Soil fauna: key to new carbon models

Authors

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Abstract

Soil organic matter (SOM) is key to maintaining soil fertility, mitigating climate change, combating land degradation, and conserving above- and below-ground biodiversity and associated soil processes and ecosystem services. In order to derive management options for maintaining these essential services provided by soils, policy makers depend on robust, predictive models identifying key drivers of SOM dynamics. Existing SOM models and suggested guidelines for future SOM modelling are defined mostly in terms of plant residue quality and input and microbial decomposition, overlooking the significant regulation provided by soil fauna. The contribution of soil fauna activities to SOM decomposition can be as high as 40%, as they control almost any aspect of organic matter turnover, foremost by regulating the activity and functional composition of soil microorganisms and their physico-chemical connectivity with soil organic matter. We suggest that inclusion of soil animal activities (plant residue consumption and bioturbation altering the formation, depth, hydraulic properties and physical heterogeneity of soils) can fundamentally affect the predictive outcome of SOM models. Understanding direct and indirect impacts of soil fauna on nutrient availability, carbon sequestration, greenhouse gas emissions and plant growth is key to the understanding of SOM dynamics in the context of global carbon cycling models. We argue that explicit consideration of the soil fauna is essential to make realistic modelling predictions on SOM dynamics and to detect expected non-linear responses to global change, and we suggest that guidelines for future SOM modelling should implement the role of soil fauna. Finally, we briefly introduce the new COST (European Cooperation in Science and Technology) Action ES 1406 (KEYSOM – Soil fauna: key to new carbon models) which brings together biogeochemists and soil ecologists from 21 EU countries. KEYSOM provides a research network for improved SOM models by implementing the role of the soil fauna as a basis for sustainable soil management. An interdisciplinary platform of experimentalists and modellers will assure better access to experimental data, identify the most burning research gaps and inform decision makers.

Keywords

Carbon turnover, soil organic matter modelling, soil invertebrates, aggregate formation, spatial heterogeneity, regional differences, COST Action

1. Introduction

Despite continuous refinement over the past decades, estimates of the global carbon cycle still show large discrepancies between potential and observed carbon fluxes (Ballantyne et al., 2012; Schmitz et al., 2014). Soil organic matter (SOM) modelling is key to understand and predict changes in global carbon cycling and soil fertility in a changing environment. SOM models can facilitate a better understanding of the factors that underlie the regulation of carbon cycling and the persistence of SOM. The predictive power of current global SOM models is, however, limited, as the majority relies on a relatively restricted set of input parameters such as climate, land use, vegetation, pedological characteristics and microbial biomass (Davidson and Janssens, 2006). In an overview on the performance of SOM models, none of 11 tested models could predict global soil carbon accurately, nor were 26 regional models able to assess gross primary productivity across the US and Canada (Luo et al., 2015). Recently, Schmidt et al. (2011) proposed eight “key insights” to enrich model predictions on the persistence of SOM. However, they ignored a major component of SOM dynamics, soil fauna, which play a fundamental role in most of the insights they propose (e.g. Fox et al., 2006; Jimenez et al., 2006; Osler and Sommerkorn, 2007; De Deyn et al., 2008; Wilkinson et al., 2009). By moving through and
arious trophic groups of aboveground and belowground actors, CPMAS 13C NMR (Cross-Polarization Magic Angle Spinning) spectra showed that earthworm casts and surrounding soil were dominated by carbohydrates, with a decrease of O-alkyl C and an increase of alkyl C. Due to their prime role in most processes in soil (Briones et al., 2012; Lavelle et al., 1998; Lavelle et al., 2006; Osler and Sommerkorn, 2007; Brussaard et al., 2007; Sanders et al., 2014), without any further change in the structure of carbon models. This was partly due to a lack of communication between modellers and experimenters, but also because the magnitude of animal effects on SOM dynamics remains poorly quantified (Schmitz et al., 2014).

In a review focusing mostly on large mammals, terrestrial herbivores and aquatic ecosystems, Schmitz et al. (2014) recently called for “animating the carbon cycle”. Bardgett et al. (2013) argued that differential responses of various trophic groups of aboveground and belowground organisms to global change can result in a decoupling of plant-soil interactions, with potentially irreversible consequences for carbon cycling. Other research has shown that including soil animal activities could help clarifying the discrepancies in existing carbon models, based on a large-scale correlative field study (de Vries et al., 2013). Similar attempts to connect animal activity to carbon cycling occurred in the past (e.g. Lavelle and Martin, 1992; Lavelle et al., 1998; Lavelle and Spain, 2006; and Schmitz et al., 2014) have mostly focused on earthworms, but also give examples for other groups of soil fauna whose role in C turnover appears to be much more relevant than thought thus far (e.g. David, 2014). We point out regional differences in climate, soils and land use and derive implications for modelling. Finally, we introduce a new COST Action (ES 1406) that is working on the implementation of soil fauna into SOM models, also exploring the pros and caveats in such a process.

