Make the environment protect you from disease

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Make the environment protect you from disease: elevated CO$_2$ inhibits antagonists of the fungus-farming termite symbiosis

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Fungus-farming termite colonies maintain monoculture fungus combs in underground chambers without apparent problems with diseases. Multiple lines of defense contribute to the suppression or removal of antagonists of the symbiosis, but the role of the termite-manipulated environment within mounds has yet to be tested. Specifically, termite mounds have extremely high levels of CO$_2$ compared to atmospheric levels. We tested the effect of 5% CO$_2$ on the growth of fungal crops from Macrotermes bellicosus colonies, generalist fungi that could challenge the symbiosis, as well as a specialist stowaway fungus, Pseudoxylaria. For sporulating fungi, we also quantified the effects on conidia production. We found that elevated CO$_2$ significantly reduces mycelial growth and conidia production of the generalist fungi Aspergillus sp., Beauveria bassiana, and Metarhizium brunneum, whereas it overall had a net positive effect on the growth of the fungal crop Termitomyces and Pseudoxylaria; albeit, with variation between fungal strains within genera. Our findings point to elevated CO$_2$ being of adaptive significance to the fungus-farming termite symbiosis as an additional layer of defense that helps keep termite fungus gardens free from fungal infections. The mound-building activities that make termites ecosystem engineers may thus also generate environmental conditions that impact the fate of fungi inhabiting the extended phenotypes that massive termite mounds represent.

KEYWORDS
Termitomyces, Macrotermiteinae, Macrotermes bellicosus, extended phenotype, defense, Pseudoxylaria

Introduction

Colony constructs in social insects allow for radically different internal environments compared to the outside, allowing essential homeostasis despite fluctuating external conditions. The mounds built by fungus-farming termite species of the subfamily Macrotermiteinae (Blattodea; Termitidae) are prominent features in African and Southeast Asian ecosystems (Figure 1A). Obligate mutualism with the fungal genus Termitomyces (Basidiomycota; Lyophyllaceae) allows these termites to obtain near-complete degradation of diverse plant
The termites maintain their monoculture fungal crop on structures of partially digested plant material (fungus comb Figure 1D), and the fungus provides nutrition for the termite hosts (da Costa et al., 2019). The termites, in turn, provide the fungus with plant substrate, protection, and optimal growth conditions (Korb, 2003; Korb, 2011). Monoculture farming should attract antagonists that could spread fast in the genetically homogeneous crop, but individual colonies can persist for decades (Wisselink et al., 2020) and do not appear to be challenged with diseases (Otani et al., 2019). Furthermore, non-<i>Termitomyces</i> fungi comprise only a minute proportion of the comb biomass (Moriya et al., 2015; Otani et al., 2019), and weeds are most likely accidentally brought in by foraging termite workers (Thomas, 1987; Guedegbe et al., 2009; Bos et al., 2021). It is only when the termites are removed or the colony is dying, that antagonists and stowaway fungi, such as members of the subgenus <i>Pseudoxyalaria</i> (Ascomycota: Xylariaceae), appear (Visser et al., 2011).

The termites protect <i>Termitomyces</i> through a series of complementary defenses that collectively provide very robust protection. The termites monitor and weed the fungus comb (Katariya et al., 2017), avoid substrates that contain mycopathogens (Bodawatta et al., 2019), remove and bury <i>Pseudoxyalaria</i> to generate hypoxia that kills the stowaway (Katariya et al., 2018), and utilize antimicrobial compounds of their own (Lamberty et al., 2001) or symbiont origins (Um et al., 2013; Schmidt et al., 2022; Murphy et al., 2023). Recent work has also identified volatile terpenes within the headspace surrounding the fungus comb that could be antimicrobial (Burkhardt et al., 2019; Kreuzenbeck et al., 2022). The comb headspace can play a role in defense as the termites ensure an enclosed, homeostatic comb environment, generated through elaborate ventilation structures that maintain consistent temperature and humidity (Korb, 2003; Murphy et al., 2023).

