



# Fungi present distinguishable isotopic signals in their lipids when grown on glycolytic versus tricarboxylic acid cycle intermediates

Stanislav Jabinski<sup>1,2</sup> Vítězslav Kučera<sup>3,4</sup>, Marek Kopáček<sup>1,5</sup>, Jan Jansa<sup>4</sup>, Travis B. Meador<sup>1,2,5\*</sup>

<sup>1</sup>University of South Bohemia, Faculty of Science, Department of Ecosystem Biology, Branišovská 1760, 370 05 České Budějovice, Czechia

<sup>2</sup>Institute of Soil Biology and Biochemistry, Biology Centre CAS, Na Sádkách 7, 370 05 České Budějovice, Czechia

<sup>3</sup>Charles University, Faculty of Sciences, Albertov 6, 128 00 Praha, Czechia

<sup>4</sup>Institute of Microbiology CAS, Vídeňská 1083, 142 20 Praha, Czechia

<sup>5</sup>Institute of Hydrobiology, Biology Centre CAS, Na Sádkách 7, 370 05 České Budějovice, Czechia

*Correspondence to:* Travis B. Meador (travis.meador@bc.cas.cz)

**Abstract.** Microbial activity in soils controls both the size and turnover rates of large carbon (C) inventories stored in the subsurface, having important consequences for ~~the~~ partitioning of C between terrestrial and atmospheric reservoirs as well as ~~the~~ recycling of mineral nutrients such as nitrogen or phosphorus (often bound to the C) that support plant growth. Fungi are major decomposers of soil organic matter (SOM); however, uncertainty about the identity of ~~in the~~ predominant C substrates that fuel their respiration confound models of fungal production and SOM turnover. To further define the signals of microbial heterotrophic activity, we applied a dual hydrogen (H) and C stable isotope probing (SIP) approach on pure fungal cultures representing the phyla Ascomycetesmycota, Basidiomycetesmycota, and Zygomycetesmycota growing on monomeric (glucose, succinate) or complex substrates (tannic acid,  $\beta$ -cyclodextrin). Our findings demonstrate that the investigated species incorporated only minor amounts of inorganic C (provided as bicarbonate) into their membrane lipids, amounting to < 3% of lipid-C, with no consistent patterns observed between species or growth substrates. The net incorporation of water-derived H (i.e.,  $\delta a_w$ ) into lipids also did not differ significantly between incubations with monomeric versus complex substrates; however, growth on succinate solicited significantly higher  $\delta a_w$  values than glucose or  $\beta$ -cyclodextrin. This finding suggests that  $^2\text{H}$ -SIP assays have the potential to distinguish between microbial communities supported predominantly by substrates that are catabolized by the tricarboxylic acid cycle versus glycolytic pathway. Furthermore, the average  $\delta a_w$  value of heterotrophic fungal incubations [ $0.69 \pm 0.03$  (SEM)] is consistent with that observed for bacterial heterotrophs, and may be applied for upscaling lipid-based estimates of fungal production in environmental assays.



## Short Summary

Microbial production is a key parameter in estimations of organic matter cycling in environmental systems, and fungi play a major role as decomposers. In order to investigate fungal production and turnover times in soils, we incubated fungal pure cultures with isotopically labelled water and bicarbonate to investigate growth signals encoded into lipid biomarkers, which can be applied to improve flux estimates in environmental studies.

