Heterotrophic and Autotrophic Soil Respiration under Simulated Dormancy Conditions

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Abstract

Carbon cycling research has increased over the past 20 years, but less is known about the primary contributors to soil respiration (i.e. heterotrophic and autotrophic) under dormant conditions. It is understood that soil CO₂ effluxes are significantly lower during the winter of temperate ecosystems and assumed microorganisms dominate efflux origination. We hypothesized that heterotrophic contributions would be greater than autotrophic under simulated dormancy conditions. To test this hypothesis, we designed an experiment with the following treatments: combined autotrophic heterotrophic respiration, heterotrophic respiration, autotrophic respiration, no respiration, autotrophic respiration in vermiculite, and no respiration in vermiculite. Engelmann spruce seedlings and soil substrates were placed in specially designed respiration chambers and soil CO₂ efflux measurements were taken four times over the course of a month. Soil microbial densities and root volumes were measured for each chamber after day thirty-three. Seedling presence resulted in significantly higher soil CO₂ efflux rates for all soil substrates. Autotrophic respiration treatments were not representative of solely autotrophic soil CO₂ efflux due to soil microbial contamination of autoclaved soil substrates; however, the mean autotrophic contributions averaged less than 25% of the total soil CO₂ efflux. Soil microorganism communities were likely the primary contributor to soil CO₂ efflux in simulated dormant conditions, as treatments with the greatest proportions of microbial densities had the highest soil CO₂ efflux rates. Although this study is not directly comparable to field dormant season soil CO₂ effluxes of Engelmann spruce forest, as snowpack is not maintained throughout this experiment, relationships, and metrics from such small-scale ecosystem component processes may yield more accurate carbon budget models.

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Keywords
Dormant Autotrophic Respiration, Heterotrophic Respiration, Dormant Soil CO2 Efflux, Soil Respiration, Engelmann Spruce

1. Introduction

Soil CO2 efflux is the primary carbon efflux from terrestrial ecosystems to the atmosphere; soils and plant biomass represent the largest terrestrial carbon pools storing more than 1000 petagrams (Pg) of carbon (Bradford, Birdsey, Joyce, & Ryan, 2008; Dixon et al., 1994; Raich & Schlesinger, 1992). Soil CO2 efflux is composed of autotrophic (plant) and heterotrophic (microbial) metabolic processes, and is usually quantified at the soil surface as a single source flux (Amiro et al., 2010). Previous studies have attempted to separate autotrophic and heterotrophic fluxes, primarily utilizing three methods (i.e. root exclusion, component integration, and isotopic experiments) (Hanson, Edwards, Garten, & Andrews, 2000; Lee, Nakane, Nakatsubo, & Koizumi, 2003); however, many of these studies have limitations, as physically separating autotrophic and heterotrophic soil CO2 efflux contributions may influence available soil moisture and gas diffusion rates. Isotopic methods are a good tool for isolating heterotrophic and autotrophic respiration but are difficult to setup in situ and very costly (Coleman, 1991; Hanson et al., 2000). Thus, heterotrophic and autotrophic carbon efflux contributions to the gross soil carbon efflux are not well understood. Further, soil CO2 efflux of dormant, snow-covered forest ecosystem is a lesser studied aspect of the carbon cycle and less commonly accounted for in carbon budgets (Houghton, Hackler, & Lawrence, 1999). Here, we attempt to separate autotrophic and heterotrophic soil CO2 efflux under simulated dormant season conditions.

Accurate quantification of carbon fluxes is necessary to construct carbon budget models and accurately estimate ecosystem productivity. Soil CO2 effluxes, as most ecosystem processes, are sensitive to seasonal and environmental change; as climates and disturbance regimes become less predictable, atmospheric carbon efflux exponentially increases driving earlier snow melt further perpetuating climate variability (Buchmann, 2000; Davidson & Janssens, 2006; Friedlingstein et al., 2001; Stewart, Cayan, & Dettinger, 2004). Studies during the last decade have shown that temperate ecosystems experiencing snow cover have considerably lower soil CO2 effluxes compared to growing season fluxes (Brooks, Williams, & Schmidt, 1996; Hubbard, Ryan, Elder, & Rhoades, 2005). Because subalpine ecosystems are typically covered with snow for longer time periods than their associated growing seasons, winter soil CO2 effluxes can compose a substantial portion of the subalpine annual carbon budget. For example, observations approximate 8% - 15% of the annual soil CO2 efflux in subalpine forest occurs during the dormant season (Beverly, 2013; Hubbard et al. 2005). Recently, carbon cycle models have started to incorporate the contributions of winter soil CO2 efflux under snowpack (Brooks, McKnight, & Elder, 2005), but more data and detailed knowledge from multiple ecosystems are needed.

