Autoclaving is at least as effective as gamma irradiation for biotic clearing and intentional microbial recolonization of soil

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**Abstract**

Sterilization is commonly used to remove or reduce the biotic constraints of a soil to allow recolonization by soil-dwelling organisms, with autoclaving and gamma-irradiation being the most frequently used approaches. Many studies have characterized sterilization impacts on soil physicochemical properties, with gamma-irradiation often described as the preferred approach, despite the lower cost and higher scalability of autoclaving. However, few studies have compared how sterilization techniques impact soil recolonization by microorganisms. Here, we compared how two sterilization approaches (autoclaving; gamma-irradiation) and soil washing impacted microbial recolonization of soil from a diverse soil inoculum. Sterilization method had little impact on microbial alpha diversity across recolonized soils. For sterile soil regrowth microcosms, species richness and diversity were significantly reduced by autoclaving relative to gamma-irradiation, particularly for fungi. There was no impact of sterilization method on bacterial composition in recolonized soils and minimal impact on fungal composition (p = 0.05). Washing soils had a greater impact on microbial composition than sterilization method and sterile soil regrowth had negligible impacts on microbial recolonization. These data suggest that sterilization method has no clear impact on microbial recolonization, at least across the soils tested, indicating that soil autoclaving is an appropriate and economical approach for biotically-clearing soils.

**Importance**

Sterilized soils represent soil-like environments that act as a medium to study microbial colonization dynamics in more “natural” settings relative to artificial culturing environments. Soil sterilization is often carried out by gamma-irradiation or autoclaving, which both alter soil properties, but gamma-irradiation is thought to be the gentler technique. Gamma-irradiation can be cost prohibitive and does not scale well for larger experiments. We sought to examine how soil sterilization technique can impact microbial colonization, and additionally looked at the impact of soil washing which is believed to remove soil toxins that inhibit soil recolonization. We found that both gamma-irradiated and autoclaved soils showed similar colonization patterns when reintroducing microorganisms. Soil washing, relative to sterilization technique, had a greater impact on which microorganisms were able to recolonize the soil. When allowing sterilized soils to regrow (i.e. persisting microorganisms), gamma-irradiation performed worse suggesting that gamma-irradiation does not biotically-clear soils as well as autoclaving. These data suggest that both sterilization techniques are comparable, and that autoclaving may be more effective at biotically-clearing soil.

**Introduction**

Traditional culture-based approaches have a long history in microbiology and remain essential to microbiological research (1). Such approaches are ideal for certain research questions (e.g., physiological probing of easily captured species), but are limited in their ability to model complex microbiomes under realistic environmental conditions. In addition, most microbes remain difficult to cultivate, restricting in-culture studies to easily isolated microbes, which may be poor analogues for many of the important microbial players in complex environments, like soil (2-5).

To circumvent these limitations, soils are often “sterilized” to provide biotically-cleared environments that can be reinoculated with specific microbes or intact microbiomes (6-11). This type of approach is controllable and reproducible; for instance, we can see strong differentiation of introduced microbiomes (6, 12-14). It is also a more realistic model of an in-field soil compared to culture media, both structurally and chemically. In fact, widespread agricultural practices use biotic clearing of soil to eliminate pathogens, so in-lab “sterilization” of soil is an excellent analogue for such cases (12, 15, 16). However, the best approach for biotic clearing of soil has been debated for decades, since any sterilization method alters soil characteristics to some degree (17). As a result, biotically-cleared soils should not be treated as perfect replicates of in-field soils, but as soil-like environments that are similar to, but not equivalent to, the pre-sterilized soil. Numerous approaches, including airdrying, oven drying, addition of chemicals or antibiotics, fumigation, microwaving, gamma-irradiation, and autoclaving have been repeatedly tested as methods that can maximize sterilization while minimizing impacts on soil physicochemical characteristics (18).

Two common soil sterilization procedures are gamma irradiation and autoclaving (for long periods of time and/or multiple times over several days). Gamma irradiation requires highly specialized facilities and equipment, but has been described by some as a “gold standard”, as it is thought to cause fewer substantial changes to some soils during sterilization (17, 19, 20). Nonetheless, gamma-irradiation is not well borne out across a wide range of soil types (21) and may not biotically-clear soils better than autoclaving (19). Similar to other sterilization methods, gamma irradiation alters the physical and chemical properties of soil, including the release of highly reactive free radicals and changes to the structure of organic matter, among others (17, 22). Although less irradiation time and intensity can reduce the impact on soil properties, it may also be less effective in removing soil organisms, thus defeating its intended purpose. Autoclaving also alters a variety of soil physicochemical properties (17), but offers benefits that cannot be achieved with irradiation, such as ease of access and low financial burden, facilitating larger-scale soil processing to run experiments at scale (6, 7, 12-14, 17, 23, 24). Autoclaving closely mirrors large-scale soil management practices, such as soil steaming, which are used in agricultural settings to remove pathogens from the soil. Although numerous studies have investigated how each sterilization approach impacts soil structure and chemistry (17, 18), it is essential to understand the relative impacts of sterilization approaches on the growth of introduced microorganisms.

In this study, we aimed to determine the impacts of common sterilization approaches on the potential of soil microbiomes to colonize, establish, and persist within biotically-cleared soils. In addition to the two most common approaches for soil sterilization (i.e. gamma-irradiation and autoclaving), we included a washing step, as it has been suggested this could reduce the impact of released compounds or spores remaining in the soil after sterilization (25-32). Microbiomes from the pre-sterilized versions of two soils were introduced to assess the impact of each sterilization technique had on the ability of microbes to colonize and persist within the biotically-cleared soil environments. Based on prior in-lab observations, we hypothesized that: (i) soil type would have a greater impact on resulting microbial composition than sterilization method, and that (ii) autoclaved soils would have less diverse microbial “contaminants” (regrowth), yielding more reproducible microbial composition relative to gamma irradiated soil samples.