## 1 Introduction

Soil organic matter (SOM) is the major reservoir of carbon ( $1580 \times 10^{15}$  g C) in the biosphere, and active microbial populations act to redistribute this C to other reactive reservoirs, such as [the](#) atmosphere (Carson et al., 2001; Grinhut et al., 2007). Major uncertainties in modeling C and climate dynamics stem from insufficient knowledge on the controls of SOM degradation and transformation (Ciais et al., 2014; Lindahl and Tunlid 2015). Saprotrophic soil fungi are one of the major decomposers in soils, [who](#)-are known to degrade naturally occurring complex molecules such as lignin (Kirk & Farrell, 1987; Fioretto et al., 2005; Baldrian et al., 2011), cellulose (Šnajdr et al., 2011) and humic substances (Grinhut et al., 2007), but are also reported to compete for accessible plant photosynthate excreted by roots (De Boer et al., 2005; Högberg et al., 2001; Smith & Read, 2008). Despite the unique and important fungal niche in biogeochemical cycles, their contributions to SOM cycling remains poorly constrained (Frey 2019; Grinhut et al., 2007). Furthermore, heterotrophic organisms feeding on organic substrates to gain energy and build biomass are also known to fix a variable amount of inorganic C ([IC](#)), in order to replenish intermediates in the tricarboxylic acid (TCA) cycle (Kornberg 1965). It has been suggested that 2 - 8% of the biomass C in heterotrophs originates from [inorganic-IC](#) incorporated through anaplerotic carboxylation reactions (Romanenko 1964; Roslev et al., 2004; Braun et al., 2021). [Although, the](#) awareness of these processes has existed for decades (Kornberg 1965; Sorotkin 1966). [Yet,](#) the relevance and [the](#) metabolic controls on heterotrophic [inorganic-C-IC](#) fixation remains poorly understood, partly due to the lack of reliable estimates for most organisms and habitats (Braun et al., 2021).

Advanced analytical techniques now allow [linking](#) microbial taxa to [be linked to](#) specific processes in environmental studies by measuring [the](#) incorporation of stable isotopes into biomarkers (Boschker et al., 1998; Dumont and Murrell, 2005; Kreuzer-Martin, 2007), such as fungal and bacterial membrane lipid fatty acids (Treonis et al., 2004; Willers et al., 2015) or other biomarkers (Boschker and Middelbourg, 2002). Previous studies have demonstrated that variability in the  $^{21}\text{H}$  composition of microbial lipids is redundant with that of environmental water (Hoefs, 2018; Kopf et al., 2015), and stable isotope probing (SIP) assays applying enrichments in  $^2\text{H}_2\text{O}$  have proven to be a useful tracer of microbial activity in a diverse range of environments ([Canarini et al., 2024; Caro et al., 2023; Fischer et al., 2013; Kellermann et al., 2012; Wegener et al., 2016; Warren 2022; Wu et al., 2018](#)). Large H-isotope fractionations, yielding  $\delta\delta^2\text{H}_-$ -values between  $-400\%$  and  $+200\%$ , have been observed during biosynthetic incorporation of water hydrogen (water-H) into individual compounds within a single cell or total biomass, which can be indicative of [the underlying](#) metabolic processes (Osborn et al., 2011; Sachse et al., 2012; Zhang et al., 2009). To fully exploit the potential of SIP experiments, a dual-SIP approach was developed to track total microbial production



by adding heavy water ( $^2\text{H}_2\text{O}$ ) together with  $^{13}\text{C}$ -labeled ~~inorganic C (IC)~~, enabling simultaneous estimates of total and autotrophic metabolism, respectively (Wegener et al., 2012; Wu et al., 2020). Recently, Jabinski et al. (2024) validated an innovation of the dual-SIP assay by using rapid pyrolysis of fungal biomass to determine the stable C and H isotopic composition of fungal lipids, and demonstrated that water-H and IC assimilation signatures could successfully distinguish between fungal ecotypes growing on glucose or glutamic acid as the C source. The aim of the current study was to further assess the controls on water-H and ~~inorganic C~~ incorporation into lipids and expand our knowledge for interpreting environmental signals by applying the dual-SIP assay on a broader range of pure fungal cultures and growth substrates, including labile monomers versus more complex, high molecular weight molecules. We hypothesized that ~~(i)~~ the incorporation of ~~inorganic C~~ and water-H into the fungal fatty acid biomarker  $\text{C}_{18:2}$  will be similar for fungal species growing on the same substrate, and ~~(ii) H~~ that ~~inorganic C~~ and water-H incorporation will distinguish between growth on labile versus more complex C substrates.