The tightly controlled comb environment, together with the high metabolism of the termites and <i>Termitomyces</i> and the processing of large quantities of decaying plant material, leads to high amounts of CO<sub>2</sub> (Darlington et al., 1997; Konate et al., 2003; Murphy et al., 2023). This leads to the accumulation of CO<sub>2</sub> that can be up to 5% in the central part of the mound in <i>Macrotermes carbonarius</i> (Matsumoto, 1978) and ~1.2% in the ventilation turrets of <i>Macrotermes bellicosus</i> (Korb and Linsenmair, 1999; Korb and Linsenmair, 2000; Turner, 2001). This is vastly higher than the ~0.04% CO<sub>2</sub> in the surrounding atmosphere and a product of limited gas exchange combined with high metabolism of termites and their fungal crop (Matsumoto, 1978), which are adapted to and benefit from these conditions (Luescher, 1961; Matsumoto, 1978; Korb, 2003). High CO<sub>2</sub> negatively affects the growth of most organisms, including negative effects on the respiration of the fungal crop of leaf-cutting ants (Kleineidam and Roces, 2000; Romer et al., 2017). However, <i>Termitomyces</i> does not appear to be affected (Katariya et al., 2018). Elevated CO<sub>2</sub> could thus aid in the selective suppression of competitor or pathogenic fungi, such as the specialist <i>Pseudoxyalaria</i>. This was first proposed by Batra and Batra (1966). However, beyond recent investigations of the impact of hypoxia in low and high CO<sub>2</sub> conditions (Katariya et al., 2018), the role of elevated CO<sub>2</sub> has not been tested.

To test this hypothesis, we evaluated the effect of elevated CO<sub>2</sub> on the growth of <i>Termitomyces</i>, <i>Pseudoxyalaria</i>, and other ecologically relevant fungi associated with fungus-farming termites. After confirming high CO<sub>2</sub> levels within <i>Macrotermes bellicosus</i> colonies, we subsequently quantified growth of 15 fungal strains from six genera under ambient (0.04%) and elevated (5.00%) CO<sub>2</sub> concentrations in <i>vitro</i>. We predicted that <i>Termitomyces</i> and <i>Pseudoxyalaria</i>, which are specialized to life in these conditions, would

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**FIGURE 1**

In situ CO<sub>2</sub> measurements in Côte d’Ivoire. <i>Macrotermes</i> termites construct elaborate mounds to ensure optimal microclimate for cultivation of <i>Termitomyces</i>. (A) A termite mount of <i>Macrotermes bellicosus</i> colony IC0039 in Côte d’Ivoire. (B) Schematic depiction of the setup for in situ measurements of CO<sub>2</sub> levels within termite mounds. (C) Field measurement. (D) Belowground comb structure comprised of plant and fungus biomass. (E) Box plot of CO<sub>2</sub> levels in the morning (left), at midday (middle) and in the evening (right) with colors represents different termite colonies. Whiskers extend to 1.5 * Interquartile Range (IQR) (n=6; significant difference are shown as *p<0.05; ***p<0.0001).
be unaffected or benefit from elevated CO₂, while opportunistic fungi should exhibit reduced growth or conidia production when exposed to mound CO₂ concentrations.

Materials and methods

**In situ measurements of CO₂ concentration, humidity, and temperature**

We identified two *Macrotermes bellicosus* colonies (IC0039 and IC0054) close to the Lamto Ecological Research Station in Côte d’Ivoire (Supplementary Table 1) and confirmed termite species with PCR amplification of the COXII gene (Zaman et al., 2022). We measured CO₂ concentrations using a 58 Miniature 5% CO₂ sensor. The sensor was placed in a falcon tube with holes to protect the probe from the termites while ensuring airflow. We placed the CO₂ probe as close to the central fungus chamber as possible via a small hole in the mound wall and avoided the probe touching anything that would agitate the termites (Figure 1B). We closed the hole with mound soil and left the termites to seal their mound properly and re-establish comb headspace conditions. The following day, we started CO₂ measurements in real-time in the morning (07:45–08:15), mid-day (13:30–14:00), and in the evening (19:30–20:00) by connecting the USB to the computer using Gaslab® (Figures 1B,C). We simultaneously measured temperature and humidity using iButtons (Hygrochron iButton®), which were placed in the falcon tube with the CO₂ sensor. For colony IC0039, it was possible to register CO₂ for four consecutive days; however, the termites in colony IC0054 covered the falcon tube within a few days, restricting the airflow to the CO₂ probe, which left us with only 2 days of measurements (Supplementary Table 2).