**Materials and Methods**

**Soil collection and substrate conditions**

Two physicochemically distinct soils within a standard agricultural pH range were collected from The Pennsylvania State University Russell E. Larson Agricultural Research Center at Rock Springs (Supplementary Table 1). Soil 1 was collected from an unmanaged, forested area, while Soil 2 was collected from a plot previously used for field trials of agricultural crops. Debris was cleared from the ground and a shovel was used to collect soil from the top 20 cm. Samples were sieved to 2 mm using a standard brass sieve and mixed 2:1 (v/v) with Quikrete All-Purpose Silica Sand to improve aeration and drainage. Between each stage (i.e. collection, sieving, sand addition, sterilization, and microcosm construction) soils were stored in sealed bags at 2°C. A fraction of each soil was left unmixed for use as negative controls

**Autoclave-based sterilization**

A portion of each prepared substrate was placed into standard biohazard bags (double-bagged) that are permissive to autoclaving and a soil depth of 2-3 cm. Each soil was autoclaved for 45 min at 121°C and 20 psi, with this process repeated three times total, including 24 h of room temperature incubation between cycles. This room temperature incubation is intended to allow spores that survived the first round of autoclaving an opportunity to germinate and become more vulnerable to a second round to improve biotic clearing. Autoclaving soils is a common technique that is generally performed for 2-3 cycles, with 24-48 h of incubation between each cycle, and allows for large-scale processing of soil (e.g. 6, 10, 11, 14, 17, 23-25, 33, 34, 35).

**Gamma irradiation-based sterilization**

A portion of each prepared substrate was sent to The Pennsylvania State University’s Radiation Science and Engineering Center’s Gamma Irradiation Facility. Samples were double-bagged in polypropylene and irradiated at a dose of 50 kGy (17, 18, 36, 37), using Co60 neutron sources in a Gammacell 220 dry cell irradiation chamber. Samples were stored at 4°C while queued at the facility and stored at 2°C upon return (~10 days).

**Soil saturation test and wash**

Portions of the sterilized substrate samples underwent a wash procedure, as it has been suggested that washing soil may eliminate compounds released during the sterilization process (25-30). In order to estimate the saturation point of each substrate, we used a modified version of the approach used in O. Priha and A. Smolander (38). We placed 5 g of unsterilized soil, 5 g of autoclaved substrate, and 5 g of irradiated substrate for each of the two soils onto separate layers of thin nylon. Nylon was used to allow drainage while retaining the bulk of the soil sample. Water was poured over the soil until the physical structure appeared to begin breaking down (max. added = 5 mL). Samples sat for 1 h to facilitate drainage. To estimate the saturation point, any excess water not added from the first pour step was combined with the drain-through for each sample. The amount of water retained by the soil was determined to be the measured volume of the leftover and drained water subtracted from the initial 5 mL added.

Once this point was determined, autoclave-sterilized DI water (20 min at 121°C and 20 psi) was used to wash the soils. For each sterilized substrate, portions were spread over the top of a standard 250 µm brass sieve. Water was poured over the substrate (1:1 v/v ratio) to exceed the determined saturation points for each substrate, so that water would drain through the soil rather than being completely absorbed. Substrates were drained to remove excess water for 45 min. Sieves were wiped and ethanol-sterilized between each sample to avoid contamination. Once washed, each substrate was divided into sterile containers with loose fitting lids and stored at room temperature for 72 h to continue removing excess moisture through evaporation.

**Microcosm assembly and inoculation**

For all microcosms, ~25 g of each type of soil or substrate was added to a small Petri dish (~50 mm x 10 mm). For recolonization (henceforth known as the natural recolonization method), small portions of unsterilized soil (~2 g) were placed in the center of small, ethanol-sterilized nylon squares. The nylon was gathered around the soil, pinched, and held closed with 24-gauge beading wire to create inoculation sachets. Excess nylon was cut from the top to ensure Petri dishes could still close with the sachets inside. To establish these microcosms, sachets were placed several mm in from the edge of a small Petri dish. Each substrate was added to surround the sachets on all sides, with the wire closure remaining visible above the surface of the surrounding soil. The soil inside the sachets had the same origin as the surrounding soil (i.e. if Soil 2 was placed inside the sachet, Soil 2 was also used to fill the microcosm). The sachets facilitate direct soil-to-soil contact (8), allowing microorganisms present in the unsterilized soil to pass into the surrounding sterilized soil, while minimizing the transfer of large soil particles.

Microcosms containing soil which was not washed prior to construction were saturated using autoclave-sterilized water. Volumes were chosen based on the previously described saturation estimates. No additional water was added to washed soils, as they were saturated during the washing procedure. Some microcosms were constructed with sterile soil only (no inoculum introduced) to allow for potential microbial regrowth in the absence of microbial inoculation. The microcosms were then wrapped with a thin layer of parafilm to prevent desiccation and incubated in a growth chamber for 8 weeks at 20°C and 85% relative humidity. Microcosms were randomly scattered across a single shelf within the growth chamber to minimize location bias.

**Microcosm sampling and vitality testing**

After the 8-week incubation, microcosms were removed from the growth chamber and aseptically sampled. For each microcosm, two 1.5 mL Eppendorf tubes were filled with the soil farthest away from the marked inoculation point and stored at -80°C for future analysis.