## 2 Methods

### 2.1 Cultivation & Harvesting

Fungal pure cultures of two Basidiomycetes ~~mycota~~ [~~*Paxillus involutus* (PI, strain SB-22); *Phanerochaete chrysosporium* (PC, strain CCM8074)~~], two Zygomycetes ~~mycota~~ [~~*Mortierella* sp. (MO, strain RK-38); *Umbelopsis* sp. (UM, strain RK-43)~~] and two Ascomycetes ~~mycota~~ [~~*Penicillium janczewskii* (PJ, strain BCCO20\_0265); *Paecilomyces lilacinus* (PL, strain DP-23)~~] were incubated in 500 mL Schott bottles at 25 °C in the dark. ~~The Liquid mineral media (in 500 mL Schott bottles containing 50 mL) was adapted after of a mineral media described previously (Bukovská et al. (2018) with the vitamins left out, which and~~ was inoculated with approximately  $10^6$  spores, ~~or, for Basidiomycetes mycota~~, a hyphal block  $< 0.5 \text{ cm}^3$  (~~Basidiomycetes~~) recovered from a previous culture using the same cultivation medium solidified with agar (1.5%).

The growth medium contained per liter: 4 g organic C in various forms ( $\text{C}_6\text{H}_{12}\text{O}_6$  glucose;  $\text{C}_4\text{H}_6\text{O}_4$  succinic acid;  $\text{C}_{42}\text{H}_{70}\text{O}_{35}$   $\beta$ -Cyclodextrin or  $\text{C}_{76}\text{H}_{52}\text{O}_{46}$  tannic acid), 0.01 g  $\text{FeSO}_4 \cdot 7\text{H}_2\text{O}$ , 2 g  $\text{KH}_2\text{PO}_4$ , 0.5 g  $\text{MgSO}_4 \cdot 7\text{H}_2\text{O}$ , 0.1 g NaCl, 0.1 g CaCl, 2.5 g  $(\text{NH}_4)_2\text{SO}_4$ , 0.45 g  $\text{NaHCO}_3$  and 1 mL of a mixed solution (per liter: 0.5 g  $\text{H}_3\text{BO}_3$ , 0.04 g  $\text{CuSO}_4 \cdot 5\text{H}_2\text{O}$ , 0.1 g KI, 0.4 g  $\text{MnSO}_4 \cdot 5\text{H}_2\text{O}$ , 0.2 g  $\text{NaMoO}_4 \cdot 2\text{H}_2\text{O}$ , 0.4 g  $\text{ZnSO}_4 \cdot 7\text{H}_2\text{O}$ ). The pH of the medium was adjusted to 4.5 before inoculation. Dual-SIP experiments were performed using  $^{13}\text{C}$ -bicarbonate ( $^{13}\text{C}$ -DIC,  $\text{NaH}^{13}\text{CO}_3$ ) and deuterated water ( $\text{D}_2\text{O}$ ). Each fungal strain was grown in triplicate with non-labeled substrates (Treatment I), with  $\delta^2\text{H}$  of the medium water adjusted to 100‰ and  $\text{AT}^{13}\text{C} = 10\%$  of  $^{13}\text{C}$ -DIC (Treatment II), 200‰  $\delta^2\text{H}$  and 10%  $^{13}\text{C}$  -DIC (Treatment III), and 400‰  $\delta^2\text{H}$  and 10%  $^{13}\text{C}$  -DIC (Treatment IV). ~~The concentration of the bicarbonate in the cultivation medium was  $0.1 \text{ g L}^{-1}$ .~~ The Schott bottles were closed with a rubber stopper in order to ~~keep-prevent~~ the labeled  $^{13}\text{C}$ -DIC from outgassing, and ample headspace was provided to maintain oxic conditions throughout the growth experiment. Fungal growth was monitored via the accumulation of  $\text{CO}_2$  in the