**Fungal isolations and barcoding**

We tested the impact of elevated CO₂ in vitro on 15 fungal strains from six genera (Supplementary Table 1). We obtained *Termitomyces* isolates from six colonies (hosting two *Termitomyces* species; see Results) in Côte d’Ivoire by aseptically placing nodules (asexual fruiting structures) on potato dextrose agar (PDA; VWR, 39 g/L) and subculturing until pure. We also obtained strains from three species of the stow-away fungus *Pseudoxylaria* from South Africa (X802) and Côte d’Ivoire (IC0040-PS and IC0057-PS). These were isolated by leaving fungus comb in moist conditions for 1–3 days without termites present, transferring emerging *Pseudoxylaria* mycelium to PDA, and subculturing until pure. Strains from two locations were tested as *Pseudoxylaria* species appear to be generalists of fungus-farming termites (Visser et al., 2009). We also included three *Aspergillus* strains (three species) isolated from dead *Macrotermes natalensis*, *M. bellicosus* and *Trinervitermes* sp. termites in South Africa or Côte d’Ivoire two entomopathogenic fungi (*Metarhizium brunneum* and *Beauveria bassiana*) that were originally isolated from *Cydia pomonella* larvae (Lepidoptera: Tortricidae) in Austria and an adult of *Anthocoris nemorum* (Hemiptera: Anthocoridae) in Denmark (Meyling et al., 2009), respectively and finally a mycopathogen, *Trichoderma harzianum* T22, originally produced by the fusion of *T. harzianum* strains T12 and T95 (Ahmad and Baker, 1988; Sivan and Harman, 1991).

**Strain genotyping**

We verified species identities of the fungal strains by sequencing the Internal Transcribed Spacer (ITS) region of the nuclear ribosomal DNA (Schoch et al., 2012). For DNA isolation, we used a Chelex protocol as described in Conlon et al. (2022). We used either the basidiomycete-specific primers ITS1F and ITS4B (Gardes and Bruns, 1993) or the ascomycete-specific primers ITS5F and ITS4B (White et al., 1990). PCR reactions were run with the following conditions: 94°C for 4 min followed by 35 cycles of 94°C for 30s, 58°C for 30s, and 72°C for 30s with a final extension step at 72°C for 4 min. PCR products were checked using agarose gel electrophoresis and purified using ExoSAP-IT™ (Affymetrix Inc., United States). Purified PCR products were sent to Eurofins MWG Operon (Ebersberg, Germany) for sanger sequencing. Sequences were identified using NCBI BLAST to estimate the number of species used for each fungal genus, with species delimitation based on a similarity threshold of >97% between ITS sequences, as this has been shown to be suitable for estimating the number of species within most fungal genera (Blaalid et al., 2013).

**In vitro effects of colony CO₂ condition on fungal growth and sporulation**

All fungi were grown on PDA supplemented with agar (10 g/L, VWR). First, plates of each fungus were grown until harvestable. Then, a ~2 cm² area of fungal biomass was scraped off the agar and deposited in a 150 mL solution of 0.05 g/L Tween 80 (MERCK) and 0.05 g/L Agar (Bie & Berntsen). After vortexing, a sterile inoculation needle was dipped in the solution, and then poked onto the middle of a 90 mm petri dish with PDA. We did 10 biological replicates for each fungal strain and each CO₂ treatment, considering a single plate per condition a replicate. This was repeated twice on separate dates, resulting in 20 replicates (with the exception of strain M17-03 (*M. brunneum*), for which we only measured growth once). Plates were incubated at 30°C and 70% relative humidity at either ambient CO₂ or a Digital incubator (INCU-Line®, VWR, Denmark) or at 5.00% CO₂ in a Midi 40 CO₂ Incubator (Thermo Scientific, Germany). Due to differences in growth rate, the strains were incubated for either 9 days (*Termitomyces* and *Pseudoxylaria*), 4 days (*M. brunneum*, *B. bassiana* and *Aspergillus*), or 2 days (*T. harzianum*). The difference in incubation time was set to ensure that the fungus would not overgrow the Petri dish and thus prevent correct measurements, while ensuring that there was enough growth to measure area. To compare growth across strains, we obtained the mycelial growth area by measuring the area in ImageJ (Rasband, 2023). If the colony was circular, we measured the diameter twice and used the average of the two measurements to calculate the growth area. If the colony was not circular, we obtained the growth area by manually drawing around the colony. Due to a systematic
effect of date, we standardized the measurements by subtracting the mean area of all replicates from the individual replicate area and then divided with the standard deviation of all replicates. We verified that CO₂ concentrations within Petri dishes were consistently at ambient or 5.00% using the S8 Miniature 5% CO₂ sensor. All individual measurements and calculations are available in Supplementary Table 3.