A third Eppendorf tube was partially filled for one sample in each treatment; this sample was mixed 1:1 with sterile water, diluted to 10-3, and 25 µL was plated onto Tryptic Soy Agar. Plates were incubated at room temperature for 48 h. This allowed a preliminary visualization of whether microbes had survived the incubation. Once the viability of the samples was confirmed, samples were processed for amplicon sequencing.

**DNA extraction and amplicon sequencing**

DNA was extracted from ~300 mg of each harvested soil sample using NucleoSpin 96 Soil kits (Machery-Nagel, Germany). Microbial composition was characterized with 16S rRNA gene (515F and 806R; V4 region) and ITS region (ITS1F and 58A2R) amplicon sequencing. The PCR ingredients for both reactions were as follows: 12 µL of 5Prime HotMasterMix, 1.5 µL of each primer (10 µM), 1.5 µL template DNA and 13.5 µL molecular grade water for a final PCR volume of 30 µL. Bacterial 16S rRNA gene PCR cycling conditions were as follows: 3 min at 94°C, 25 cycles of: 45 sec at 94°C, 60 sec at 50°C, and 90 sec at 72°C, and a final elongation step of 10 min at 72°C. Fungal ITS PCR cycling conditions were as follows: 3 min at 94°C, 35 cycles of: 20 sec at 94°C, 30 sec at 45°C and 45 sec at 72°C, and a final elongation step of 5 min at 72°C. The resulting amplicons were cleaned using Mag-Bind TotalPure NGS magnetic beads (Omega Bio-Tek; catalogue: M1378-01) and cleaned amplicons were indexed, normalized, concentrated and purified. Sequencing was performed at the Cornell University Biotechnology Resource Center Genomics Facility on an Illumina MiSeq using the 2×250 cycle v2 kit. Raw reads are available in the NCBI Sequence Read Archive under BioProject number PRJNA727754.

**Sequence analysis**

Raw demultiplexed 16S rRNA gene and ITS region data were processed using the Quantitative Insights into Microbial Ecology (QIIME 2 version 2020.11) pipeline (39). Briefly, paired-ended 16S rRNA gene and ITS region DNA sequences were imported and trimmed, and denoised using DADA2 which also removes chimeras (40). The classify-sklearn qiime feature classifier was used to assign taxonomy against the Silva v132 (41) or UNITE v04.02.2020 (v8.2) database (42) at the single nucleotide threshold (ZOTUs; zero-radius OTUs). The dataset was further cleaned by removing sequences identified as chloroplasts or mitochondria. The cleaned 16S rRNA gene and ITS data were then rarefied at 5,732 and 2,430 sequences per sample, respectively.

**Statistical analysis**

Statistical comparisons were performed in the R statistical environment (43) version 4.2.1. To identify differences in alpha diversity metrics (i.e., Chao1 species richness and Shannon species diversity) for the natural recolonization treatment, we used ANOVA followed by a posthoc TukeyHSD test from the *stats* package (version 4.2.1). Homogeneity of variance and normality of residuals were tested using the Levene’s and Shapiro-Wilk’s tests in the *car* (44) (version 3.1-2) and *stats* packages, respectively, and data were transformed when needed. A Kruskal-Wallis test was used to compare sterilization method for the soil regrowth soils. Bacterial and fungal compositional matrices were imported into the *phyloseq* R package (45) (version 1.4). We chose rarefaction and proportional transformation (i.e., relative abundance), as these produce the most accurate representations of community-level patterns and account for uneven sequencing depth (46, 47). Comparisons of microbial composition at the ASV level were performed with a Principal Coordinates Analysis (PCoA) ordination with a Bray-Curtis dissimilarity index and with a PERMANOVA (Adonis2) from the *vegan* package (version 2.6-4) with 999 permutations (48). To examine shifts in taxa according to sterilization and wash treatment, we performed a SIMPER analysis from the *vegan* package on data summarized at the Phylum and Class levels for bacteria and fungi, respectively. To determine the degree of dissimilarity between sterile soil regrowth and microbial recolonization, we calculated Bray-Curtis dissimilarity values of recolonized soil relative to soil regrowth, which were compared using a Kruskal-Wallis test from the *stats* package and a posthoc Dunn test from the *FSA* package (49) (version 0.9.4).

**Assessment of microbial regrowth in the absence of inoculation across autoclave conditions**

As we were limited in the number of soils and sample treatment conditions that we could test here, we performed a limited-scope follow-on study, in which we solely assessed the extent of microbial regrowth post-autoclaving in the absence of any microbial reinoculation. We tested regrowth across 7 distinct soils collected from Koffler Scientific Reserve at Jokers Hill (King City, ON, Canada) and the Campus Farm at the University of Toronto - Scarborough, as well as 6 modified treatment conditions in a single soil. Standard conditions were 3 autoclave cycles, with soil kept to 1.5” depth, and field-level moisture retained, with the following modifications for treatments: 1) pre-drying of soil; 2) reduction from 3 autoclave cycles to 1; 3) reduction from 1.5” to 0.5” depth; 4) increase from 1.5” to 3” depth; 5) 1:1 mixture of soil:peat (v:v); and 6) 1:1 mixture of soil:perlite. After 7 weeks of incubation in closed microcosms, DNA was extracted and quantified with a Qubit 4.0 fluorometer (Invitrogen, Carlsbad, CA, USA) across two separately incubated microcosms for each condition.