headspace, and we aimed to harvest when CO<sub>2</sub> levels reached 10%; however, without preliminary knowledge of the fungal growth dynamics, some cultivations exceeded this level more quickly than they could be ~~sampled and analyzed~~ harvested. To harvest the fungal biomass, ~~M~~mycelia were separated from the growth medium via vacuum filtration through 5 µm Isopore polycarbonate filters (47 mm diam, Merck catalogue number TMTP04700) ~~using vacuum filtration device allowing to and the collect the~~ cultivation medium was collected into a sterile 50 mL tube. Thereafter, the mycelium was washed with ample MilliQ water, transferred to pre-weighed, sterile 50 mL tubes; ~~the~~ fresh weight of the biomass was recorded, and the samples were frozen at -80 °C until lyophilization. A subsample of the cultivation medium was also frozen at -80 °C and the rest was used to determine pH post-incubation. After lyophilization, the dry weight of each sample was determined and stored at -20 °C until further analysis.

## 2.2 Measurements

### 2.2.1 Headspace CO<sub>2</sub> concentration and isotope composition

Samples of headspace (0.3 mL) were collected weekly from each bottle into helium flushed 12 mL exetainer vials (Exetainer, Labco Limited, UK) and analyzed for their CO<sub>2</sub> concentration and <sup>13</sup>C/<sup>12</sup>C isotopic ratio using Gasbench II equipped with a single cryo-trap connected to Delta V Advantage isotopic ratio mass spectrometer (IRMS) via Conflow IV (Thermo Scientific, Bremen, Germany). Ambient air (with ~~the its~~ CO<sub>2</sub> concentration measured using LiCor 850 gas analyzer previously) was used as a standard for CO<sub>2</sub> concentration measurements, whereas a laboratory cylinder with 0.1% CO<sub>2</sub> in helium gas (~~δ<sup>13</sup>C = -2.86 ‰~~) was used as a standard for ~~the~~ isotopic composition (δ<sup>13</sup>C = -2.86 ‰) ~~of C in the the C~~. The analytical precision error was below 1‰. Data were analyzed and exported using ~~the~~ Isodat 3.0 software.

### 2.2.2 Medium water (²H₂O)

Liquid samples were transferred into 1.5 ml glass vials (32 x 11.6 mm, Fischer Scientific) and then measured using Triple Liquid Water Isotope Analyzer (Los Gatos Research), which is based on the principle of high-resolution laser absorption spectroscopy. Samples were dispensed into the instrument using an autosampler (PAL3 LSI, ABB company) and a 1.2 µL syringe (Hamilton). Samples were measured and evaluated against prepared laboratory standards of known isotopic composition. The isotopic ratios of these laboratory standards were verified by measuring against international standards (VSMOW2, SLAP2) made by the IAEA. For quality control purposes, the measurements of the samples were also interspersed with periodic measurements of the prepared verification samples with known isotopic composition. The final isotopic composition (δ²H) was determined using LIMS software. Analytical precision error of δ²H was <1.5‰.

Water sampled from incubations with tannic acid could not be measured using the laser, as described above, due to its high organic carbon content, and was rather measured via a GasBench II devicesystem (Thermo Scientific, Bremen, Germany; Application Note: 30049). Medium water samples (200 µL) were added with a platinum catalyst to a 12 mL exetainer vials



(Exetainer, Labco Limited, UK). The headspace was flushed with 1% H<sub>2</sub> in He at approximately 100 mL min<sup>-1</sup> ~~with~~ for 6 min. After an equilibration time of over 40 min, the samples were measured by purging the exetainer using a double-holed needle with helium into a 250 µL sample loop. The sample was then injected and separated via a Carboxen PLOT 1010 (0.53 mm ID; Supelco, Bellefonte, USA) held at 90 °C with a flow rate ~~set at of 0.75 bar~~ 2.2 mL min<sup>-1</sup>, and then introduced into the MAT253 Plus IRMS via a Conflo IV interface. Each sample was injected three times during one analysis. The isotopic composition was determined using Isodat 3.0 software against the corresponding H<sub>2</sub> working gas (-239‰ for  $\delta^2\text{H}$ ) and the values were corrected and normalized using international standards VSMOW2 (0‰ for  $\delta^2\text{H}$ ), SLAP2 (-427.5‰ for  $\delta^2\text{H}$ ), USGS53 (+40.2‰ for  $\delta^2\text{H}$ ) and GFLES-2 (159.9‰ for  $\delta^2\text{H}$ ). The analytical ~~precision~~ error was around 1‰.

