetated treatment compared to bare soil after 6 months of growth**



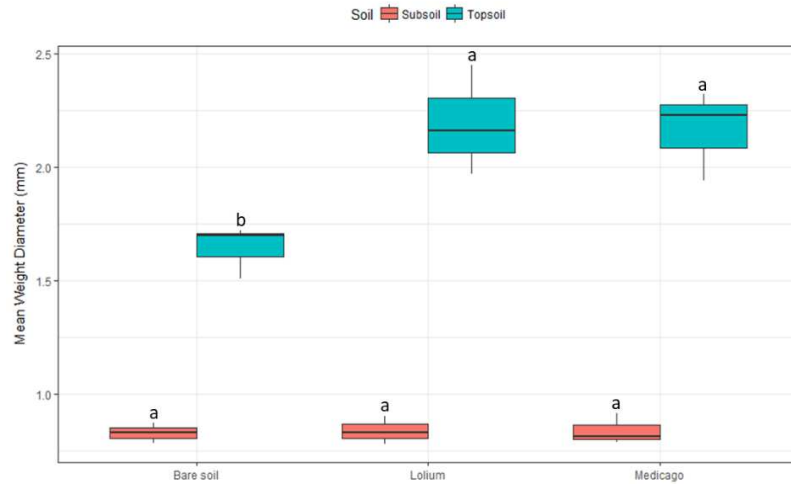
217

218 **Figure A3:** Evolution of microstructural voids in 6 months of soil vegetated with *M. sativa* and *L. perenne* compared to bare soil control. Figure (a) shows the pore size
 219 density function of topsoil in bare soil control (red boxes), *M. sativa* (black boxes), and *L. perenne* (green box) after 6 months of growth. (b) shows the total intruded
 220 void ratio for different diameter classes in topsoil. (c) shows the pore size density function of subsoil in bare soil control (orange boxes), *M. sativa* (grey boxes), and
 221 *L. perenne* (dark green boxes) after 6 months of growth. (d) shows the total intruded void ratio for different diameter classes in subsoil.

- 222
- *M. sativa* show an increase in total porosity in both subsoil and topsoil, while *L.perenne* decrease the total porosity (Figure A3b,d).
- 223
- *M.sativa* increase the macroporosity in both soils (> 10 μm), while *L.perenne* show a decrease in macroporosity in bare soil (Figure A3a,c).
- 224
- Both *M. sativa* and *L. perenne* decrease the microporosity of the system (<0.1 μm) (Figure A3a,c).
- 225
- The increase in macroporosity due to *M.sativa* might be correlated with increased aggregate formation and interactions, and root
- 226
- channeling effect. The decrease of microporosity is imputable to exudates from microbial activity and plants that clog the micropores in
- 227
- aggregates. However, replicates are needed to verify these hypothesis.