**Results and Discussion**

*More taxa were detected in gamma-irradiated soil regrowth microcosms*

We compared autoclaving and gamma irradiation to determine whether sterilization method had a significant impact on soil recolonization by microorganisms or microbial regrowth. For the natural recolonization method, we generally did not detect differences between sterilization method and wash treatment for species richness and species diversity, except for fungal alpha diversity in Soil 1 (Supplementary Table 2 and Supplementary Table 3) which was primarily driven by poor fungal recolonization (low alpha diversity values) in the washed autoclaved soils. These data indicate that sterilization method did not substantially impact alpha diversity in recolonized soils.

In a number of prior studies, we have seen that autoclaved soil had no detectable DNA or amplifiable PCR product shortly after sterilization and even after a period of incubation (e.g. L. M. Kaminsky et al. (12)). Other studies have also shown a lack of viable organisms following both sterilization approaches (e.g. A. E. Berns et al. (17)). However, we have noted that the diversity and rate of microbial regrowth in sterilized soils differs by soil type, which is an area of interest for future studies. In our sterile regrowth controls, there was significantly greater bacterial species richness (Soil 1: H = 4, p = 0.04; Soil 2: H = 5, p = 0.02), and bacterial diversity in Soil 2 (H = 5, p = 0.02), regrowth in the gamma-irradiated soils relative to the autoclaved soils (Supplementary Table 2). For fungal composition, we detected no significant differences for either species richness or species diversity in either soil. However, autoclaving had a particularly strong impact on the fungal soil regrowth controls (Figure 1) which may explain the lack of significance. In Soil 1, we observed two replicates with Chao1 values of 2 and 3 and an outlier replicate at 50. In Soil 2, only one replicate made it through the sequence analysis, as two replicates were removed during cleaning (only 3 to 8 input reads) and an additional replicate removed during rarefaction (only 280 cleaned sequences). As a consideration, we did not account for differences in total DNA and our observations are based on the relative abundance of those samples that could amplify. There was 17x and 7x greater species richness in gamma-irradiated regrowth soils when compared to autoclaved regrowth for bacteria and fungi, respectively (Figure 1). In support of our data, autoclaving has been shown to have a stronger negative impact on bacterial counts and CO2 production (19) and microbial cell lysis (17) relative to gamma irradiation. In this study, we used a gamma irradiation dose of 50 kGy. The efficacy of gamma irradiation can be impacted by soil properties, such as moisture content and organic matter content (18). Fungal composition is sensitive to gamma irradiation and can be impacted with doses of 0.01 kGy, with 10 and 50 kGy doses optimal to completely clear all viable fungi and spores in low and high organic matter soils, respectively (18). Further, doses of 20-25 kGy have been suggested as sufficient to clear bacteria (18, 20). While our chosen dose should have adequately eliminated bacteria and fungi from our soils, in both soils we observed trends towards substantially higher alpha diversity in the gamma-irradiated soil regrowth relative to the autoclaved soil regrowth control.



Figure 1: Alpha diversity measures for bacteria and fungi. ‘Unsterilized’ is untreated soil; ‘Sterile’ is sterile soil regrowth control; ‘Natural’ is natural recolonization of biotically-cleared soils from the unsterilized version of the same soil.

*Wash treatment had a greater impact than sterilization method on soil recolonization*

We assessed the impact of both sterilization method and soil washing (i.e., autoclaving or gamma irradiation) on microbial composition following natural recolonization (Figure 2; Supplementary Figure 1). For bacterial composition, we observed a significant effect of soil-type (F1,30 = 4, R2 = 0.1, p ≤ 0.001), wash treatment (F1,30 = 2, R2 = 0.05, p ≤ 0.001), and their interaction (F1,30 = 2, R2 = 0.05, p = 0.004), but observed no significant effect of sterilization method (p > 0.05; interaction with soil-type p > 0.05; Table 1). For fungal composition, we generally saw a similar pattern with a significant effect of soil-type (F1,30 = 4, R2 = 0.11, p ≤ 0.001), wash treatment (F1,30 = 3, R2 = 0.08, p ≤ 0.001) and their interaction (F1,30 = 4, R2 = 0.09, p ≤ 0.001) and no significant effect of sterilization method. We did observe a marginally significant interaction between soil-type and sterilization method (F1,30 = 2, R2 = 0.04, p = 0.05), and a three-way interaction between soil-type, sterilization method and wash treatment (F1,30 = 2, R2 = 0.04, p = 0.05), but they only explained a minor part of the composition variance (R2 = 0.04). These data suggest that sterilization method does not have a sizeable impact on soil recolonization when using soil as an inoculum source, while wash treatment plays a relatively larger role in influencing microbial recolonization. Soil washing is a common practice with numerous studies using soil washing to remove inhibitory compounds (25-30). In particular, soil washing can remove various soil toxins that inhibit bacterial growth (26) and has been shown to effectively remove fungal spores (31, 32). However, microbial colonization is generally not compared between washed and unwashed soils. Regardless, our data highlights that soil washing can impact soil microbial colonization and consideration should be applied when interpreting colonization patterns in washed soil and when extrapolating to field settings.



Figure 2: Principal Coordinates Analysis (PCoA) of bacterial and fungal composition following recolonization from a soil source. Points are coloured according to sterilization method and the shapes refer to different soil-types. Solid and hollow points refer to the no soil wash and soil wash treatments, respectively. 90 % ellipses are shown and are grouped according to sterilization method per soil-type. Ordinations per soil are shown in Supplementary Figure 1.

Table 1: Statistical analysis of beta diversity for bacterial and fungal composition after natural recolonization. The Adonis formula for both comparisons was Soil type \* Sterilization method \* Wash treatment. Comparison does not include the unsterilized soil.