### 2.2.3 Carbon ( $\delta^{13}\text{C}$ ) substrate analysis

Substrates (~100 µg) were weighed into tin capsules (8 ~~×~~ 5 mm, Sercon, Crewe, UK) and placed in a helium-flushed carousel autosampler, then introduced to an Elemental Analyzer IsoLink device (EA IsoLink CNSOH, Thermo Scientific, Bremen, Germany) equipped with a CHN/NC/N EA combustion/reduction reactor (Sercon, Crewe, UK) heated to 1020 °C. A pulse of oxygen was introduced to the reactor simultaneously with the sample. The sample gases were quantified via a thermal conductivity detector (TCD) and then introduced to a MAT 253 Plus isotope ratio mass spectrometer (IRMS; Thermo Scientific; Bremen, Germany) via the open split of a Conflo IV interface, with helium as the carrier gas. The isotopic composition was determined using Isodat 3.0 software against the corresponding CO<sub>2</sub> working gas (-4.191‰ for  $\delta^{13}\text{C}$ ), and the values were corrected for linearity and normalized to the VPDB scale using international reference material IAEA-600 (-27.771‰ for  $\delta^{13}\text{C}$ ). The analytical ~~precision~~ error was <0.04‰.

### 2.2.3 Pyrolysis GC for lipid analysis

The pyrolysis unit Shimadzu 3030D (Shimadzu, Kyoto, Japan/ Frontier Laboratories, Fukushima, Japan) was installed on top of the GC Trace1310 gas chromatograph SSL injector (Thermo Scientific, Bremen, Germany) and the GC was equipped with an SLB-IL60 column (non-bonded; 1,12-Di(tripropylphosphonium)dodecane bis(trifluoromethanesulfonyl)imide phase, 30 m, 0.25 mm ID, 0.20 µm df, Supelco, Bellefonte, USA). The furnace temperature was 650 °C and the interface temperature was 370 °C. The injector temperature was 360 °C and the GC oven was held at 80 °C for 1 min then ramped to 175 °C at 15 °C min<sup>-1</sup>, then ramped to 195 °C at 2 °C min<sup>-1</sup>, then ramped to 300 °C at 10 °C min<sup>-1</sup>, and finally held at 300 °C for 7 min. Helium was used as ~~the~~ carrier gas, with a constant flow of 1.5 mL min<sup>-1</sup> ~~with~~, a split ratio of 40, and a split flow of 26.7 mL min<sup>-1</sup>. The column flow was split via a multichannel device to acquire MS and isotopic data simultaneously from one injection. The GCMS (ISQ QD; Thermo Scientific, Bremen, Germany) ion source was set to electron impact ionization mode (EI) at 70 eV and a scan range of m/z 50 – 500 with a scan time of 0.2 sec<sup>-1</sup> ~~was applied~~. Scanning started after 8 min to avoid the solvent peak in the MS. The ~~t~~-Transfer line temperature was set to 300 °C and the ~~i~~on source was set to 250 °C.



The samples (~~lyophilized freeze-dry~~ biomass, 0.1 ~~mg~~– 1.3 mg) were weighed into an ultra-clean stainless steel Eco-Cup LF (Frontier Laboratories, Fukushima, Japan), which were burned with a torch before usage to ~~ensure no remove~~ contamination. ~~FAMES signals were acquired in the same run.~~ Immediately prior to ~~sample injection the measurement~~, 30 µL of trimethylsulfonium hydroxide (TMSH) was added on the sample to increase the volatilization of the fatty acids and improve measurement sensitivity. Identification ~~of fatty acid methyl esters (FAMES)~~ was performed using fragmentation patterns and the NIST 14 library.