228

229 *Aggregate stability: Mean weight diameter (MWD)*



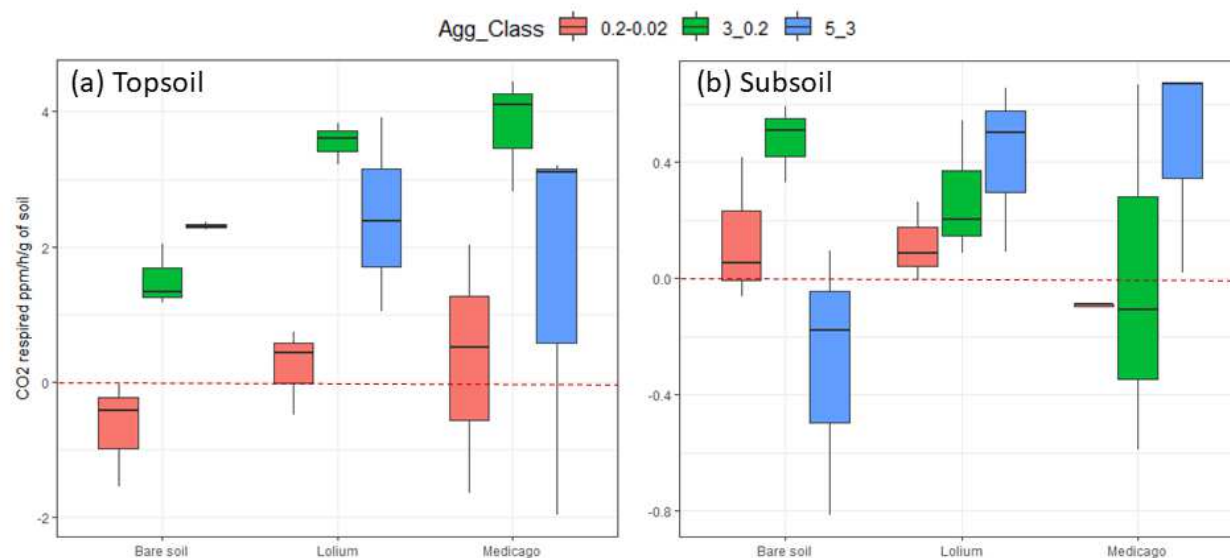
230

231 **Figure A4:** Mean weight diameter (MWD) in topsoil (blue boxplot) and subsoil (red boxplot) for the three different treatments (*M.sativa*, *L.perenne* and bare soil
232 control) after 6 months of revegetation. In each boxplot, the lower edge of the box corresponds to the 25th percentile data point, while the top edge of the box
233 corresponds to the 75th percentile data point. The line within the box represents the median and black dots indicate outliers. Different letters above the boxplots
234 indicate statistically significant differences ($p < 0.05$) between families and controls according to Tukey HSD test.

- 235
- Subsoil and topsoil have a significant different MWD after 6 months of revegetation, with a higher aggregate stability in topsoil.
- 236
- No significant effect of vegetation in subsoil.
- 237
- In topsoil vegetated treatment have a significantly higher stability compared to bare soil control. However, we didn't observe any effect of
- 238 different vegetation.

239

240 *Protected C in aggregates*



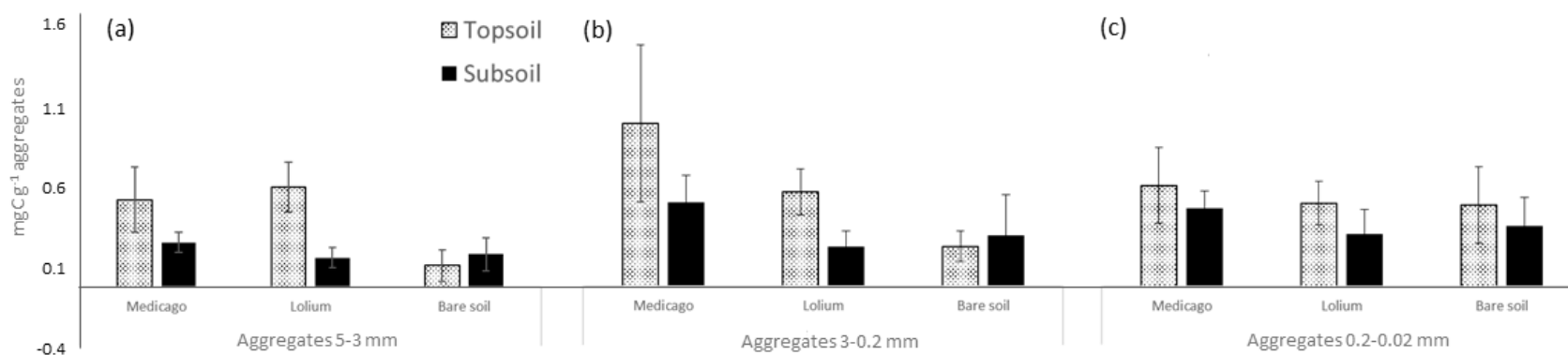
241

242 **Figure A5:** amount of CO₂ (respired ppm h⁻¹ g⁻¹ of soil) protected inside different aggregate classes (5-3 mm clogs in blue, 3-0.2 mm macroaggregates in green, 0.2-0.02
 243 mm microaggregates in red) in (a) topsoil and (b) subsoil. The protected C is calculated as the difference between the respired CO₂ deriving from incubation of
 244 undisturbed aggregates (CO₂ deriving from unprotected C) and CO₂ deriving from incubation of crushed aggregates (CO₂ deriving from consumption of protected
 245 and unprotected C). In each boxplot, the lower edge of the box corresponds to the 25th percentile data point, while the top edge of the box corresponds to the 75th
 246 percentile data point. The line within the box represents the median and black dots indicate outliers. The red dotted line is the 0 line, meaning no protection of C in
 247 aggregates.

- 248
- In topsoil the higher C protection is found in the macroaggregates, with vegetation that increase the amount of protected C underlining
- 249 the reactivity of this aggregate class to revegetation. In topsoil clots no effect of vegetation can be found, with increased standard variation
- 250 in vegetated treatment but no differences with bare soil.

- 251 • In subsoil we observe a very different trend, with vegetation decreasing the C protection in the 3-0.2mm macroaggregates, while increasing
- 252 the amount of protected C in the 5-3 clots, that did not presented any protection in bare soil control.
- 253 • In both topsoil and subsoil, microaggregates do not have any role in C protection, since no changes are observed when crushing them.
- 254 However, since the crushing was done by hand in an agate mortar it is not sure that the structures were efficiently disaggregated, leaving
- 255 a possible bias in the methodology.