Autotrophic respiration during the growing season results from active transport of nutrients, growth of new biomass, tissue maintenance, and exchange of allocated carbon for nitrogen between plants and heterotrophic soil microbes (Bowden, Nadelhoffer, Boone, Melillo, & Garrison, 1993; Ryan, 1991). However, less is known about soil CO2 effluxes while many of these autotrophic processes are down-regulated during dormancy. The correlations of soil CO2 efflux with soil temperature and moisture during the growing season are well understood, making them valuable predictors for climate and ecosystem productivity models. However, for unfrozen, dormant soils, conditions are relatively stable under snowpack (Iwata, Hayashi, & Hirota, 2008; Ryan, 1991; Sutinen, Holappa, Ritari, & Kujala, 1999); e.g., once snowpack insulates soil, soil temperature remains approximately 0°C as depth of snowpack increases (Monson et al., 2006). Because of the stability of the environment under snowpack compared to the growing season, soil CO2 efflux contributions from autotrophic and heterotrophic sources may have less predictive covariables in the winter.

Separating autotrophic and heterotrophic contributions could help parameterize carbon cycle models to more accurately predict carbon fluxes under changing conditions, especially if plants and microorganisms respond differently to the changes (e.g., soil warming). However, such separation is tedious and difficult (Anderson & Domsch, 1973; Bowden, Nadelhoffer, Boone, Melillo, & Garrison, 1993; Hanson et al., 2000). Soil CO2 efflux quantified at the soil surface is limited to inferences on the complex microbial and root interactions occurring...
within the rhizosphere (Johnson & Jost, 2011). The majority of data is collected from field-based experiments making accurate quantification of soil CO₂ efflux contributors difficult (Hanson et al., 2000), especially with mycorrhizal symbiotic species (Paterson et al., 1997; Rouhier, Billès, Billès, & Bottner, 1996). Bowden et al. (1993), using root and litter exclusion methods, estimated that two-thirds of summer soil respiration was contributed by root respiration; microbial soil efflux only contributed one-third from growth and decomposition.

Three techniques for quantifying autotrophic and heterotrophic contributions dominate the literature; component integration (i.e. gross flux is derived through addition of isolated contributors) (Garrett & Cox, 1973; Johnson et al., 1994), root exclusion (Bowden, Nadelhoffer, Boone, Melillo, & Garrison, 1993; Comstedt, Boström, & Ekblad, 2011; Ewel, Cropper, Jr., & Gholz, 1987), and stable and isotopic labeling (Andrews, Harrison, Matamala, & Schlesinger, 1999; Biasi, Pitkämäki, Tavi, Koponen, & Martikainen, 2012; Gomez-Casanovas, Matamala, Cook, & Gonzalez-Meler, 2012) experiments. Isotopic methods require extensive sampling protocols, expensive laboratory equipment, and invasive sampling developing a below ground reference gas concentration for computing flux rates. Component integration and root exclusion methods are substantially cheaper and have been utilized in a variety of ecosystems and vegetation types; however, overestimation of soil CO₂ efflux rates can occur due to soil disturbance increasing soil CO₂ flushing (Hanson et al. 2000); thus, studies using these methods should be applied in experiments lasting over a week allowing for efflux rates to equilibrate. Many component integration experiments do not attempt to physically separate autotrophic and heterotrophic contributors (e.g., soil and root sterilization); rather, experiments infer that rhizosphere respiration and autotrophic respiration are not different, though heterotrophic contributions are known to be present.