Stable carbon and hydrogen isotope compositions of FAMES ~~and ergosterol~~ were determined by splitting the flow from the GC column to a GC-Isolink II reactor, coupled to a MAT253 Plus IRMS via a ConFlo IV interface. Values are expressed in standard delta notation ( $\delta^{13}\text{C}$  and  $\delta^2\text{H}$ ). MS information was simultaneously acquired by use of the multi-channel device described above. For conversion of FAMES and ergosterol to  $\text{CO}_2$ , the combustion reactor (nickel oxide tube with  $\text{CuO}$ ,  $\text{NiO}$ , and Pt wires) was set to 1000 °C. For conversion of FAMES and ergosterol to  $\text{H}_2$ , the pyrolysis reactor (aluminum tube) was set to 1420 °C.

The FAMES were identified by their retention times and fragmentation patterns. The isotopic composition was determined using Isodat 3.0 software against the corresponding  $\text{CO}_2$  or  $\text{H}_2$  working gas (-4.191‰ for  $\delta^{13}\text{C}$ , -239.5‰ for  $\delta^2\text{H}$ ). Isotope corrections for instrument drifts, linearity, and normalization to the VPDB or VSMOW scales were performed according to the response of USGS70 (-30.53‰ for  $\delta^{13}\text{C}$ , -183.9‰ for  $\delta^2\text{H}$ ) and USGS72 (-1.54‰ for  $\delta^{13}\text{C}$ , 348.3‰ for  $\delta^2\text{H}$ ) reference standards. The analytical ~~precision error~~ was < 0.5‰ and < 10‰ for  $\delta^{13}\text{C}$  and  $\delta^2\text{H}$ , respectively.

## 2.3 Calculations

~~Carbon use efficiency (CUE) for the growth experiments was calculated by normalizing the amount of C in biomass by that plus C that accumulated as  $\text{CO}_2$  [CUE = biomass-C / ( $\text{CO}_2$ -C + biomass-C)].~~

~~The  $\delta^{13}\text{C}$  values of fungal biomarker  $\text{C}_{18:2}$  was determined as described in section 2.2.4 and is reported in this section as standard  $\delta^{13}\text{C}$  values (‰). The inorganic-C incorporation into the biomarker (%IC) was calculated based on the following equation:~~

$$\%IC_{(\text{assimilation})} = \frac{{}^{13}F_{\text{lipids}} - {}^{13}F_{\text{lipid, control}}}{{}^{13}F_{\text{DIC}(\text{medium})} - {}^{13}F_{\text{substrate}}} \times 100$$

~~$F_{13C}$~~

$$\%IC_{(\text{assimilation})} = \frac{F_{13C}^{\text{lipids}} - F_{13C}^{\text{lipid, control}}}{F_{13C}^{\text{DIC}(\text{labeling medium})} - F_{13C}^{\text{lipid, substrate control}}} \times 100\%$$

(Eq. 1)





**Equation 1: %Inorganic carbon (%IC) assimilation was calculated as the difference in the  $^{13}\text{C}$  atom fraction of the lipids from harvested at the end of the  $^{13}\text{C}$  labeling experiment ( $F_{\text{lipid}}$ ) - compared to the lipids harvested at the end of the natural abundance treatment ( $F_{\text{lipid,control}}$ ), relative to the difference between the mixing-weighted average  $^{13}\text{C}$  atom fraction  $F_{\text{DIC}}$  of dissolved inorganic C ( $F_{\text{DIC}}$ , cf. Text S1) and the  $F_{\text{substrate}}$  of the substrate. F was calculated as  $F = (R^{13\text{C}/12\text{C}}_{\text{lipid}} / (R^{13\text{C}/12\text{C}}_{\text{DIC}} + 1))$ , where R is  $\delta^{13}\text{C}$  ratios as measured with the reported by Isodat Software following measurement by IRMS equipment using the reverse of the  $\delta$  notations ( $\delta^{13}\text{C} = 1000 \cdot (R^{13\text{C}/12\text{C}}_{\text{sample}} / R^{13\text{C}/12\text{C}}_{\text{ref}} - 1)$  \* 1000 (modified after Boschker & Middelburg 2002; Wegener et al., 2012).**