2. Key insights

The eight ‘key insights’ compiled by Schmidt et al., 2011 are shown in Fig. 1, together with the most important activities of soil animals affecting them. As many animal-mediated processes are tightly interconnected, they also matter for several of these insights. For instance, aggregate formation in faeces simultaneously affects molecular structure, humic substances, physical heterogeneity and soil microorganisms. In the following text we briefly summarize the role of animal activities for each of the ‘key insights’. As a more detailed example of animal impacts on SOM turnover, we consider their role on soil aggregate formation in a separate section. Table 1 contains quantitative examples of animal activity taken from different biomes and land use types, showing that earthworms alone strongly affect each of the ‘key insights’. However, much smaller soil animals can also have substantial effects on them (Table 1).

2.1. Molecular structure

Many soil animals ingest and process SOM (and accompanying microorganisms) in their gut system, where it is partly assimilated with the help of mutualistic gut microflora and partly egested. Metabolisation alters the chemical structure of ingested SOM (Jiménez and Lal, 2006; Hedde et al., 2005; Coulis et al., 2009; Frouz, 2015b; Schmitz et al., 2014) and consequently, the decomposition dynamics of animal faeces, which can be a substantial component of SOM (humification). For instance, earthworm casts have species-specific NIR spectral signatures, indicating presence of specific organic compounds (Hedde et al., 2005). Under grass/legume pasture they are characterized by significant enrichment of slightly-altered plant residues in the sand particle size (> 53 μm). CPMAS 13C NMR (Cross-Polarization Magic Angle Spinning Carbon-13 Nuclear Magnetic Resonance) spectra showed that earthworm casts and surrounding soil were dominated by carbohydrates, with a decrease of O-alkyl C and an increase of alkyl C.
merkorn, organic matter in young soils and humic substances are mainly faeces of earthworms and diplopods (see also Dolmer et al., 2014; Malmström, 2012), and that earthworms likely possess a unique capability of neutralizing plant polyphenols that otherwise strongly decrease decomposition rates of fresh plant litter (Liebeke et al., 2015). Micro- and mesofauna excrete ammonium or dissolved organic carbon (Filser, 2002; Fox et al., 2006; Osler and Sommerkorn, 2007), and affect the quantity of microbial metabolites (Bonkowski et al., 2009). Gut passage, defecation, excretion together with bioturbation by macro- and mesofauna facilitate humification and decomposition, altering also nutrient stoichiometry (Bohren et al., 2004). These modifications in the molecular structure of SOM due to soil fauna activity have significant effects on its dynamics.

2.2. Humic substances

The term “humic substances” nowadays is considered outdated: neither is the concept itself clear, nor is there any evidence that the often mentioned highly complex large organic molecules play any relevant role under natural conditions (Schmidt et al., 2011; Lehmann and Kleber, 2015). However, here we stick to it when referring to the “insights”, simply for reasons of consistency with the article our argumentation is based on (Schmidt et al., 2011). We acknowledge that “humus” or “humic substances” represent a continuum of more or less decomposed dead organic matter of which energy content and molecule size mostly should decrease over time, and that water solubility, sorption to the mineral matrix and accessibility for microorganisms are highly relevant for OM turnover (Lehmann and Kleber, 2015).

Humic substances are formed during the gut passage: organic matter in young soils and humic horizons almost completely consists of soil animal faeces (Lavelle, 1988; Martin and Marinissen, 1993; Brussaard and Juma, 1996). Humus forms mainly comprise animal casts, e.g. casts of ants, isopods, millipedes, beetle larvae or termites in deadwood, of insect larvae and spiders in leaf litter, or casts of collemobolans, mites and enchytraeids in raw humus. In his review, David (2014) considered macroarthropod casts partially stabilized organic matter rather than hotspots of microbial activity. The dark colour of casts (compared to the ingested organic material) visually demonstrates the strong chemical modification in animal guts, which is accompanied by a substantial physical modification. Clay-humus complexes, physically protecting organic matter (Jiménez and Lal, 2006), are mainly faeces of earthworms and diplopods (see also Sect. 2.5 on physical heterogeneity). Due to differences in feeding preferences, gut microflora, SOM quantities consumed etc. of soil animals, their faeces vary in size, shape and quality not only between fauna groups, but also between species within one group (see Sect. 3 on aggregate formation). Discovering the important role of animal faeces in humification is essential to improve our understanding of carbon dynamics in soil.