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| Gene | Term | F-value | R2 | Significance |
| 16S rRNA gene | SoilSterilization methodWash treatmentSoil\*Sterilization methodSoil\*Wash treatmentSterilization method\*Wash treatmentSoil\*Sterilization method\*Wash treatment | F1,30 = 4F1,30 = 1F1,30 = 2F1,30 = 1F1,30 = 2F1,30 = 1F1,30 = 1 | 0.10.030.050.030.050.030.04 | \*\*\*NS\*\*\*NS\*\*NSNS |
| ITS region | SoilSterilization methodWash treatmentSoil\*Sterilization methodSoil\*Wash treatmentSterilization method\*Wash treatmentSoil\*Sterilization method\*Wash treatment | F1,30 = 4F1,30 = 1F1,30 = 3F1,30 = 2F1,30 = 4F1,30 = 2F1,30 = 2 | 0.110.040.080.040.090.040.04 | \*\*\*\*\*\*\*\*\*\*\*\*\* |

\*\*\* is ≤ 0.001, \*\* is < 0.01, \* is < 0.05

To explore whether sterilization method and soil washing had a significant impact on microbial recolonization for particular taxa, we performed a SIMPER analysis on data summarized at the Phylum and Class level for bacteria and fungi, respectively. For bacteria, we observed a greater relative abundance of the Verrucomicrobia and Chloroflexi phyla in the gamma irradiated recolonized soil (Table 2). For fungi, an unidentified fungal class and the Agaricomycetes were significantly higher in the gamma irradiated soils. Wash treatment had a larger impact on the recolonization of individual taxa relative to sterilization method, particularly for fungi (Table 2).

Table 2: SIMPER analysis of taxa between sterilization methods and wash treatments. Used data were summarized at the Phylum and Class levels for bacteria and fungi, respectively.

|  |  |  |  |
| --- | --- | --- | --- |
| Taxa | Autoclaved soil relative abundance (%) | Gamma-irradiated soil relative abundance (%) | p-value |
| Verrucomicrobia | 1.3 | 2.9 | 0.005 |
| Chloroflexi | 0.2 | 0.8 | 0.02 |
| Unidentified fungi | 16.8 | 41.4 | 0.003 |
| Agaricomycetes | 0.01 | 1.3 | 0.001 |
|  |
| Taxa | Not washed soil relative abundance (%) | Washed soil relative abundance (%) | p-value |
| Proteobacteria | 65.8 | 77.0 | 0.012 |
| Firmicutes | 15.4 | 4.2 | 0.002 |
| Unidentified fungi | 16.0 | 42.4 | 0.003 |
| Unclassified Fungi | 14.5 | 41.5 | 0.004 |
| Sordariomycetes | 29.4 | 4.1 | 0.03 |
| Pezizomycetes | 8.8 | 0 | 0.001 |
| Leotiomycetes | 3.3 | 0.03 | 0.003 |
| Rhizophydiomycetes | 0 | 2.3 | 0.001 |
| Unidentified Chytridiomycota | 0.03 | 2.2 | 0.02 |

Because of the strong influence of soil-type on microbial recolonization and to further explore the impact of soil washing on microbial recolonization, we examined the impact of washing on each individual soil (Figure 3). For each soil and for both bacterial and fungal composition, the wash treatment explained more of the compositional variance relative to the sterilization method used (Supplementary Table 4). We did not detect any differences for species richness (Chao1) between soil washing treatments for the recolonized soils, but there appeared to be a trend for lower species richness in recolonized washed soils. When compared to the unsterilized soils, we saw significant differences with the recolonized washed soils for Soil 1 (bacteria: autoclave q = 0.05, gamma q = 0.03) and Soil 2 (bacteria: autoclave q = 0.04; fungi: autoclave q = 0.002). To determine whether washed or unwashed soils more closely resemble the unsterilized soil, we extracted and compared Bray-Curtis dissimilarity values. For bacteria in each soil, and fungi in Soil 1, we observed no significant differences between the wash treatments. These data agree with our previous observations that the pool of recolonizing microorganisms is usually quite distinct from the bulk soil (14, 50). However, for Soil 2 and for fungi, we did observe a significant difference (H = 18, p ≤ 0.001) which was primarily driven by a greater similarity between the unsterilized soil and the washed soils for each sterilization method (average Bray-Curtis dissimilarity: autoclave wash = 0.81; autoclave no wash = 0.98, gamma wash = 0.87; gamma no wash = 0.99; q-values: autoclave wash vs. no wash = 0.002, gamma wash vs. no wash = 0.02).



Figure 3: Principal Coordinates Analysis (PCoA) of bacterial and fungal composition clustered according to the Wash treatment, and a boxplot of species richness (Chao1) for sterilization method and wash treatment. For the PCoA: the points are coloured according to sterilization method, shapes represent different soil-type, the fill color separates the wash treatment, and the solid and hollow points additionally refer to the no soil wash and soil wash treatments, respectively. 90 % ellipses are shown and are grouped according to the wash treatments.