256 *Plant derived NewC input in aggregates*



257

258 **Figure A6:** newC (mg new C g⁻¹ aggregates) deriving from plant input in aggregates for gram of aggregates of different classes for the three analyzed treatment

259 (*M.sativa*, *L.perenne*, and bare soil control) after 6 months in topsoil (light dotted bars) and subsoil (solid black bars). (a) Show the C concentration in the clots (5-

260 3mm diameter), (b) in the macroaggregates (3-0.2 mm), and (c) in the microaggregates (0.2-0.02mm). Bars represent the standard deviation.

- 261 • In clots (Figure 5A,a), vegetated treatment store more new C compared to bare soil, while in subsoil no significant differences seems to
- 262 occur.
- 263 • The higher increase in C seem to be in macroaggregates (Figure 5A,b), higher in *M.sativa* compared to *L.perenne* and bare soil.

- 264
- In microaggregates, no effect of vegetation seem to influence the quantity of new C moved (Figure 5A,c).
- 265
- Bare soil control shows increase in NewC, probably due to mosses colonization that mineralized enriched CO₂. The analysis of deeper layer
- 266
- of soil is needed to avoid the contamination.

267 **FUTURE WORK**

- 268 1. Replication of MIP results to confirm the trends.
- 269 2. MIP results will be separated in different pore classes to have cumulative data for void ratio in
270 different pore classes relative to different aggregate sizes.
- 271 3. Implement principal component analysis and Pearson's correlations between i) aggregate stability
272 MWD, ii) amount of C protected in different aggregates classes in CO₂ equivalent, iii) new C input
273 in different aggregate classes, and iv) void ratio in different pore classes representing
274 microaggregates, macroaggregates and clots, in the two soils and for the different species.
- 275 4. Discussion on the effect of vegetation on soil structure and relationship between aggregate
276 characteristics and soil structure in terms of void ratio. Use of the data to link the C input in
277 aggregates with the soil structure formation, and the feedback between the structure and the C
278 protection.
- 279 5. Results will help to understand the differences in C fluxes in aggregates in topsoil and subsoil
280 brought to the surface and test the aggregate hierarchy theory of Tisdall and Oades (1982), widely
281 accepted for topsoil, on subsoil brought to the surface. Preliminary results suggest that subsoil
282 brought to the surface might not share the same aggregation processes than topsoil.

283

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320

321

Résumé exhaustif: Objectifs, résultats, conclusions générales

267 OBJECTIFS GENERAUX ET HYPOTHESES

268 Les objectifs généraux appliqués de la thèse sont :

269 i. Comprendre l'effet des caractéristiques des plantes et du sol sur la séquestration du carbone dans
270 le sol en termes de quantité et de qualité (objectif fondamental)

271 ii. Identifier les pratiques possibles en matière de plantes et de sols qui peuvent être mises en œuvre
272 pour augmenter le stockage du carbone dans les remblais des routes et ferroviaires et,
273 éventuellement, dans les sols gris des travaux géotechniques (objectif appliqué)

274 Différentes questions spécifiques concernant les mécanismes fondamentaux du cycle C ont été abordés
275 dans chaque chapitre de la thèse.

276

277 *Chapitre II : Voie de la persistance : les caractéristiques des racines des plantes modifient*
278 *l'accumulation de C dans différents réservoirs de carbone du sol par médiation microbienne*

279 i. Objectif 1 : Comprendre les relations entre les caractéristiques des racines et l'accumulation de C dans
280 différents bassins de carbone du sol pour 12 espèces herbacées différentes couramment utilisées
281 dans la revégétalisation des talus.

282 Hypothèse 1 : Les caractéristiques liées à l'apport de C labile (taux d'allongement des racines, teneur en
283 hémicellulose, biomasse racinaire) favorisent l'accumulation de C dans les bassins protégés de limon
284 grossier et de limon fin + argile par activité microbienne. Les caractères racinaires liés à la récalcitrance
285 (teneur élevée en lignine et en cellulose, rapport C:N élevé) favorisent l'accumulation de C dans le mélange
286 de matière organique en particules (POM) grossier non protégé.

287 ii. Objectif 2 : Quel est l'effet de la sélection des espèces sur la séquestration du carbone dans différents
288 bassins de carbone du sol ?