For Aspergillus strains N1IC19 and DT1W, and for M. brunneum M17-03, asexual spores (conidia) could be harvested after incubation, so we counted conidia production per plate after 4 days of growth for both CO₂ concentrations. We did this by depositing 10 mL 0.05% Triton-X (Sigma) on each fungal plate and rubbed the culture with a Drigalsky spatula to obtain a suspension of conidia, which was washed twice in 0.05% Triton-X by centrifugation and then discarding of the supernatant. We subsequently did a dilution series and counted conidia using a hemocytometer (Fuchs-Rosenthal). To test if prior exposure to elevated CO₂ affected sporulation even under ambient CO₂ levels, we pipetted 100 µL of a 1,000x dilution from plates grown at 0.04% or 5.00% CO₂ onto new PDA plates, spread conidia using a Drigalsky spatula, and counted the number of live (germinated) vs. dead (not germinated) conidia after overnight incubation. We did this for at least two replicates per strain. Neither Termitomyces or Pseudoxylaria produce conidia in our lab conditions, precluding testing of the impact of CO₂ on conidia production.

Statistical analyses

All analyses were performed in R 3.6.3 (RCoreTeam, 2021). The in situ CO₂ measurements of termite mounds were compared with a two-way ANOVA testing for colony and time of day effects. Subsequently we determined effect sizes by using Cohen’s f and then conducted pairwise comparisons of time points with TukeyHSD post hoc testing. The effect of CO₂ level on fungal growth was analyzed using a linear model with the standardized growth area as the dependent variable and CO₂ treatment and replicate as fixed effects. Effects of date, we determined effect sizes by using Cohen’s f and then conducted pairwise comparisons with TukeyHSD showed that CO₂ levels varied across different times of day; morning vs. mid-day: adj-p = 0.0007, morning vs. evening: adj-p = 0.0122, mid-day vs. evening: adj-p = 0.2510. The average temperature within mounds was 26.6°C (SD ± 0.98°C) and relative humidity was 98.2% (SD ± 1.14%). The temperature decreased approximately 0.5°C from 8:00–11:00 and again from 18:00–23:00.

Fungal growth in different CO₂ conditions

Combining the six Termitomyces strains, we found a significant positive effect of elevated CO₂ on growth (post hoc paired t-ratio test; t = −2.339; df = 211; p = 0.0200), with an overall medium effect of CO₂ (Cohen’s f = 0.407). However, when evaluated individually, three strains exhibited significantly increased growth, two were unaffected, and one strain exhibited significantly decreased growth when exposed to elevated CO₂ (Figure 2A and Table 1). The positive effect of growth was most significant on the Termitomyces species associated with colonies IC0010 and IC0034, whereas we observed more variable effects of elevated CO₂ on the species associated with IC0027 and IC0033 (unaffected), IC0032 (negative), and IC0031 (positive). The three Pseudoxylaria species were overall positively affected by elevated CO₂ (t = −0.741; df = 89; p = 0.0318) with a medium effect (Cohen’s f = 0.397), driven by a significant positive effect on one of the strains (Figure 2B and Table 1). Two of the three Aspergillus strains were negatively affected by elevated CO₂ (N1IC19: t = 2.806; df = 27; p = 0.0052; GF10t1: t = 3.459; df = 23; p = 0.0006), as was B. bassiana (t = 5.319; df = 43; p < 0.0001) and M. brunneum (t = 3.360; df = 17; p = 0.0008), but not T. harzianum (t = −0.5614; df = 63; p = 0.5749) (Figure 2C and Table 1). The individual p-values and effect sizes are given in Table 1.