2.3. Fire-derived carbon

Microbial recolonisation of burned sites is mediated by wind and soil animals that survived in soil or emigrated from neighbouring areas, e.g. by macro- and mesofauna, birds and mice (Zaitsev et al., 2014; Malmström, 2012). Besides, soil fauna also ingest the charcoal particles (Eckmeier et al., 2007; see Table 1). Due to animal activity, charcoal is sorted by size and translocated down the soil profile. Mice and earthworms (Eckmeier et al., 2007) and the tropical earthworm Pontoscolex corethrus (Topolianz and Ponge 2003; Topolianz et al., 2006) had been suggested as responsible for rapid incorporation of charcoal into the soil. Quantitative data are however scarce (Table 1). In spite of potentially great importance, the effect of soil animals on the fate of the “black carbon” in soil remains practically unknown (Ameloot et al., 2013).

2.4. Roots

Plant roots preferably grow in existing soil cavities (Jiménez and Lal, 2006), mostly formed by soil fauna (Wilkinson et al, 2009). Both burrowing and non-burrowing soil animals have a strong
impact on root growth, allocation, length and density (Brown et al., 1994; Bonkowski et al.,
2009; Arnone and Zaller, 2014). Animal grazing of root bacteria and mycorrhiza affects their
activity and community composition, and animal excreta are enriched in micronutrients and
selectively affect plant nutrition (Brown, 1995; Filser, 2002; Brussaard et al., 2007). Root
herbivores and rhizosphere grazers affect C allocation of roots (Wardle et al., 2004) and largely
regulate nutrient acquisition and plant productivity (Bonkowski et al., 2009). Not only root
herbivores, but also saprotrophic/microbivorous soil animals may obtain a significant proportion
of energy from plant roots (Pollierer et al., 2007). This suggests an animal-mediated regulatory
loop that connects plant roots and SOM.

2.5. Physical heterogeneity

Animals fragment any organic residues, perform bioturbation, distribute organic matter and
generate smaller and larger organic matter hot spots (e.g. faecal pellets, ant and termite mounds).
According to body size, they act at various spatial scales, from micro-aggregates to landscapes
(Éttema & Wardle, 2002; Jouquet et al., 2006). Mounds and burrows made by soil fauna are
obvious signs of physical heterogeneity created by ecosystem engineers (Meysmann et al., 2006;
Wilkinson et al., 2009; Sanders et al., 2014), which significantly affect microorganisms, plants
(Chauvel et al., 1999; Frelich et al., 2006), aggregate stability (Bossuyt et al., 2005; 2006),
hydraulic properties (Bottinelli et al., 2015; Andruzzoli et al., 2016), sorption and degradation of
sparingly soluble organic compounds (Edwards et al., 1992; Bolduan and Zehe, 2006) and C
emissions (Wu et al., 2013; Lopes de Gerenyo et al., 2015). Earthworms in particular feed on
organic and mineral parts of the soil and mix them (Eckmeier et al., 2007; Wilkinson et al.,
2009). The resulting clay-organic matter complexes considerably increase SOM retention of soils
(Jiménez and Lal, 2006; Fox et al., 2006; Brussaard et al., 2007), although C loss from fresh casts
is much higher than from surrounding soil (Zangerlé et al., 2014). The impact on soil processes
and physical heterogeneity varies considerably between different groups of ecosystem engineers
(Jouquet et al., 2006; Bottinelli et al., 2015). For instance, some earthworm species strongly
affect their physical environment while others are more affected by the soil organic matter
content (Jiménez et al. 2012).

2.6. Soil depth

A considerable part of physical heterogeneity are animal burrows that can reach several meters
deep. Bioturbation (e.g. by earthworms, termites, ants, beetle and Diptera larvae, spiders, solitary
bees and wasps, snails, isopods and amphipods, ground owls, lizards, porcupines, pigs, moles,
voles, rabbits, foxes, or badgers) is a key process to the formation of soil depth, soil structure and
associated C translocation, as shown by several examples in Table 1 and reviewed e.g. by
Wilkinson et al. (2009).

2.7. Permafrost

Due to unfavourable environmental conditions (resulting in low animal biomass, activity and
diversity), there is only little impact of fauna in permafrost soils (De Deyn et al., 2008). However,
fauna invasions, especially of above-mentioned soil engineers, due to soil melting in tundra and
boreal forests are likely to have drastic effects (Frelich et al., 2006; Van Geffen et al., 2011). Data
on earthworm invasions in North American forests (Bohlen et al., 2004; Frelich et al., 2006;
Eisenhauer et al., 2007) show that they must be taken into consideration in carbon-rich soils,
particularly in melting permafrost soils (Frelich et al., 2006; Schmidt et al., 2011) where they
may affect many soil functions.
2.8. Soil microorganisms