289 Hypothèse. 2 : Les espèces fixatrices de diazote (N₂) favorisent l'accumulation de C dans les bassins
290 protégés de limon fin+argile car elles ont des caractéristiques plus liées à l'apport de C labile, tandis que
291 les espèces non fixatrices de N₂ favorisent l'accumulation de C dans la fraction POM.

292

293 *Chapitre III : Les destins du carbone du sol nouveau et ancien diffèrent dans le sol superficiel et le*
294 *sous-sol nouvellement exposé et s'expliquent par les traits racinaires, des microbes et des particules*
295 *du sol.*

296 i. Objectif 1 : Quantifier les flux de nouveau C introduit par les plantes et de vieux C préexistant dans
297 différents bassins de sol;

298 Hypothèse 1 : les fractions granulométriques du pétrole associées aux fractions de taille des particules du
299 pétrole peuvent réguler les destins de l'ancien C et du nouveau C dans le processus de séquestration du C

300 ii. Objectif 2 : Rechercher l'effet synergique de la nouvelle entrée C et des changements de l'ancienne
301 entrée C dans les différents bassins de carbone.

302 Hypothèse 2 : Le sort du nouveau C et de l'ancien C montrera des modèles indépendants.

303 iii. Objectif 3 : Étudier si les différents acteurs impliqués dans le stockage du carbone et l'influence que
304 la plante et le sol ont sur eux peuvent expliquer les schémas des nouveaux flux de carbone et des
305 anciens flux de carbone dans différents bassins de carbone du sol.

306 Hypothèse 3 : Nous faisons l'hypothèse que les traits racinaires liés à la composition chimique et à la
307 récalcitrance entraîneront une nouvelle accumulation de C dans la POM, tandis que les traits liés à un
308 apport élevé en C entraîneront le stockage dans des fractions protégées par consommation et dépôt
309 microbiologiques. Je m'attends à ce que la stabilité des agrégats soit corrélée positivement avec la
310 nouvelle accumulation de C total et dans le POM fin et les fractions grossières de limon en raison de la

311 protection physique des agrégats. Nous nous attendons à ce que la teneur en N du sol soit positivement
312 corrélée avec la nouvelle teneur en C. Nous pensons que la fraction fine dans le sol est corrélée
313 positivement avec le nouveau stockage du carbone dans la fraction de limon fin + argile en raison des
314 interactions organominérales, et que le nouveau stockage du carbone dans le limon fin + argile est plus
315 élevé dans le sous-sol que dans le sol superficiel en raison des niveaux inférieurs de saturation en carbone
316 du sol. Enfin, je m'attends à ce que l'activité, la diversité et l'abondance microbiennes soient fortement
317 liées à la quantité de nouveau C déposé dans les fractions de limon grossier et de limon + argile protégées,
318 et à la consommation et à la transformation du nouveau C dans les fractions grossières et fines non
319 protégées du POM et du POM fin en raison des minéralisations des communautés microbiennes.

320

321 *Chapitre IV : La qualité du sol détermine le 'priming effect' et les espèces végétales l'affinent : le rôle*
322 *de la préférence du substrat et de la concurrence dans le sol superficiel et le sous-sol*

323 i. Objectif 1 : Quantifier les changements dans le C et l'apport de nouveau C dans le sol pour
324 déterminer les pertes de l'ancien C dans le sol superficiel et le sous-sol remontés à la surface
325 et revégétalisé et le 'priming effect' de la revégétalisation avec des espèces fixant N₂
326 (*Medicago sativa*) et une espèce non fixant N₂ (*Lolium perenne*)

327 Hypothèse 1 : Notre hypothèse est que le sol superficiel aura des pertes plus élevées de vieux C en raison
328 de la biomasse et de l'activité microbienne plus élevées. Cependant, en raison de la plus grande protection
329 du vieux C dans le sous-sol et des changements des conditions environnementales dus à la
330 revégétalisation, nous émettons l'hypothèse que le sous-sol aura des pertes de vieux C plus élevées que
331 le sol nu, ce qui signifie un 'priming effect' positif plus élevé que le sol de surface.

332 ii. Objectif 2 : Quantifier le 'priming effect' dans différents bassins C liés aux fractions
333 granulométriques du sol.

334 Hypothèse 2 : Étant donné la protection plus élevée de C dans la fraction plus fine du sol (fractions limon
335 et limon + argile), nous supposons que le 'priming effect' se produira dans les fractions de matière
336 organique particulaire non protégée (POM et POM fin).

337 iii. Objectif 3 : Étudier l'évolution dans le temps des sources de C respiré dans le système (représentées
338 par l'abondance du ¹³C) et ses corrélations avec les nouvelles pertes de C, les nouvelles entrées de
339 C et le priming effect.