In this study, we attempted to quantify soil CO₂ efflux contributions of dormant Engelmann Spruce (Picea engelmannii Parry ex Engelm.) seedlings and soil microorganisms in dormant simulated conditions. We hypothesized that reduced autotrophic activity of dormant seedlings results in heterotrophic respiration composing the majority of the dormant soil CO₂ efflux. To our knowledge, this is the first study trying to isolate dormant soil CO₂ efflux with a controlled lab approach.

2. Material and Methods

2.1. Study Design

A chamber respiration experiment was developed using root exclusion (i.e. seedling and no seedling) and component integration (i.e. soil microorganisms and no soil microorganisms) methods for three soil types. A two factor (i.e. with and without seedling; and soil: autoclaved, non-autoclaved, and autoclaved vermiculite) study was designed for quantifying contributions of soil CO₂ efflux. Factorial treatments included five replications of each treatment: autotrophic and heterotrophic respiration (AHR), autotrophic respiration (AR), heterotrophic respiration (HR), no organism respiration in soil (NR), autotrophic respiration in vermiculite (VAR), and no organism respiration in vermiculite (VNR). Vermiculite was used as an additional control (i.e. negative control) to remove soil effects on soil CO₂ efflux. Seedlings were obtained from the Colorado State Forest Service Nursery (Fort Collins, CO), seedlings were grown outdoors for six months where they set bud and entered dormancy. The six treatments, of thirty chambers (15 with seedlings), were brought into the laboratory one month prior to first measurement while being stored in a cold room at 4°C simulating seedling dormancy and reducing metabolic activity of soil microbes. Grow lights in the cold room provided photosynthetic active radiation (PAR) of 86 - 102 µmol m⁻² s⁻¹ with an 8:16 light to dark ratio mimicking winter day length. Soils for this study were collected in April 2014 from P. engelmannii stands in the Poudre Canyon approximately 60 kilometers west of Fort Collins, Colorado (N 40.637983, W 105.806136). The field site was mixed forest of P. engelmannii and Abies lasiocarpa. Snow was excavated to reach the soil surface where the top 20 cm of soil were collected from a 2 × 3 m plot. Soils were sifted with a 2 mm screen producing a uniform soil substrate composed of fine sands and silts. Prior to filling soil chambers, vermiculite was thoroughly mixed with sifted soils in a 3:2 vermiculite to soil ratio to reduce soil compaction in the chambers. Soil and vermiculite substrates to be used in treatments with no heterotrophic respiration were placed in stainless steel autoclavable trays approximately 5 cm deep, and autoclaved three consecutive times prior to filling the soil respiration chambers (Trevors, 1996; Wolf, Dao, Scott, & Lavy, 1989).

2.2. Respiration Chambers

Thirty respiration chambers were constructed from 5.08 × 28 cm polyvinylchloride (PVC) plumbing pipe fitted
with a perforated end cap (allowing for water drainage) that was sealed prior to measuring soil CO$_2$ effluxes. Respiration chambers with seedlings had small openings (1 cm$^2$) on the rim of the chamber to divert the seedling stems, and separate roots and crown mass by the chamber wall; small openings were sealed with silicone post-planting of the seedling. Respiration measurements were collected through the 81.0 cm$^2$ opening at the top of the chamber.

Soil chambers were sterilized with 70% ethanol and then filled with 2050 cc of autoclaved soil substrate, non-autoclaved soil substrate, or autoclaved vermiculite. Seedling roots were removed of residual soil and washed; seedling root volume was measured by volume displacement in a graduated cylinder. Prior to planting, seedling roots were rinsed, surface sterilized with 70% ethanol for 30 seconds (protocol from macropropagation; Bhojwani and Razdan, 1986) then rinsed again with ultra-pure water (Milli-Q Millipore water; EMD Millipore Inc.). At the end of the experiment, seedlings were removed from chamber, cleaned, and measured for final root volumes.