Soil fauna comprise ecosystem engineers as well as an armada of mobile actors connecting elements of the soil system, mediating microbial processes (Briones, 2014). Countless isopods, ants, termites, enchytraeids, microarthropods, nematodes or protozoans make large contributions to SOM turnover underground (Persson, 1989; Filser, 2002; Wardle et al., 2004; Fox et al., 2006; Osler and Sommerkorn, 2007; Wilkinson et al., 2009; Wu et al., 2013). They affect the activity and community composition of soil microorganisms in multiple ways such as feeding, burrowing, facilitating the coexistence of different fungal species (Crowther et al., 2011) or by modifying micro-habitat conditions. Litter comminution by detritivores increases SOM accessibility for microorganisms, and propagules are dispersed with body surface and casts. Gut incubation provides protected microsites with modified environmental conditions and increases bacterial abundance substantially – e.g. three orders of magnitude in earthworm guts (Edwards and Fletcher, 1988). Grazing affects microbial biomass, activity and community composition, and animal excreta modify nutrient availability for microorganisms (Brown, 1995; Filser, 2002).

3. Aggregate formation

The modern view on the stability of organic matter on soils requires a thorough understanding of aggregate structure and formation including the role of soil biota (Lehmann and Kleber, 2015). Soil aggregation is the process by which aggregates of different sizes are joined and held together by different organic and inorganic materials. Thus, it includes the processes of formation and stabilization that occur more or less continuously, and can act at the same time. With clay flocculation being a pre-requisite for soil aggregation, the formation of aggregates mainly occurs as a result of physical forces, while their stabilization results from a number of factors, depending in particular on the quantity and quality of inorganic and organic stabilizing agents (Amézketa, 1999).

By bioturbation, feeding and dispersal of microbial propagules soil animals regulate all of the above forces and agents, and are therefore a crucial factor in the formation and stabilisation of soil aggregates. Earthworms, many insect larvae and other larger fauna may stabilize aggregate structure by ingesting soil and mixing it intimately with humified organic materials in their guts, and egesting it as casts or pellets (Tisdall and Oades, 1982; Oades, 1993). Earthworms have a direct and fast impact on microaggregate formation and the stabilization of new C within these microaggregates (Bossuyt et al., 2005) (Table 1). There are several mechanisms to explain the increase of micro- and macroaggregate stability by earthworms, but no mechanism has been quantified in relation to population size yet. Effects are related to ecological groups of earthworms, associated with feeding habit, microhabitat in the soil profile, and burrow morphology. However, irrespective of this classification, species may enhance or mitigate soil compaction (Blanchart et al., 1997; Guéi et al., 2012). The tensile strength of casts appears to be species dependent: for example, the casts of Dendrobaena octaedra have a lower tensile strength compared to those of L. terrestris (Flegel et al., 1998). Similarly, organic carbon and water-stable aggregation was significantly higher in casts of L. terrestris than in casts of A. caliginosa (Schrader and Zhang, 1997).

Some research, however, suggests that earthworm activity can also evoke soil degradation. Shipitalo and Protz (1988) proposed that ingestion of soil by earthworms results in disruption of some existing bonds within micro-aggregates and realignment of clay domains. Therefore, fresh casts are more dispersible than uningested soil, contributing to soil erosion and crusting. Significant improvement in the water stability of fresh, moist casts only occurs when incorporated organic debris from the food sources is present and when moist casts are aged or dried. Nevertheless, in the long term, casting activity enhances soil aggregate stability.

Our understanding of the contribution of soil fauna to aggregate formation and stabilization is limited, however, and mostly qualitative in nature. Different methodologies complicate the
Comparison among aggregate stability data (Amézketa, 1999). Data in terms of functional response to density are limited as many studies have been conducted in arable systems, where the diversity and abundance of soil animals is reduced as a consequence of tillage, mineral fertilizers and pesticide use. Recently, some studies have emerged. A negative correlation between earthworm abundance and total macroaggregates and microaggregates within macroaggregates in arable treatments without organic amendments could be linked to the presence of high numbers of Nematogenia lacuum, an endogenic species that feeds on excrements of other larger epigeic worms and produces small excrements (Ayuke et al., 2011). Under the conditions studied, differences in earthworm abundance, biomass and diversity were more important drivers of management-induced changes in aggregate stability and soil C and N pools than differences in termite populations. Another study highlighted that in fields converted to no-tillage earthworms incorporated C recently fixed by plants and moved C from soil fragments and plant residues to soil aggregates of >1 mm (Arai et al., 2013). Thus, soil management practices altering fauna activities may have a significant effect on the re-distribution of soil organic matter in water-stable aggregates, impacting agronomically favourable size fractions of water-stable macro-aggregates, and water-stable micro-aggregates which are the most important source of carbon sequestration (Šimanský and Kováčik, 2014).