340 Hypothèse 3 : La source de respiration dans le système sol-plante se tournera davantage vers les intrants
341 végétaux marqués au fil du temps, avec le développement des plantes. Les nouveaux apports de C seront
342 positivement corrélés avec l'abondance du ¹³C dans le CO₂ respiré (A13C). Cependant, je m'attends à des
343 comportements différents dans les deux sols en ce qui concerne les pertes de nouveau C. Dans le sol
344 superficiel, je suggère que l'A13C sera corrélé négativement avec les pertes de nouveau C, en raison d'un
345 apport élevé de nouveau C dans le système et de l'utilisation accrue de nouveau C comme substrat pour
346 la croissance microbienne (reflétée par un A13C supérieur). Dans le sous-sol, je fais l'hypothèse d'une
347 corrélation positive entre l'A13C et les pertes de nouveau C, puisqu'un faible apport de nouveau C
348 augmentera l'activité microbienne qui, ne pouvant satisfaire leurs besoins énergétiques principalement à
349 partir de ces sources labiles, exploitera le nouveau C plus efficacement. De la même façon, le 'priming
350 effect' sera corrélé négativement à A13C dans le sol arable, tout en étant corrélé positivement dans le
351 sous-sol.

352

353 APPROCHE ET CONCEPTION EXPERIMENTALE

354 Pour atteindre ces objectifs, deux expériences ont été conçues et réalisées dans le cadre de ce projet de
355 recherche.

356 Dans la première expérience, 12 espèces herbacées différentes ont été cultivées en monoculture dans 72
357 boîtes de culture (six répétitions par espèce). Sur ces six répétitions, la moitié a été utilisée pour
358 l'échantillonnage du sol, tandis que l'autre moitié a été cultivée dans des boîtes munies de fenêtres en
359 PVC utilisées pour observer la croissance des racines. Toutes les deux semaines, chaque fenêtre de racines
360 a été photographiée pour évaluer le taux d'allongement des racines et les caractéristiques des racines.
361 Après 10 mois, des carottes de sol ont été prélevées pour évaluer 1) les caractéristiques architecturales
362 des racines, 2) la composition chimique des racines, 3) le carbone du sol dans quatre fractions de sol
363 différentes (POM <200 µm ; POM fin 50-200 µm, limon 20-50 µm, limon + argile <20 µm), 4) la respiration
364 microbienne induite (SIR) du substrat comme indicateur d'activité microbiologique.

365 Dans la deuxième expérience, deux des 12 espèces présentant des tendances aux extrémités opposées du
366 spectre économique racinaire (*Lolium perenne* et *Medicago sativa*) ont été sélectionnées et cultivées en
367 monoculture en pots. Les pots ont été cultivés dans des microcosmes avec des conditions
368 environnementales constantes et du CO₂ atmosphérique constamment enrichi en ¹³C pendant 183 jours
369 sur deux types de sol. Les deux types de sol, soit le sol superficiel (0-30 cm) et le sous-sol (110-140 cm),
370 ont été extraits du même profil de sol à Pisciotta (SA), en Italie. Les sols étaient argileux et présentaient
371 des caractéristiques contrastées (teneur en azote, stabilité des agrégats, biomasse et activité
372 microbienne). De plus, dans le sol superficiel, la teneur en argile était légèrement inférieure à celle du
373 sous-sol (-8 %) et la teneur en C nettement supérieure (sol superficiel 12 mgC g⁻¹ sol ; sous-sol 6 mgC g⁻¹
374 sol), ce qui entraîne un niveau de saturation en C supérieur. Les plantes et les sols ont été croisés et six
375 pots répliqués ont été semés. Par ailleurs, six pots témoins nus (non semés) ont été mis en place pour
376 chaque sol. Toutes les deux semaines, la respiration du sol était échantillonnée pour évaluer ¹³C% de CO₂
377 respiré et le CO₂ dérivé de la plante, et après six mois, les pots étaient collectés et le sol échantillonné pour
378 une gamme de différentes caractéristiques du sol, des racines et des caractéristiques microbiologiques.