Simulated winter conditions maintained an ambient air temperature of 4°C, but initial soil moistures varied for the three soil types. Thus, chambers were watered with autoclaved water increasing soil moisture levels to field holding capacity. To ensure that seedlings were dormant, concurrent foliar gas exchange measurements were conducted on all fifteen seedlings during three of the four measurement days using a second LI6400. Measurements were conducted using the LI6400 equipped with the conifer photosynthesis chamber (Li-Cor 6400-22L); measurement conditions included PAR at 1000 µmol$\cdot$m$^{-2}\cdot$s$^{-1}$ and reference CO$_2$ at 400 µmol$\cdot$mol$^{-1}$.

### 2.3. Respiration Measurements

Soil CO$_2$ efflux was quantified using the Li-Cor 6400 infrared gas analyzer (LI6400) (Li-Cor Inc., Lincoln NE) with a soil respiration chamber (Li-Cor 6400-09). During measurements, chambers were randomly removed from the cold room. Measurements for each chamber were conducted during the afternoon with three measurement cycles for each day of measurement; the three cycles for each chamber were averaged producing a soil CO$_2$ efflux for each day for each chamber. Soil CO$_2$ efflux measurements occurred two, five, ten and thirty-three days post experiment start. For analysis, soil CO$_2$ effluxes were averaged over the four days as we did not expect, nor see, a day effect.

Soil temperature and moisture measurements were conducted with every soil CO$_2$ efflux measurement. Soil temperatures were measured with a mercury thermometer inserted 12 cm into the soil column for 30 seconds. Soil moisture content was measured using HS2 Hydrosense TDR system (Campbell Scientific Inc., Logan, Utah) that averaged soil moistures from 12 cm to 20 cm in the soil column.

Soil CO$_2$ effluxes were natural log transformed to correct for non-normality of the data. Data analysis was conducted using a forward step-wise ANOVA model with covariables of root volume, photosynthetic rate, soil moisture and temperature, bacterial and fungal densities. A two factor model, crossing of factors with autoclaved, non-autoclaved soils, and vermiculite (soil factor) with seedling or no seedling (seedling factor), was used to determine interaction effects of the treatments. Least square means with a Tukey adjustment was used as a post-hoc analysis to evaluate differences among treatments. Five replications of each treatment had an $a$ posteriori ANOVA power of 0.839.

Because root volume was a significant covariable in analysis, autotrophic soil CO$_2$ effluxes were adjusted using root volumes for each treatment measured on day thirty-three. Adjusted soil CO$_2$ effluxes were used to determine autotrophic contributions for each treatment.

### 2.4. Microbial Quantification

After day thirty-three, soil samples from each chamber were subjected to an eight-fold serial dilution to evaluate effectiveness of microorganism reduction within the soil substrates. The first dilution was derived from 50 g of soil suspended in 450 mL purified water. Higher dilutions were derived by adding 1 mL of the previous dilution to 9 mL purified water in a sterile 15 mL tube. Dilutions were thoroughly mixed with a vortex mixer before inoculating the next higher dilution. Each replicate used 100 µL aliquots and a cell spreader was used to distribute aliquots evenly across selective mediums. Selective media plates were used to differentiate between fungal and bacterial soil fauna. Czapek-Dox (CZA) agar plates were used to select against soil bacteria. Selection against fungal contributors was completed using nutrient agar with 0.1 µL Amphotericin B. Aliquot plates for each 7th and 8th fold dilutions were replicated for each soil sample (6 nutrient and CZA agar
plates per chamber). All treatments were incubated at 26°C for five days. Colony-forming units (CFU) were counted for both bacterial and fungal plates as an estimation of microbial density for each soil chamber (Saxena & Stotzky, 2001). CFU counts between 3 and 300 per plate were used for analysis.

The six microbial CFU estimates (three fungal and three bacterial) for each chamber were averaged for both bacterial and fungal densities prior to analysis. Bacterial and fungal densities were used to adjust soil CO₂ efflux rates to determine overall heterotrophic contributions in each soil substrate. Adjustment equations for each soil treatment (microorganism soil efflux without seedlings/total microorganisms without seedlings = x/total microorganisms with seedlings) were developed to adjust soil CO₂ efflux contributions.