4. Regional differences in climate, soils and land use

Many of our examples refer to earthworms and temperate regions as they have been studied most intensively. However, we suggest that any dominant group of soil fauna, irrespective of body size or the ability to create larger soil structures, may substantially affect carbon dynamics. Table 1 gives a number of respective case studies. More information is needed on how existing abiotic and biotic constraints to SOM decomposition will vary with changing climate and in different regions (Davidson and Janssens, 2006). The key players and specific effects of soil animals vary across space (Fig. 2), with increasing importance for SOM dynamics in humid-warm and nutrient-limited conditions (Perssson, 1989; Filser, 2002; Wardle et al., 2004; Fox et al., 2006; Osler and Sommerkorn, 2007; De Deyn et al., 2008; Briones, 2014). Also ecosystem engineers differ between soil types, biomes and land-use types, from rodents and ants in dry areas to termites, earthworms and millipedes in tropical rainforests. They consume different types of organic matter, make deep or shallow, narrow or wide burrows, and differ in aggregation behaviour (e.g. more or less regularly distributed earthworms versus distinct ant nests and termite mounds). Accordingly, their role in SOM re-distribution and turnover differs as well.

In cold ecosystems – where, together with wetlands and peatlands, the majority of terrestrial carbon is stored (Davidson and Janssens, 2006) – the response of detritivores to climatic change is expected to be most pronounced (Blankinship et al., 2011). Melting of permafrost soil might lead to northward expansion of soil macro-invertebrates, associated with accelerated decomposition rates (van Geffen et al., 2011). Further examples are shown in Table 1.

5. Implications for modelling

As there is no scientific support for the widespread belief in “humic substances” (here defined as very large and highly complex, poorly degradable organic molecules with manifold aromatic rings; Lehmann and Kleber, 2015), the question how long organic carbon remains in soil is largely related to a) physical protection and b) how often the once photosynthesized dead organic matter is recycled in the soil food web. For both processes soil animals are of great importance, as we have shown above. The thermodynamic viewpoint makes the issue even more relevant: reaction speed increases with temperature, but most soil organisms are rather adapted to relatively
cool conditions and might thus be pushed beyond their niche limits – with negative consequences on their activity.

Changes in climate (Blankinship et al., 2011), land use (Tsiafouli et al., 2014), resource availability and biotic interactions (De Vries et al., 2012) alter the distribution, community composition, activity and associated impact of soil animals on distribution and turnover rate of SOM (Wall et al., 2008) to the extent that underlying assumptions of SOM models may no longer be valid (Swift et al., 1998; Bardgett et al., 2013; Schmitz et al., 2014). Therefore it is opportune to include approaches that have been developed during the past decades (Filser, 2002; Jiménez and Lal, 2006; Osler and Sommerkorn, 2007; Brussaard et al., 2007; Meysmann et al., 2006; Wall et al., 2008; Sanders et al., 2014). For instance, implementing earthworm activity in the CENTURY model (Lavelle et al., 2004) revealed a 10% loss of the slow C pool without earthworms within 35 years.

Without considering the role of animals, models are less accurate: in a field study spanning four countries from Sweden to Greece, soil food web properties were equally important as abiotic factors and predicted C and N cycling processes better than land use (De Vries et al., 2013). In their study, earthworms enhanced CO₂ production whereas Collembola and bacterivorous nematodes increased leaching of dissolved organic carbon. Mechanistic experiments confirm that earthworms have a detrimental effect on the greenhouse gas balance under nitrogen-rich conditions (Lubbers et al., 2013) and under no-till (Lubbers et al., 2015). Inclusion of group-specific diversity of mesofauna in models of global-scale decomposition rates increased explained variance from 70% to 77% over abiotic factors alone (Wall et al., 2008). These examples indicate that the actors that play an important role in SOM dynamics should be considered in SOM models.

Model parameters are often measured in situ at relatively large spatial scales – at least compared with the size or activity range of most soil animals. As a result, the fauna effect is de facto included, although not appreciated (Swift et al., 1998). However, in many cases parameters are measured or extrapolated by combining in situ methods (e.g. monitoring of gas flux or litterbag experiments) and ex situ techniques such as laboratory experiments at controlled, highly simplified conditions. Especially the results of the latter may be sensitive to neglecting soil fauna. On the other hand, not taking explicitly into account the spatial heterogeneity created by soil fauna in field measurements might lead to substantial errors in calculating carbon budgets (Wu et al., 2013; Lopes de Gerenyy et al., 2015). It is thus crucial to develop sound (and biome-specific) strategies for combining in- and ex-situ measurements as parameters in more realistic SOM models.

Another fundamental issue in the large-scale approach is often neglected: When including the effects of the soil fauna implicitly, this assumes that the soil fauna will always have the same effects under the same conditions, and hence that the soil fauna is essentially static. This assumption is increasingly unrealistic in a fast-changing world where both biodiversity and the climate are changing at accelerated paces, and where we are likely to witness major reorganisations of plant, animal and microbial communities. Therefore explicit representation of the soil fauna, where possible, should increase the predictive ability of soil models.