Conidia production and viability

Our evaluation of the impact of elevated CO₂ on conidia production and viability revealed that the two Aspergillus strains (N1IC19 and DT1W) produced fewer conidia after incubation at elevated CO₂ (post hoc paired t-ratio test; DT1W: t = 2.967; df = 15; p = 0.0053; N1IC19: t = 13.45; df = 15; p < 0.0001), but this was not the case for M. brunneum (t = 1.457; df = 9; p = 0.1538) (Figures 3A,B). Fungal strain did not significantly affect the number of viable conidia (Pearson χ² = 11.63; df = 6; p = 0.0709). We therefore pooled all results, and a subsequent test of the pooled results did not reveal an impact on the proportion of live vs. dead conidia under subsequent CO₂ conditions of strains originally grown at ambient vs. elevated CO₂ concentrations (one-way ANOVA; F1 = 0.4652; p = 0.7074) (Figure 3C).
Discussion

We confirmed extraordinarily high CO$_2$ concentrations within the mounds of *Macrotermes bellicosus*, and found consistent patterns between colonies, which allowed us to set an appropriate CO$_2$ level for our *in vitro* studies. These measurement align with previous findings in conspecifics (Matsumoto, 1978) and underline how specialists of the symbioses are adapted to these relatively
As expected for specialists of the symbiosis, five of the six *Termitomyces* strains and all *Pseudoxylaria* strains were either unaffected or exhibited increased growth with elevated CO$_2$, with some variability. The remaining strain of *Termitomyces* showed reduced growth under elevated CO$_2$, however only with a low to medium effect size. The mound microclimate may thus influence growth differently across species of specialists. This general overall effect underlines how these fungal genera are adapted to comb conditions and suggests that mound CO$_2$ conditions do not aid in the normal suppression of the stowaway fungus. In line with this, the absence of increased *Pseudoxylaria* growth at ambient (compared to elevated) CO$_2$ concentrations indicates that decreased CO$_2$ concentration when mound wall integrity is compromised is unlikely to be the signal that triggers rampant growth of *Pseudoxylaria* (Visser et al., 2011). Other defenses must therefore be more important to suppress this fungus, such as antifungals (Visser et al., 2012; Um et al., 2013; reviewed by Schmidt et al., 2022), termite weeding and grooming of the comb (Katariya et al., 2017; Murphy et al., 2023), or the competitive advantage that *Termitomyces* obtains through its dense inoculation during comb formation (Badertscher et al., 1983; Leuthold et al., 1989).

Elevated CO$_2$ suppressed the growth of several of the entomopathogens and generalist contaminants that may enter mounds with the plant substrates (Bos et al., 2021) and persist in trace amounts within fungus combs (Otani et al., 2019). *Aspergillus* spp. are ubiquitous fungi found in a variety of environments and substrates, such as soil, air, and decaying plant material, where they thrive as saprophytes (Mousavi et al., 2016). The *Aspergillus* strains exhibited reduced growth and conidia production for the two strains for which this could be quantified (Figures 2, 3). This implies that although the fungi may enter combs, they are unlikely to be adapted to the internal nest conditions. This suggests that tradeoffs exist that preclude adaptations to the environment within termite mounds. The two generalist entomopathogens also exhibited reduced growth, implying that elevated CO$_2$ may help prevent their spread between termite hosts within colonies. The discrepancy in the impact on conidia production on the fungal genera may imply that mechanisms affecting sporulation differ, which should be further explored, ideally across more strains and species per genus. In contrast to entomopathogens, the mycopathogen *T. harzianum* was unaffected by elevated CO$_2$. *Trichoderma* spp. are found in nearly all soil types worldwide, infect termite fungal combs (Mathew et al., 2012), and respond in species-specific ways to CO$_2$ conditions (Danielson and Davey, 1973). Since the mycopathogen was not impacted by elevated CO$_2$, it is likely critical that workers avoid plant material containing the fungus (Bodawatta et al., 2019), and that combinations of defenses collectively allow the termites to maintain robust defenses toward diverse fungi with different susceptibilities to environmental conditions.