*Microbial recolonization with both sterilization methods are distinct to soil regrowth*

Due to observing far fewer taxa in regrowth controls, especially following autoclaving, we sought to identify whether soil regrowth had a discernable impact on microbial composition in recolonized soils (Figure 4). In all instances, microbial recolonization was distinct from the soil regrowth (Supplementary Table 5). To identify whether recolonization in autoclaved or gamma-irradiated samples was more similar to microbial composition in regrowth controls, we extracted Bray-Curtis dissimilarity values relative to sterile soil regrowth. Generally, both bacteria and fungi in both soils were very distinct from sterile soil regrowth controls (average Bray-Curtis dissimilarity: bacteria = 0.98; fungi = 0.99) and we did not detect differences between sterilization methods and wash treatments for bacteria. For fungi, we detected significantly lower Bray-Curtis dissimilarity values for gamma-irradiated and washed soils (Supplementary Table 6). However, and while significant, the actual Bray-Curtis dissimilarity values were quite similar (Autoclaved and no wash = 0.9998; Gamma irradiated and no wash soil = 0.9958; Autoclaved and wash = 0.9937; Gamma irradiated and wash = 0.9881) suggesting that the impact of soil regrowth on microbial recolonization with either sterilization technique or wash treatment is minimal.



Figure 4: Principal Coordinates Analysis (PCoA) of bacterial and fungal composition for recolonized soils and sterile soil regrowth, and a boxplot of extracted Bray-Curtis dissimilarity values. For the ordination, points are colored according to sterilization method and the shapes refer to recolonized soil or sterile soil. Solid and hollow points refer to the no soil wash and soil wash treatments, respectively. 90 % ellipses are shown. For the boxplot, Bray-Curtis dissimilarity values relative to the sterile regrowth soil are shown.

To identify taxa that were common in our sterile regrowth, we examined the most abundant taxa in the autoclaved and gamma-irradiated sterile soils (Figure 5). Each sterilization technique permitted the regrowth of different taxa. While gamma-irradiation permitted more species richness of sterile regrowth (Figure 1), autoclaved sterile regrowth appeared to have fewer, but more dominant, taxa (Figure 5 and Supplementary Table 7). For example, bacteria assigned to the Bacillus genus made up 93 % relative abundance in one autoclaved regrowth sample but was 1.5 % in the next highest replicate (Supplementary Table 7). Gamma-irradiated soil regrowth appeared to be more consistent across replicates (Supplementary Table 7).



Figure 5: Average relative abundance of abundant bacterial (A) and fungal (B) genera in sterile regrowth soils. The top five most abundant taxa for each sterilization technique are shown.

 Future work will need to assess how soil characteristics and treatment conditions impact the extent of microbial regrowth in post-autoclaved soils. We have anecdotally observed differences in the degree of regrowth across the soils we have worked with. We never observe detectable DNA in soils directly post-autoclaving and in some cases, DNA stays below detection after many weeks of incubation. As a limited-scope follow-on study to this work, we assessed the amount of detectable DNA via Qubit following incubation of microcosms in the *absence* of any microbial reinoculation (Supplementary Figure 2 and Supplementary Table 8). Relative to DNA concentrations in initial soils, regrowth DNA was always very low, but varied across soil conditions and treatment conditions. For instance, it appeared that lower soil thickness at autoclaving and more autoclave cycles (3 vs. 1) led to less regrowth after 7 weeks. Future work that more robustly assesses the conditions affecting regrowth will improve our ability to generate diverse and effective model soil systems.

*Conclusions*

Biotically-cleared soils have been used for decades as a medium to study microbial colonization and persistence and relationships between biodiversity and community function (10, 11, 51-53). Biotically-cleared soils act as soil-like environments that more closely mimic natural conditions compared to culturing using commercially available media. Sterilization methods are still hotly debated, with gamma-irradiation often touted as the gold standard. In this study, we have shown that: (i) gamma-irradiated soils permitted greater microbial regrowth following sterilization, (ii) sterilization method had minimal impact on natural microbial recolonization, (iii) soil washing had a greater impact on microbial colonization relative to sterilization method, and (iv) fungi were particularly sensitive to autoclaving when examining our soil regrowth controls. The greater efficacy of autoclaving could be due to the repeated autoclaving steps which permit microbial regrowth and sporulation before autoclaving again (17). These data suggest that autoclaving and gamma-irradiation are both effective techniques for biotically clearing soil, and that autoclaving performed marginally better when considering soil regrowth.

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**References**

1. Overmann J, Abt B, Sikorski J.2017. Present and Future of Culturing Bacteria. Annual Review of Microbiology 71:711-730.

2. Stefani FOP, Bell TH, Marchand C, de la Providencia IE, El Yassimi A, St-Arnaud M, Hijri M.2015. Culture-Dependent and -Independent Methods Capture Different Microbial Community Fractions in Hydrocarbon-Contaminated Soils. PLOS ONE 10:e0128272.

3. Amann RI, Ludwig W, Schleifer KH.1995. Phylogenetic identification and in situ detection of individual microbial cells without cultivation. Microbiological Reviews 59:143-169.

4. Hugenholtz P, Goebel BM, Pace NR.1998. Impact of culture-independent studies on the emerging phylogenetic view of bacterial diversity. J Bacteriol 180:4765-74.

5. Pace NR.1997. A Molecular View of Microbial Diversity and the Biosphere. Science 276:734-740.

6. Swenson W, Wilson DS, Elias R.2000. Artificial ecosystem selection. Proceedings of the National Academy of Sciences 97:9110.

7. Panke-Buisse K, Poole AC, Goodrich JK, Ley RE, Kao-Kniffin J.2015. Selection on soil microbiomes reveals reproducible impacts on plant function. The ISME Journal 9:980-989.

8. Trexler RV, Bell TH.2019. Testing sustained soil-to-soil contact as an approach for limiting the abiotic influence of source soils during experimental microbiome transfer. FEMS Microbiology Letters 366:fnz228.

9. Lau JA, Lennon JT.2012. Rapid responses of soil microorganisms improve plant fitness in novel environments. Proceedings of the National Academy of Sciences 109:14058.