6. Conclusions and outlook

Understanding and modelling SOM is essential for managing the greenhouse gas balance of the soil, for land restoration from desertification, for sustaining food production and for the conservation of above- and belowground biodiversity and associated ecosystem services (Nielsen et al., 2015). Soil animal abundance, biodiversity, species traits and interactions are crucial for SOM turnover (Chauvel et al., 1999; Bohlen et al., 2004; Wardle et al., 2004; Wall et al., 2008; Uvarov, 2009). In Table 2 we give recommendations how the known impact of soil fauna on SOM turnover could be used for improving carbon models. Due to the pronounced differences with respect to climate, soil and land use outlined above, it is important that these
recommendations are considered region-specific, taking into account the key players and their specific activities in the respective area.

Based on the arguments compiled here, a COST Action entitled “Soil fauna - Key to Soil Organic Matter Dynamics and Modelling (KEYSOM)” was established in March 2015 (http://www.cost.eu/COST_Actions/essem/ES1406). An interdisciplinary consortium of soil biologists and biogeochemists, experimenters and modellers from 21 European countries plus the Russian Federation and the USA cooperates to implement soil fauna in improved SOM models as a basis for sustainable soil management. A number of workshop will address key challenges in experimentation and modelling of SOM and soil fauna and support research exchange and access to experimental data. Special attention will be given to education of young scientists. The Action comprises four Working Groups with the following topics:

1. Knowledge gap analysis of SOM – soil fauna interactions;
2. Potentials and limitations for inclusion of soil fauna effects in SOM modelling;
3. Data assemblage and data sharing;
4. Knowledge management and advocacy training.

After an intensive and enthusiastic workshop held in Osijek, Croatia in October 2015, first activities included compilation of literature, the setup and permanent update of a website (http://keysom.eu/). Meanwhile short-term scientific missions for early-career scientists have been launched (http://keysom.eu/stsm/KEYSOM-STSMS-are-open-for-application). The second workshop was held in Prague in April 2016. One major outcome was the plan to conduct a common European-wide field study related to the overall objective.

**Author contribution**

J. Filser wrote the article, prepared Fig. 1 and the tables and compiled the contributions from all co-authors. These are listed according to their quantitative and qualitative impact on the manuscript, except for J.J. Jimenez who was placed last as he is the chair of KEYSOM. L. Brussaard suggested including Fig. 2.

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### Tables

#### Table 1. Quantitative examples of the impact of earthworms and selected groups of other soil fauna on soil properties and processes involved in soil organic matter (SOM) turnover