Our findings do not determine the mechanism that causes growth suppression, but elevated CO$_2$ may reduce comb pH as CO$_2$ reacts with H$_2$O to form carbonic acid (Wang et al., 2016). This is consistent with comb pH levels being 4–5 in *Macrotermes* and *Odontotermes* spp. compared to 6–7 in the surrounding soil (Muwawa et al., 2014). Lower pH causes weak acid stress in *Saccharomyces cerevisiae* (Piper et al., 2001), while alkaline conditions have been shown to increase fungal virulence (Selvig and Alspaugh, 2011; Vylkova, 2017). The elevated CO$_2$ inside mounds could thereby not only affect the growth but also the virulence of potential pathogens. Species of *Metarhizium* prefer alkaline conditions and can increase surrounding pH (Lovett and St. Leger, 2015) to optimize growth and conidia production (Gao et al., 2009).
Lowered pH could thus limit proliferation of certain fungi, but as many fungi inhabit mildly acidic environments, such as soils and plant and animal surfaces (Vylkova, 2017), low pH in itself is most likely not enough to rid unwanted fungi. Increased CO₂ concentration also stimulates the CAM/PProtein kinase A pathway, which can regulate fungal growth, development, and conidia and secondary metabolite production (Azzam et al., 2010; Sun et al., 2022). We did not evaluate whether elevated CO₂ affected this pathway, but it could explain the observed effects. Irrespective of the causal effects of elevated CO₂ on growth and reproduction, our findings imply that comb CO₂ should impact fungal proliferation and ultimately reproduction, potentially in species-specific ways. The variation we observe implies that more elaborate comparisons are needed to establish how variable responses are to elevated CO₂ levels, and that different optima perhaps exist for specialists in the symbiosis vs. generalist pathogens and saprophages. Once established, a natural next step would be manipulation experiments to test the impacts of compromised CO₂ environments on colony health to establish the adaptive benefits of CO₂ in defense.

We guided our experiments based on CO₂ concentrations in the massive mounds of M. bellicosus, which were comparable to levels in another species that also builds large mounds, such as M. carbonarius (Matsumoto, 1978). CO₂ concentrations in species with comparably large colonies and ventilation turrets, i.e., members of the genera Macrotermes and Odontotermes that represent ~33% of the known species within the Macrotermitinae (Bánki et al., 2022) are likely similar. However, ventilation and the propensity for CO₂ to play a role in defense is conceivably different in termite genera with different colony structures. For example, species of the genera Microtermes and Ancistrotermes build small interconnected fungal comb chambers within the soil (Jouquet et al., 2003; van de Peppel and Aanen, 2020). Insights into CO₂ levels here are absent, and future work should take a comparative approach to establish in detail the environment within fungal combs as well as the potential for (or limitations of) CO₂ to act in defense.

The extended phenotype of the termite mound is a cornerstone in termite ecology. Our findings point to an additional adaptive role emerging from generating a comb environment that aids in the suppression of alien fungi. This speaks to the need to view disease suppression in a more holistic view than previously considered. It also implies that other complementary parameters may play roles in defense, such as reduced oxygen availability or volatile compounds present within the comb headspace. Preliminary work on the latter suggests that this is indeed plausible (Kreuzenbeck et al., 2022, 2023). In addition, the ability to tolerate the extreme environment within mounds is likely an adaptive homeostatic trait that the termite can fine-tune to provide not only a defensive barrier but also optimal growth conditions for their specific crop species (and strains). Furthering our understanding of how termite farmers manage this complex environment for optimal farming of their fungal crop for food, while effectively suppressing antagonists, should thus integrate comb abiotic conditions.

Data availability statement

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article/Supplementary material.

Author contributions

SS, NB, and MP: conceptualization and writing—original draft. SS, NB, KS, NK, and NM: data curation. SS, NB, MP, NM, and RM: methodology. SS, NB, and RM: formal analysis and visualization. SS, NB, RM, NM, KS, NK, and MP: writing—review and editing. MP and NK: supervision. MP: funding acquisition. All authors contributed to the article and approved the submitted version.

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Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Supplementary material

The Supplementary material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fevo.2023.1134492/full#supplementary-material

References


Bacillus sp. that inhibit potentially antagonistic fungi. *Sci. Rep.* 3:3250. doi: 10.1038/srep03250