10. King WL, Richards SC, Kaminsky LM, Bradley BA, Kaye JP, Bell TH.2023. Leveraging microbiome rediversification for the ecological rescue of soil function. Environmental Microbiome 18:7.

11. Wagg C, Bender SF, Widmer F, van der Heijden MGA.2014. Soil biodiversity and soil community composition determine ecosystem multifunctionality. Proceedings of the National Academy of Sciences 111:5266-5270.

12. Kaminsky LM, Esker PD, Bell TH.2021. Abiotic conditions outweigh microbial origin during bacterial assembly in soils. Environmental Microbiology 23:358-371.

13. King WL, Kaminsky LM, Gannett M, Thompson GL, Kao-Kniffin J, Bell TH.2021. Soil salinization accelerates microbiome stabilization in iterative selections for plant performance. New Phytologist doi:10.1111/nph.17774.

14. King WL, Kaminsky LM, Richards SC, Bradley BA, Kaye JP, Bell TH.2022. Farm-scale differentiation of active microbial colonizers. ISME Communications 2.

15. López-Aranda JM, Miranda L, Medina JJ, Soria C, de los Santos B, Romero F, Pérez-Jiménez RM, Talavera M, Fennimore SA, Santos BM.2009. Methyl Bromide Alternatives for High Tunnel Strawberry Production in Southern Spain. HortTechnology hortte 19:187-192.

16. Hanson K, Mahato T, Schuch UK.2014. Soil Solarization in High Tunnels in the Semiarid Southwestern United States. HortScience horts 49:1165-1170.

17. Berns AE, Philipp H, Narres HD, Burauel P, Vereecken H, Tappe W.2008. Effect of gamma-sterilization and autoclaving on soil organic matter structure as studied by solid state NMR, UV and fluorescence spectroscopy. European Journal of Soil Science 59:540-550.

18. McNamara NP, Black HIJ, Beresford NA, Parekh NR.2003. Effects of acute gamma irradiation on chemical, physical and biological properties of soils. Applied Soil Ecology 24:117-132.

19. Ramsay AJ, Bawden AD.1983. Effects of sterilization and storage on respiration, nitrogen status and direct counts of soil bacteria using acridine orange. Soil Biology and Biochemistry 15:263-268.

20. Lensi R, Lescure C, Steinberg C, Savoie JM, Faurie G.1991. Dynamics of residual enzyme activities, denitrification potential, and physico-chemical properties in a γ-sterilized soil. Soil Biology and Biochemistry 23:367-373.

21. Hassi U, Hu J, Sleutel S, De Neve S.2023. Recovery of soil microbial community structure and activity following partial sterilization with gamma irradiation. Applied Soil Ecology 187:104839.

22. Bank TL, Kukkadapu RK, Madden AS, Ginder-Vogel MA, Baldwin ME, Jardine PM.2008. Effects of gamma-sterilization on the physico-chemical properties of natural sediments. Chemical Geology 251:1-7.

23. Arruda B, George PBL, Robin A, de L. C. Mescolotti D, Herrera WFB, Jones DL, Andreote FD.2021. Manipulation of the soil microbiome regulates the colonization of plants by arbuscular mycorrhizal fungi. Mycorrhiza 31:545-558.

24. McHaffie MB, Maherali H.2020. Variation in mycorrhizal growth response influences competitive interactions and mechanisms of plant species coexistence. Oecologia 192:755-765.

25. Kremer JM, Sohrabi R, Paasch BC, Rhodes D, Thireault C, Schulze-Lefert P, Tiedje JM, He SY.2021. Peat-based gnotobiotic plant growth systems for Arabidopsis microbiome research. Nature Protocols 16:2450-2470.

26. Greig-Smith R.1914. Contributions to our knowledge of soil fertility. Nos. vii to xi. Proceedings of the Linnean Society of New South Wales 38:725-746.

27. Lewis IM.1929. Bacterial antagonism with special reference to the effects of *Psuedomonas Fluorescens* on spore forming bacteria of soils. J Bacteriol 17:89-103.

28. Hou S, Thiergart T, Vannier N, Mesny F, Ziegler J, Pickel B, Hacquard S.2021. A microbiota–root–shoot circuit favours Arabidopsis growth over defence under suboptimal light. Nature Plants 7:1078-1092.

29. Wolinska KW, Vannier N, Thiergart T, Pickel B, Gremmen S, Piasecka A, Piślewska-Bednarek M, Nakano RT, Belkhadir Y, Bednarek P, Hacquard S.2021. Tryptophan metabolism and bacterial commensals prevent fungal dysbiosis in *Arabidopsis* roots. Proceedings of the National Academy of Sciences 118:e2111521118.

30. Getzke F, Hassani MA, Crüsemann M, Malisic M, Zhang P, Ishigaki Y, Böhringer N, Jiménez Fernández A, Wang L, Ordon J, Ma K-W, Thiergart T, Harbort Christopher J, Wesseler H, Miyauchi S, Garrido-Oter R, Shirasu K, Schäberle Till F, Hacquard S, Schulze-Lefert P.2023. Cofunctioning of bacterial exometabolites drives root microbiota establishment. Proceedings of the National Academy of Sciences 120:e2221508120.

31. Williams ST, Parkinson D, Burges NA.1965. An examination of the soil washing technique by its application to several soils. Plant and Soil 22:167-186.

32. Parkinson D, Williams ST.1960. A method for isolating fungi from soil microhabitats. Plant and Soil 13:347-355.

33. Howard MM, Bell TH, Kao-Kniffin J.2017. Soil microbiome transfer method affects microbiome composition, including dominant microorganisms, in a novel environment. FEMS Microbiol Lett 364:fnx092.