379 PRINCIPAUX RESULTATS

380 Au chapitre II, nous n'avons observé aucun effet significatif des espèces sur l'accumulation de C dans les
381 différents gisements de C associés aux fractions du sol. Cependant, lorsque nous avons observé l'effet de
382 la famille, les espèces de Fabaceae fixatrices de N₂ ont accumulé plus de C dans la fraction de limon fin
383 protégée, tandis que les espèces de Poaceae non fixatrices de N₂ dans la fraction POM. Les caractéristiques
384 des racines différaient significativement entre les deux familles, les Poaceae ayant des tissus plus
385 récalcitrants (lignine et cellulose élevées, et rapport C:N élevé), une biomasse racinaire plus faible et un
386 taux d'allongement des racines plus faible. Les espèces de Fabaceae présentaient des tissus plus labiles
387 (hémicellulose élevée et faible rapport C:N), une biomasse plus élevée et un taux d'élongation des racines
388 plus élevé. Les espèces de Fabaceae ont également augmenté l'activité microbienne. Grâce à l'analyse en
389 composantes principales et aux corrélations de Pearson, nous avons montré l'effet d'un apport élevé de C
390 labile (typique des espèces acquisitrices à croissance rapide) entraînant une accumulation plus élevée dans
391 la fraction de limon protégée. Les espèces conservatrices à croissance lente, à l'autre extrémité du spectre
392 économique racinaire, augmentent l'accumulation de C dans la POM non protégée. Cette différence dans
393 les stratégies d'accumulation de C a confirmé la corrélation entre le spectre économique racinaire et le
394 stockage de C dans différents bassin de C. Cette différence était due à l'effet de l'espèce sur l'activité
395 microbienne. Une activité microbienne élevée chez les espèces de Fabaceae a favorisé la minéralisation
396 de l'intrant C et son entombage dans la fraction limoneuse, tandis que l'activité microbienne plus faible
397 chez les espèces de Poaceae a diminué la décomposition et la minéralisation du C introduit par rotation
398 des racines et augmenté sa stabilité et son accumulation dans la fraction POM.

399 Dans le chapitre III, nous avons montré comment l'apport de C dérivé de nouvelles plantes et les pertes
400 de C préexistant étaient en synergie, avec un apport plus élevé de nouveau C diminuant les pertes d'ancien
401 C. Les espèces plantées en surface ont considérablement augmenté l'apport de nouveau C dans le sol et

402 diminué le rendement du C ancien. En particulier, *M. sativa* avait un apport plus élevé et des pertes plus
403 faibles que *L. perenne*. L'apport de nouveau C s'est principalement fait au niveau des fractions POM et
404 limons fins+argile dans les sols. Dans le sol superficiel, l'ancien C a diminué dans tous les gisements sauf
405 dans la fraction limons fins+argile, où il s'est accumulé. Dans le sous-sol, l'ancien C a diminué dans tous les
406 gisements sauf dans le POM, où la diminution n'a pas été significative. Les différentes caractéristiques des
407 racines, des microbes et du sol étaient mieux corrélées avec l'apport de nouveau C dans les fractions que
408 les changements de l'ancien C. Les anciennes pertes de C semblaient plus liées au choix du sol et "intrinsic"
409 au système du sol. Cette entrée de nouveau C était principalement corrélée positivement
410 avec la production de biomasse racinaire, tandis que le rapport C:N était corrélé négativement avec la
411 nouvelle entrée de C dans les fractions POM et limons fins+argile. Les caractères racinaires sont mal
412 corrélés aux variations de quantité de l'ancien C. Les caractéristiques microbiologiques ont été le principal
413 facteur à l'origine des nouveaux apports de C, corrélés positivement à l'augmentation du nouveau C dans
414 chaque fraction. Ils étaient aussi positivement corrélés avec l'ancienne accumulation de C dans la fraction
415 limons fins+argile. En ce qui concerne les caractéristiques du sol, la stabilité des agrégats et la teneur en N
416 étaient en synergie et en corrélation positive avec les nouveaux apports de C dans le système et
417 l'accumulation des anciens C dans la fraction limons fins+argile. La fraction fine du sol (<20µm) a été
418 corrélée négativement avec la nouvelle entrée de C et, étonnamment, l'ancienne accumulation de C dans
419 la fraction limons fins+argile. Ces résultats ont clairement montré comment le type de sol constitue le
420 facteur principal influant sur le stockage et le cycle du carbone dans le sol car la fertilité et l'activité
421 microbienne du sol constituent le moteur de la séquestration du carbone. Les espèces végétales ont un
422 effet secondaire sur le stockage et le cycle du carbone dans le sol. *M. sativa* est l'espèce la plus influente
423 parmi les 12 étudiées et agit en augmentant l'apport de nouveau C grâce à une production plus élevée de
424 tissus labiles et une activité microbienne accrue. Une faible saturation en C du sol ne semble pas avoir
425 d'influence positive sur le stockage du C dans la fraction limons fins+argile. Cependant, lorsque l'apport de

426 nouveau C dans la fraction limons fins+argile est normalisé par la biomasse racinaire (pour estimer l'apport
427 de nouveau C par g de racine), le sous-sol a un rendement de stockage C supérieur à celui de la terre
428 végétale, et *L. perenne* a un apport supérieur par g de biomasse produite. Une saturation plus faible en C
429 pourrait donc avoir un effet positif sur le stockage du C dans le sol, mais cet effet est atténué par la fertilité
430 du sol (qui détermine la production de biomasse et l'apport de C dans le système) et l'activité microbienne
431 (qui transforme l'apport de C et le transfère dans la fraction limons fins+argile par métabolisme
432 microbien). Dans ce chapitre, nous montrons clairement la puissance du couplage des techniques de
433 marquage isotopique avec le fractionnement du sol pour décrire efficacement les changements de C dans
434 le sol et étudier leurs corrélations avec les différents acteurs impliqués.