<table>
<thead>
<tr>
<th>Insight*</th>
<th>Examples</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Molecular structure</td>
<td>An indicator of lignin degradation in earthworm casts was twice that of the surrounding soil</td>
<td>Guggenberger et al., 1995</td>
</tr>
<tr>
<td>2. Humic substances</td>
<td>Introduced earthworms can double microaggregate formation and the stabilization of new C in the topsoil</td>
<td>Marashi and Scullion, 2003; Six et al., 2004</td>
</tr>
<tr>
<td></td>
<td>C protection is promoted by microaggregates within large macroaggregates, and earthworms can add 22% anew to this C pool</td>
<td>Bossuyt et al., 2005</td>
</tr>
<tr>
<td></td>
<td>Exclusion of earthworms reduced SOC accumulation by 0 (at 0-10 cm depth) to 75% (at 30-40 cm depth), associated with a decrease in percentage of water-stable aggregates</td>
<td>Albrecht et al., 2004, cited in Schmidt et al., 2011</td>
</tr>
<tr>
<td></td>
<td>In organic layers of a Canadian aspen forest, in locations with earthworms, N (1.5–0.8%) and especially C concentrations (25.3–9.8%) were strongly reduced, together with C/N ratio (16.7–13.2) and soil pH (6.5–6.1). This suggests a shift towards a faster cycling system, resulting in a net loss of C from the soil and turning Northern temperate forests from C sinks into C sources</td>
<td>Eisenhauer et al., 2007</td>
</tr>
<tr>
<td>3. Fire-derived carbon</td>
<td>Small charcoal particles from burned plots after one year increased by 21% in 0-1cm depth. One year later they were concentrated in earthworm casts at the soil surface, after 6.5 years such casts were found at 8 cm depth</td>
<td>Eckmeier et al., 2007</td>
</tr>
<tr>
<td>4. Roots</td>
<td>Presence of earthworms in a continuous maize plot in Peruvian Amazonia increased the organic C input from roots by 50%</td>
<td>Jiménez et al., 2006</td>
</tr>
<tr>
<td>5. Physical heterogeneity (see also insights no. 2, 3, 6 and 7)</td>
<td>Up to 50% of soil aggregates in the surface layer of temperate pastures are earthworm casts</td>
<td>Van de Westeringh, 1972</td>
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<td>Mull-type forest soil top layers and wooded savanna soils consist almost entirely of earthworm casts</td>
<td>Kubiena, 1953; Lavelle, 1978</td>
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<td>Earthworm inoculation in pastures on young polder soils completely removed within 8-10 years the organic surface layer, incorporated it into deeper layers, creating an A horizon. This affected manifold measures, increasing e.g. grass yield by 10%, root content in 0-15% from 0.38 to 1.31 g dm$^{-3}$, C content in 0-20 cm from 1.78 to 16.9 kg C $\times 10^{3}$ ha$^{-1}$, and water infiltration capacity from 0.039 to 4.6 m 24 h$^{-1}$. In turn, penetration resistance at 15 cm depth decreased from 35 to 22 kg cm$^{-2}$.</td>
<td>Hoogerkamp et al., 1983</td>
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<td></td>
<td>In average temperature pasture and grasslands, earthworms cast 40-50 t ha$^{-1}$ year$^{-1}$ on the surface and even more below surface</td>
<td>Lee, 1985</td>
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<td>Passage of a tropical soil through the gut of the invading earthworm <em>Pontoscolex corethrurus</em> reduced macroporosity from 21.7 to 1.6 cm$^{3}$ g$^{-1}$, which exceeded the effect of mechanically compacting the same soil at 10$^{3}$ kPa (resulting macroporosity: 3 cm$^{3}$g$^{-1}$)</td>
<td>Wilkinson et al., 2009</td>
</tr>
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<td>In a Canadian aspen forest a thick layer of earthworm cast material (thickness up to 4 cm) on top of organic layers was developed</td>
<td>Eisenhauer et al., 2007</td>
</tr>
<tr>
<td>6. Soil depth</td>
<td>Burrows of anecic earthworms are up to several meters deep and last for many years</td>
<td>Edwards and Bohlen, 1996</td>
</tr>
<tr>
<td>7. Permafrost and boreal areas</td>
<td>Earthworm invasions in boreal forests completely transformed mor to mull soils and significantly altered the entire plant community</td>
<td>Frellich et al., 2006</td>
</tr>
<tr>
<td>8. Soil microorganisms</td>
<td>Earthworms may lower actual microbial activity (by 11-23%) but markedly (by 13-19%) optimize microbial resource utilization.</td>
<td>Scueh et al., 2002</td>
</tr>
</tbody>
</table>