34. Li K, DiLegge MJ, Minas IS, Hamm A, Manter D, Vivanco JM.2019. Soil sterilization leads to re-colonization of a healthier rhizosphere microbiome. Rhizosphere 12:100176.

35. Gao D, Zhou X, Duan Y, Fu X, Wu F.2017. Wheat cover crop promoted cucumber seedling growth through regulating soil nutrient resources or soil microbial communities? Plant and Soil 418:459-475.

36. Zhang S, Cui S, Gong X, Chang L, Jia S, Yang X, Wu D, Zhang X.2016. Effects of gamma irradiation on soil biological communities and C and N pools in a clay loam soil. Applied Soil Ecology 108:352-360.

37. van Elsas JD, Chiurazzi M, Mallon CA, Elhottovā D, Krištůfek V, Salles JF.2012. Microbial diversity determines the invasion of soil by a bacterial pathogen. Proceedings of the National Academy of Sciences 109:1159.

38. Priha O, Smolander A.1999. Nitrogen transformations in soil under *Pinus sylvestris*, *Picea abies* and *Betula pendula* at two forest sites. Soil Biology and Biochemistry 31:965-977.

39. Bolyen E, Rideout JR, Dillon MR, Bokulich NA, Abnet CC, Al-Ghalith GA, Alexander H, Alm EJ, Arumugam M, Asnicar F, Bai Y, Bisanz JE, Bittinger K, Brejnrod A, Brislawn CJ, Brown CT, Callahan BJ, Caraballo-Rodríguez AM, Chase J, Cope EK, Da Silva R, Diener C, Dorrestein PC, Douglas GM, Durall DM, Duvallet C, Edwardson CF, Ernst M, Estaki M, Fouquier J, Gauglitz JM, Gibbons SM, Gibson DL, Gonzalez A, Gorlick K, Guo J, Hillmann B, Holmes S, Holste H, Huttenhower C, Huttley GA, Janssen S, Jarmusch AK, Jiang L, Kaehler BD, Kang KB, Keefe CR, Keim P, Kelley ST, Knights D, et al.2019. Reproducible, interactive, scalable and extensible microbiome data science using QIIME 2. Nature Biotechnology 37:852-857.

40. Callahan BJ, McMurdie PJ, Rosen MJ, Han AW, Johnson AJA, Holmes SP.2016. DADA2: High-resolution sample inference from Illumina amplicon data. Nature Methods 13:581-583.

41. Quast C, Pruesse E, Yilmaz P, Gerken J, Schweer T, Yarza P, Peplies J, Glöckner FO.2013. The SILVA ribosomal RNA gene database project: improved data processing and web-based tools. Nucleic acids research 41:D590-D596.

42. Kõljalg U, Larsson K-H, Abarenkov K, Nilsson RH, Alexander IJ, Eberhardt U, Erland S, Høiland K, Kjøller R, Larsson E, Pennanen T, Sen R, Taylor AFS, Tedersoo L, Vrålstad T.2005. UNITE: a database providing web-based methods for the molecular identification of ectomycorrhizal fungi. New Phytologist 166:1063-1068.

43. R Core Team. 2012. R: A language and environment for statistical computing, R Foundation for Statistical Computing,

44. Fox J, Weisberg S. 2019. An R Companion to Applied Regression, Third ed. Sage, Thousand Oaks CA.

45. McMurdie PJ, Holmes S.2013. phyloseq: An R Package for Reproducible Interactive Analysis and Graphics of Microbiome Census Data. PLOS ONE 8:e61217.

46. McKnight DT, Huerlimann R, Bower DS, Schwarzkopf L, Alford RA, Zenger KR.2019. Methods for normalizing microbiome data: An ecological perspective. Methods in Ecology and Evolution 10:389-400.

47. Weiss S, Xu ZZ, Peddada S, Amir A, Bittinger K, Gonzalez A, Lozupone C, Zaneveld JR, Vázquez-Baeza Y, Birmingham A, Hyde ER, Knight R.2017. Normalization and microbial differential abundance strategies depend upon data characteristics. Microbiome 5:27.

48. Oksanen J, Blanchet FG, Kindt R, Legendre P, Minchin P, O'Hara B, Simpson G, Solymos P, Stevens H, Wagner H.2015. Vegan: Community Ecology Package. R Package Version 22-1 2:1-2.

49. Ogle DH, Wheeler P, Dinno A. 2020. FSA: Fisheries Stock Analysis,

50. Richards SC, King WL, Sutherland JL, Bell TH.2024. Leveraging aquatic-terrestrial interfaces to capture putative habitat generalists. FEMS Microbiology Letters 371:fnae025.

51. Griffiths BS, Ritz K, Wheatley R, Kuan HL, Boag B, Christensen S, Ekelund F, Sørensen SJ, Muller S, Bloem J.2001. An examination of the biodiversity–ecosystem function relationship in arable soil microbial communities. Soil Biology and Biochemistry 33:1713-1722.

52. Griffiths BS, Ritz K, Bardgett RD, Cook R, Christensen S, Ekelund F, Sørensen SJ, Bååth E, Bloem J, De Ruiter PC, Dolfing J, Nicolardot B.2000. Ecosystem response of pasture soil communities to fumigation-induced microbial diversity reductions: an examination of the biodiversity–ecosystem function relationship. Oikos 90:279-294.

53. Evans E.1954. Soil Recolonization Tube for studying Recolonization of Sterilized Soil by Micro-organisms. Nature 173:1196-1196.