435 Enfin, dans le Chapitre IV, nous montrons une fois de plus comment le sol est l'élément principal qui
436 façonne le 'priming effect', avec des pertes de carbone plus élevées dans le sol superficiel que dans le
437 sous-sol en raison de la biomasse et de l'activité microbienne accrues, mais un 'priming effect' négatif dans
438 le sol superficiel et positif dans le sous-sol. L'augmentation de l'apport de nouveau C dans le sol favorise
439 le passage de la préférence pour le substrat des plantes de l'ancien C préexistant à l'apport de nouveau C.
440 Ceci peut être observé dans les résultats de respiration du sol : dans le sol superficiel, l'augmentation de
441 la signature du ¹³C au cours des six mois est supérieure à celle de le sol superficiel, atteignant une quantité
442 plus élevée de CO₂ provenant de la minéralisation du nouveau C introduit par rapport à l'ancien C
443 préexistant dans le sol sol. La quantité totale d'ancien C consommée dans un sol végétalisé diminue par
444 rapport à un sol nu, ce qui entraîne un 'priming effect' négatif. Dans le sous-sol, l'apport de nouveau C
445 n'est pas assez élevé pour permettre le changement de préférence du substrat, et les communautés
446 microbiennes continuent d'utiliser l'ancien C préexistant pour l'acquisition des nutriments. Dans le sous-
447 sol, nous pouvons observer un effet significatif des espèces, *M. sativa* ayant un 'priming effect' positif plus
448 élevé que *L. perenne*. Ceci peut s'expliquer par la concurrence entre les communautés microbiennes et les

449 plantes pour l'acquisition d'azote. L'absorption racinaire par *L. perenne* concurrence les communautés
450 microbiennes pour l'acquisition d'azote et réduit leur activité, ce qui réduit globalement leur efficacité à
451 consommer l'ancien C et entraîne un faible 'priming effect' (pas significativement différent du sol nu). *M.*
452 *sativa* étant une espèce fixatrice de N₂, elle ne concurrence pas les microorganismes pour le N.
453 L'augmentation de l'intrant de C labile augmente en fait la biomasse et l'activité microbiennes et
454 l'extraction de l'ancien C pour l'exploitation des ressources. Pour cette raison, *M. sativa* a un 'priming
455 effect' positif plus élevé. Dans ce chapitre, nous réconcilions les théories de la préférence pour le substrat
456 et celles de la concurrence, qui déterminent le 'priming effect' et dépendent de la fertilité du sol et,
457 ensuite, des espèces végétales. Le 'priming effect' de la fertilité du sol se manifeste par la préférence du
458 substrat, le sol fertile permettant aux communautés microbiennes de changer de substrat et ayant pour
459 résultat un 'priming effect' négatif. Les sols pauvres ne permettent pas le changement de substrat et
460 donnent lieu à un 'priming effect' positif, dont l'ampleur est déterminée par l'absence de concurrence
461 microbienne pour l'azote par les plantes. Nous avons également observé que le 'priming effect' dans le
462 sous-sol était plus élevé dans les fractions limon et limon fin + argile, remettant en question la stabilité
463 effective de ces fractions.

464

465 CONCLUSION ET APPLICATION PRATIQUE

466 Avec cette recherche, nous avons mis en évidence comment les espèces fixant N₂ sont plus efficaces pour
467 la séquestration du C grâce à un apport plus élevé de C labile qui augmente le stockage total du C, plus
468 particulièrement dans les bassins de C stables limon et limon + argile. L'apport plus élevé dans les bassins
469 protégés limon et limon + argile est lié aux caractéristiques racinaires liées à la labilité (en particulier les
470 caractéristiques chimiques des racines) qui augmentent l'activité microbologique. Dans cette perspective,
471 l'étude du spectre économique racinaire est un outil prometteur pour établir un lien entre les traits
472 racinaires et la séquestration du carbone. La symbiose avec la bactérie *Rhizobium* joue également un rôle
473 important en augmentant la production et le dépôt d'exopolysaccharides dans les fractions fines du sol.