3. Results

3.1. Foliar Gas Exchange

Foliar gas exchange measurements averaged over the three measurement days for AHR and AR suggested respiration, respectively $-2.26 \pm 1.94$ and $-1.22 \pm 1.17$ µmol CO₂ m⁻² s⁻¹ (Figure 1). Seedlings in vermiculite soil substrates had very low photosynthetic gas exchange rates averaging $0.58 \pm 2.44$ µmol CO₂ m⁻² s⁻¹ (Figure 2). The high variability in foliar gas exchange resulted in no significant differences of the photosynthetic rates among the three soil substrates.

3.2. Soil Respiration

Soil CO₂ efflux showed a significant interaction between seedling and soil substrate (Table 1). Root volume was the only significant covariable in the forward step-wise ANCOVA model (R-square = 0.32; p-value = 0.029); soil moisture and temperature, photosynthetic rate, and microbial densities were not significant covariables. After accounting for covariables, treatments containing a seedling had significantly higher soil CO₂ efflux rates than without a seedling for all three soil types, but the proportions of autotrophic contribution to the total soil CO₂ efflux was less compared to the heterotrophic contribution in autoclaved soil (8.9%, p-value = <0.0001), non-autoclaved soil (20.1%, p-value = 0.0246), and vermiculite soil substrate (40.1%, p-value = 0.0036) (Figure 2 and Figure 3). The greatest soil CO₂ efflux rates were in the autoclaved soil substrate with an autotrophic con-
Table 1. Analysis of covariance examining effects of soil substrates (autoclaved, non-autoclaved, and vermiculite) and seedling (presence, absent) on soil CO₂ efflux in respiration chambers under simulated dormant (winter) conditions. Covariables included root volume, photosynthetic rate, soil moisture and temperature, bacterial and fungal densities.

<table>
<thead>
<tr>
<th>Source</th>
<th>DF</th>
<th>Sum of Squares</th>
<th>Mean Square</th>
<th>Expected Mean Square</th>
<th>Error Term</th>
<th>Error DF</th>
<th>F Value</th>
<th>p Value</th>
</tr>
</thead>
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<tr>
<td>Tree</td>
<td>1</td>
<td>0.221149</td>
<td>0.221149</td>
<td>Var (Residual) + Q (Tree, Tree*Soil Substrate)</td>
<td>Ms (Residual)</td>
<td>23</td>
<td>5.75</td>
<td>0.025</td>
</tr>
<tr>
<td>Soil substrate</td>
<td>2</td>
<td>3.518912</td>
<td>1.759456</td>
<td>Var (Residual) + Q (Soil Substrate, Tree*Soil Substrate)</td>
<td>Ms (Residual)</td>
<td>23</td>
<td>45.76</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Tree*Soil Substrate</td>
<td>2</td>
<td>1.146466</td>
<td>0.573083</td>
<td>Var (Residual) + Q (Tree*Soil Substrate)</td>
<td>Ms (Residual)</td>
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<td>14.9</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Residual</td>
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<td>0.884356</td>
<td>0.03845</td>
<td>Var (Residual)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Figure 2. Soil CO₂ effluxes were significantly higher with the presence of dormant seedlings independent of soil substrate. No respiration (Hobson et al.) and vermiculite no respiration (VNR) had significantly larger soil CO₂ effluxes due to soil microbial contamination.

Seedling root volumes of the autoclaved soil substrate treatments were significantly larger than vermiculite and non-autoclaved soil seedlings; 28.3% and 48.3% larger, respectively (t-value = -2.47, p-value = 0.0296) (Figure 4). Vermiculite and non-autoclaved soil substrate seedlings were not significantly different (t-value = -1.02, p-value = 0.3287). Root volumes consisted of less than 2% of the total chamber volume; mean root volume for all soil substrates was 1.4% of total chamber volume.

3.3. Microbial Density

The result of this study lasting over a month was that all treatments contained soil microorganisms; autoclaved soil substrates resulted in rapid microorganism colonization with ample nutrient availability and the absence of competitive controls for growth. Autotrophic respiration (AR) and autoclaved no respiration (NR) treatments...
Figure 3. Adjusted soil CO$_2$ efflux contributions for non-autoclaved soil (HR and AHR), autoclaved soil (NR and AR), and vermiculite soil substrates. Autotrophic contributions (black) were weighted on root volumes for AHR, AR, and VAR, contributing 20.1%, 8.9%, and 40.1%, respectively.