* According to Schmidt et al. (2011)
<table>
<thead>
<tr>
<th>Insight</th>
<th>Examples</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>2. Humic substances</td>
<td>In a degraded marsh in NE China, ant mounds were CH₄ sinks, contrary to the control soils which were CH₄ sources (-0.39 – -0.19 mg vs. 0.13 – 0.76 m² h⁻¹)</td>
<td>Wu et al., 2013</td>
</tr>
<tr>
<td>5. Physical heterogeneity</td>
<td>Ant and termite mounds can occupy up to 25% of the land surface</td>
<td>Bottinelli et al., 2015</td>
</tr>
<tr>
<td>5. Physical heterogeneity and</td>
<td>Underground nests of leafcutter ants (e.g. <em>Atta</em> spp.) can cover up to 250 m² and extend down to 8 m, which is associated with a massive impact on forest vegetation</td>
<td>Correa et al., 2010</td>
</tr>
<tr>
<td>6. soil depth</td>
<td></td>
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<tr>
<td>8. Soil microorganisms</td>
<td>Grazing by Collembola affected community composition of ectomycorrhizal fungi and on average reduced ¹⁴CO₂ efflux from their mycelia by 14%</td>
<td>Kanters et al., 2015</td>
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<td>Grazing by <em>Protaphorura armata</em> at natural densities on AM fungi disrupted carbon flow from plants to mycorrhiza and its surrounding soil by 32%</td>
<td>Johnson et al., 2005</td>
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<td>The presence of a single Collembola species may enhance microbial biomass by 56%</td>
<td>Fibser, 2002</td>
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<td>At elevated temperature, litter decay rates were up to 30% higher due to Collembola grazing</td>
<td>A’Bear et al., 2012</td>
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<td></td>
<td>Various or mixed groups</td>
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<tr>
<td>1. Molecular structure</td>
<td>Microbial grazing by Collembola or enchytraeids alone enhanced leaching of NH₄⁺ or DOC by up to 20%</td>
<td>Fibser, 2002</td>
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<td></td>
<td>Feeding by millipedes and snails reduced the content of condensed tannins in three Mediterranean litter species from 9–188 mg g⁻¹ dry matter to almost zero</td>
<td>Coulis et al., 2009</td>
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<td></td>
<td>Long-term mineralisation of fauna faeces may be slower than the mineralisation of litter from which the faeces were produced. This decrease in decomposition rate corresponds to a decrease in the C:N ratio and in the content of soluble phenols.</td>
<td>Frouz et al., 2015a,b</td>
</tr>
<tr>
<td>2. Humic substances</td>
<td>In a laboratory experiment, activity of earthworms, Collembola, enchytraeids and nematodes in coarse sand liberated &gt;40% from the insoluble C pool as compared to the control</td>
<td>Fox et al., 2006</td>
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<td>The quantitative contribution of invertebrates (mainly beetles and termites) to wood decomposition ranges between 10-20%</td>
<td>Ulyshen, 2014</td>
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<td></td>
<td>Depending on fungal and animal species (Collembola, isopods and nematodes), grazing on fungi colonising wood blocks altered (mostly decreased) their decay rates by more than 100%. Isopods and nematodes had opposite effects in this study.</td>
<td>Crowther et al., 2011</td>
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<td>Carbon and nitrogen losses from soil followed by drought and rewetting were substantially affected by microarthropod richness, which explained 42% of the residual variance.</td>
<td>De Vries et al., 2012</td>
</tr>
<tr>
<td>5. Physical heterogeneity</td>
<td>Bioturbation rates of soil animal groups typically range between 1 and 5 Mg ha⁻¹ y⁻¹ but may reach up to 10 (crayfish, termites), 20 (vertebrates), 50 (earthworms) and &gt; 100 Mg ha⁻¹ y⁻¹ (earthworms in some tropical sites), which is equivalent to maximum rates of tectonic uplift</td>
<td>Wilkinson et al., 2009</td>
</tr>
<tr>
<td>8. Soil microorganisms</td>
<td>In the course of a 2.5-yr succession, fauna activities (especially of nematodes and mesofauna during the first year, and later of earthworms) accelerated microbial decomposition of clover remains in an arable soil by 43%</td>
<td>Uvarov, 1987</td>
</tr>
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<td></td>
<td>Depending on vegetation, animal group and climate, soil animals directly or indirectly increased C mineralisation between 1% and 32%. However, intense grazing by fungal feeders may even reduce C mineralisation</td>
<td>Persson, 1989</td>
</tr>
</tbody>
</table>
Table 2. “Insights” (compiled after Schmidt et al., 2011) for future soil organic matter models and recommendations for further improvements by implementing effects of soil fauna

<table>
<thead>
<tr>
<th>SOM modelling element (“Insight”)</th>
<th>Recommendations*</th>
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<tbody>
<tr>
<td>1. Molecular structure</td>
<td>Incorporate the knowledge on the structure of organic substances and element concentrations in faunal casts and excreta in SOM decay rate models. Consider linkage between C and N cycling mediated by fauna. See 8.</td>
</tr>
<tr>
<td>2. Humic substances</td>
<td>Add physical and chemical stability of casts, patterns of their microbial colonisation and degradation dynamics. See 1, 3, 5, 6, 7, 8.</td>
</tr>
<tr>
<td>4. Roots</td>
<td>Add activity of bioturbators, rhizosphere microbial grazers and root herbivores. See 1, 5, 6, 8.</td>
</tr>
<tr>
<td>5. Physical heterogeneity</td>
<td>Consider spatial and physicochemical heterogeneity created by soil fauna, including consequences of soil aggregation and dis-aggregation (e.g. bulk density, infiltration rate, preferential flow, casts). See 1, 2, 6, 8.</td>
</tr>
<tr>
<td>6. Soil depth</td>
<td>Incorporate burrowing depth and annual transport rates of bioturbators and animal-induced spatial heterogeneity of old and young carbon in the deep soil. See 5.</td>
</tr>
<tr>
<td>7. Permafrost</td>
<td>For warming scenarios, take into account short- and long-term invasion effects, particularly of earthworms and enchytraeids.</td>
</tr>
<tr>
<td>8. Soil microorganisms</td>
<td>Add microbial grazer effects, effects on microorganisms during gut passage and faunal impact on C and N coupling. See 1-7.</td>
</tr>
</tbody>
</table>

* Recommendations refer to site-specific keystone groups of animals (dominating in terms of biomass or impact; see Fig. 2). Their prevalence is determined by climate, bedrock and land use (e.g. rodents or ants in deserts, earthworms in temperate grasslands or microarthropods and enchytraeids in acidic Northern forests).
Figure Captions

Figure 1. Main animal-mediated processes (boxes) affecting the eight insights (symbols) identified by Schmidt et al. (2011) that should be considered for improving SOM models.

Figure 2. Dominant soil types and characteristic soil forming invertebrates across biomes (major global change threats are shown in italics). MAT = mean annual temperature, MAP = mean annual precipitation. Sources for data and biomes see Brussaard et al. (2012).

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Figures

Figure 1
Figure 2