474 Le sol est le principal facteur qui influe sur le stockage du C, et l'analyse des bassins de carbone liés aux
475 fractions du sol couplé à l'expérience de l'étiquetage isotopique est une méthodologie puissante pour
476 démêler les mécanismes du cycle C. Le sol superficiel a un apport plus élevé en C en raison d'une fertilité
477 plus élevée et d'une activité microbienne plus élevée, ce qui augmente le dépôt de C dans la fraction
478 protégée de limon et de limon + argile. Le sol superficiel a également moins de pertes de carbone ancien
479 grâce au passage de la consommation préférentielle de substrat de l'ancien C vers le nouveau C des
480 communautés microbologiques. Globalement, l'effet de la saturation en C sur le stockage du C dans la
481 fraction limon + argile semble être soumis à la qualité du sol en termes de teneur en N et d'activité
482 microbologique. Cependant, lorsque ces exigences sont satisfaites, il peut stocker C plus efficacement,
483 comme le suggère la quantité plus élevée de C déplacée dans les fractions de limon + argile par g de racine
484 dans le sous-sol (faible saturation en C) par rapport à la terre végétale (saturation en C élevée).

485 Le sol est également le principal moteur de le 'priming effect', le sol superficiel présentant un 'priming
486 effect' négatif en raison du passage des communautés microbologiques de l'ancien C au nouveau C. Dans
487 le sous-sol, le 'priming effect' est positif et la concurrence détermine son ampleur : *L. perenne* diminue le

488 'priming effect' positif (presque aucun 'priming effect') grâce à la compétition pour l'azote qui inhibe
489 l'activité microbologique. *M. sativa*, d'autre part, augmente l'azote du sol grâce à sa capacité de fixation
490 de l'azote et augmente l'activité microbologique, ce qui augmente globalement le 'priming effect' positif.

491 Une de nos principales indications pratiques est de ne pas considérer le potentiel de stockage du C du sol
492 uniquement du point de vue de la minéralogie, de la teneur en argile ou de la saturation en C, mais de
493 faire attention à la santé du sol. Plus spécifiquement, pour évaluer ses niveaux de fertilité (teneur en N),
494 la stabilité des agrégats (MWD) et le développement des communautés microbiennes (évaluation de leur
495 biomasse et/ou activité). La diversité microbienne pourrait également être un indicateur important. Ces
496 indicateurs sont liés à un apport plus élevé de C dans le sol par le biais d'une production accrue de
497 biomasse, d'un transfert vers un bassin limons fins+argile protégé et d'un 'priming effect' négatif dû à un
498 changement d'utilisation du substrat.

499 L'utilisation de le sol superficiel fertile augmente l'accumulation de carbone par rapport à un sous-sol
500 pauvre et il est donc souhaitable pour la revégétalisation des sols géotechniques. Lors de la
501 revégétalisation de le sol superficiel fertile, les espèces à croissance rapide qui fixent l'azote (c.-à-d. les
502 légumineuses) avec un apport élevé de C labile sont plus efficaces pour stocker le C dans un bassin protégé
503 de limons et de limons fins+argile par un apport racinaire et un renouvellement microbien *in vivo* plus
504 élevés. De plus, la revégétalisation de le sol superficiel a un 'priming effect' négatif, ce qui augmente la
505 stabilité du C préexistant.

506 Cependant, l'utilisation de terre végétale n'est pas toujours possible. Certaines conditions particulières
507 peuvent nécessiter la revégétalisation du sous-sol ; par exemple, en cas d'indisponibilité de le sol
508 superficiel fertile, l'impact écologique de l'enlèvement de le sol superficiel fertile d'une zone différente,
509 ou en raison de vastes zones excavées qui seraient trop coûteuses économiquement et écologiquement

510 pour être couvertes de terre végétale fertile (comme de vastes carrières). Dans ce cas, nous conseillons

511 de:

512 1) Fertiliser le sol pour augmenter la production de biomasse et le stockage du C.

513 2) Ensemencer avec des communautés microbiennes.

514 De plus, la saturation en C basse n'augmente pas le stockage du C protégé dans le sous-sol dans notre

515 expérience, mais elle est encore prometteuse pour le stockage potentiel du C si la fertilité et les exigences

516 microbiennes sont respectées. Si la fertilisation et l'inoculation microbienne sont impossibles, nous

517 suggérons d'éviter l'utilisation d'espèces fixant l'azote, car l'augmentation de la biomasse microbienne liée

518 à ces espèces entraînerait une minéralisation plus importante de l'ancien C.

519