Figure 4. Root volume was the only significant variable in the global ANCOVA for predicting soil CO$_2$ efflux. Root volumes were significantly larger in the autoclaved soil substrate than non-autoclaved soil substrate; root volumes in the vermiculite soil substrate were not significantly different from either soil substrate.
had significantly greater densities of soil fungal growth than other treatments ($F = 4.17, p\text{-value} = 0.0337$) (Figure 5). All soil substrates had substantial bacterial growth, albeit not significantly different among treatments ($F = 0.15, p\text{-value} = 0.8621$) (Figure 6). The greatest proportions of total soil microbial densities were in the soil chambers that had autoclaved soils and vermiculite; autotrophic treatments (AR) contained 37% of all soil microbial biomass. Separating soil microbial groups (bacteria and fungi) shows the same trend with the autoclaved treatments containing the greatest proportions of fungal and bacterial colonization (Figure 6). Linear regression models predicting soil CO$_2$ efflux from proportions of total soil microbial and fungal densities for all six treatments explained greater than 70% of the variability of soil CO$_2$ effluxes; $R^2$ values of 0.717 and 0.860, respectively; the model for bacterial proportions explained less of the variability with an $R^2$ value of 0.362.

4. Discussion

Contributors to soil CO$_2$ effluxes remain poorly understood due to the difficulties of separating autotrophic and heterotrophic components; and very little is known about these contributors during the dormant season under snowpack. Temperate and subalpine forest soils are not completely frozen as snowpack insulates soil allowing for microbial and root activity (Brooks & Williams, 1999; Hubbard et al., 2005). Winter soil CO$_2$ efflux contributions have been estimated to be 8% - 15% of the annual soil respiration for subalpine forests, though the organismal contribution is unknown; thus, carbon fluxes during winter conditions cannot be ignored and need to be incorporated into annual carbon budgets (Hubbard et al., 2005; Beverly, 2013), as has been shown by Wang et al. (2013).

Autotrophic respiration is the primary contributor to growing season soil CO$_2$ efflux, accounting for two-thirds of the gross soil CO$_2$ efflux (Bowden, Nadelhoffer, Boone, Melillo, & Garrison, 1993), but is variable and ranges between 10% - 90% depending on ecosystem (Hanson et al., 2000). During our simulated dormancy, autotrophic respiration accounted for 10% - 40% (mean = 23.0%) of the total soil CO$_2$ efflux, supporting our hypothesis that soil CO$_2$ efflux in simulated dormant conditions results primarily from microbial activity (Figure 3). Engelmann spruce, as many other conifer species, are known to rapidly exit dormancy under ideal conditions leading to increased autotrophic soil CO$_2$ efflux rates (Borken, Xu, Davidson, & Beese, 2002); however, spruce species typically respond to increases in both light and air temperature (Granhus, FlØistad, & SØgaard, 2009). There was potential for rapid release from dormancy during the soil CO$_2$ efflux measurements that occurred out-
Figure 6. Soil CO₂ effluxes increased with treatments containing the greatest proportions of fungi (a) and bacteria (b); R² values of 0.86 and 0.362, respectively. Autotrophic respiration (AR) had the greatest levels of soil microbial contamination resulting in the highest soil CO₂ effluxes.

side of the cold room, but our results suggest that the duration of soil CO₂ efflux measurements was too short to initiate release from dormancy. Foliar gas exchange rates suggested seedlings were dormant and carbon allocation did not occur; thus, autotrophic soil CO₂ effluxes were likely a result of minimal root biomass maintenance rather than new growth or nutrient acquisition. Indeed, root biomass actually decreased over the study period (data not shown) suggesting stored carbohydrates were being catabolized. The autotrophic contribution was substantial, even with low soil CO₂ efflux rates during dormant conditions, and was positively related to root biomass (R² = 0.32) suggesting a potential metric for carbon flux models.
Microorganism contamination of respiration chambers deserves further discussion for both the insight it gives to carbon fluxes and lessons it provides for such chamber experiments. Autoclave sterilization of soils can alter physical soil conditions, potentially affecting soil CO₂ efflux (Berns et al., 2008); however, both autoclaved and non-autoclaved Douglas-fir soils from a chamber experiment similar to the present one had no significant differences in soil pH or concentrations of organic matter (Beverly, 2013). Contrarily, mineral nutrients were significantly higher in the autoclaved soils than the non-autoclaved soils; this pulse of nutrient content released into the soils was likely from microbial destruction (Alef & Nannipieri, 1995), creating a priming effect that induced rapid fungal and bacterial growth, analogous to nutrient addition experiments that show an increase of soil CO₂ efflux rates between 44% and 169% (Nottingham, Griffiths, Chamberlain, Stott, & Tanner, 2009). Thus, autoclaving either did not fully sterilize soils or microbial contamination resulted following sterilization, so future studies need to sample soil microbial densities throughout the experiment. Substantial fungal contamination was a possibility due to known mycorrhizae associations with seedlings. Microbial contamination may also have occurred from the cold room ventilation systems over the course of the month during the experiment; however, contamination source needs further investigation.

We argue that the elevated soil CO₂ effluxes of autotrophic only and no organism respiration treatments resulted from high levels of microorganism densities within all treatments, especially fungi. While it is impossible to determine the source of contamination, several species of mycorrhizal fungi are known to contaminate container-grown Engelmann spruce (Hunt, 1989), and substantial mycorrhizae were observed in all chambers. These results match with field data that showed peak fungal biomass was three times greater in the winter under snowpack compared to the summer growing season (Schadt, Martin, Lipson, & Schmidt, 2003). Indeed, samples with the highest proportions of soil microbial contamination, primarily fungal contamination, yielded the largest soil CO₂ efflux rates. This partially explains why the autoclaved soil substrate with seedling (AR) chambers had a heterotrophic contribution that was greater than 90% of the soil CO₂ efflux (Figure 6).

While soil microorganism densities were substantially larger in the autoclaved soil substrates, densities were within known natural variation of soil microorganisms densities of pine forests (mean = 2.5 × 10⁷ CFU g⁻¹ of soil; (Moore; 1983, Vazquez et al., 1993). Fungal densities are likely the main contributor to soil CO₂ effluxes under simulated dormant conditions, as fungal densities in this study were two-fold greater than natural pine forests likely increasing soil CO₂ effluxes (Vázquez, Acea, & Carballas, 1993). However, further understanding of soil microbial population dynamics of subalpine forest during dormant season is needed.

5. Conclusion

Our data suggest that fungal and bacterial soil communities are the major contributors to dormant season (simulated winter conditions) soil CO₂ efflux, contributing approximately 75% of the total carbon efflux. This relationship is the direct inverse of summer estimates (Bowden, Nadelhoffer, Boone, Melillo, & Garrison, 1993). Soil CO₂ efflux studies from temperate ecosystems that experience extended durations of snow cover likely underestimate winter carbon budgets for several reasons. First, many long-term sampling (e.g. eddy covariance) strategies have greater error in measurements during the winter seasons (Goulden, Munger, FAN, Daube, & Wofsy, 1996). This variability can result in a 10% - 30% underestimation of energy budgets, decreasing accuracy of measurements in long-term carbon budgets (Twine et al., 2000). Second, winter soil CO₂ efflux is not well studied, so the knowledge of autotrophic versus heterotrophic contribution is lacking. With some uncertainties of current ecosystem models, lab and field chamber studies might be incorporated to strengthen future ecosystem flux models. For example, dormant soil CO₂ efflux was positively correlated with seedling root volume (R² = 0.32), and such metrics may improve carbon models. In addition, seedlings significantly increased soil CO₂ efflux of all three soil substrates making up the other c. 25% of winter soil respiration. While this study confirms dormant season estimates need to be incorporated into carbon budgets and ecosystem models, field comparisons are needed to develop dormant autotrophic estimates at the forest stand level. As a final note, this study suggests that autoclaving soil does not necessarily result in sterilization throughout a study and microbial densities should accompany such studies.

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