The water H assimilation efficiency ( $a_w$ ) — values for fungal biomarker  $\text{C}_{18:2}$  was determined approximated from the regressions of the hydrogen isotope composition of individual fatty acids  $^2F_{\text{FA}}$  and that of medium water ( $^2F_{\text{water}}$ ), according to Kopf et al., (2015) from the regressions of  $^2F_{\text{lipid}}$  and  $^2F_{\text{water}}$  (Zhang et al., 2009; Kopf et al., 2015). Briefly, the H hydrogen isotopic compositions of microbial fatty acids produced by an organism generally follow the isotopic composition of environmental water, and is described are related to in terms of the mole fraction of H contributed from spiked water in the cultivation medium ( $x_w$ ) and the associated net hydrogen isotope fractionation between lipids-fatty acids and water ( $\alpha_{\text{fa/w}}$ ), and the residual contribution from substrates ( $1 - x_w$ ) and associated net H isotope fraction between lipids and substrates ( $\alpha_{\text{fa/s}}$ ), where  $a_w = x_w \times \alpha_{\text{fa/w}}$ . Whereas  $a_w$  can be determined experimentally, the latter terms cannot be independently determined for heterotrophic growth (Kopf et al., 2015). The traditional isotope effects and  $\epsilon_{\text{C18:2/water}}$  and  $\alpha_{\text{C18:2/water}}$  were calculated after Hayes (2004), where  $\alpha_{\text{C18:2/water}} = [(\delta^2\text{H}_{\text{C18:2}} + 1000) / (\delta^2\text{H}_{\text{water}} + 1000)]$  and  $\epsilon_{\text{C18:2/water}} = (\alpha_{\text{C18:2/water}} - 1) \times 1000\text{‰}$ . is defined as water assimilation constant, and it can be obtained from the regressions of  $^2F_{\text{lipid}}$  and  $^2F_{\text{water}}$  (Zhang et al., 2009; Kopf et al., 2015).

## 3 Results

### 3.1 Fungal growth and $\text{CO}_2$ production

All fungal species were pure cultures, which were incubated in a mineral medium with either glucose, succinate,  $\beta$ -cyclodextrin, or tannic acid serving as the sole organic C source. Growth was monitored by the evolution of  $\text{CO}_2$  into the headspace, which ranged from 0.36% (no respiration of substrate) to a maximum of 35%, after incubation times ranging from 5 to 160 days (Fig. 1). The pH of the media in all incubations ranged from 2 to 5.5 at the time of harvest, with a general trend of decreasing pH with increasing  $\text{CO}_2$ ; however, the trend was opposite when succinate was the carbon source, with pH increasing from 4 to 5.5. For samples that produced sufficient biomass, the dry biomass of harvested fungal hyphae ranged up to 250 mg, and at least 30  $\mu\text{g}$  dry biomass was used to analyze fungal membrane fatty acids by Pyr-GC-IRMS. Only the Ascomycetes species PL and PJ grew sufficiently on each tested substrate to produce enough biomass for stable isotope analysis. Incubations of Zygomycetes species with glucose or succinate also yielded sufficient dry biomass, and only



214 UM ~~and not MO~~ was able to grow on  $\beta$ -cyclodextrin. Zygomycetes species produced neither CO<sub>2</sub> nor biomass when  
215 incubated with tannic acid. The Basidiomycetes typically exhibited the slowest growth, and both species (PI and PC)  
216 only produced enough biomass when grown on glucose. The headspace CO<sub>2</sub> levels in Basidiomycetes incubations with  
217 succinate increased to a maximum of ~ 2%, but only PI yielded sufficient biomass for analysis. PC grew sufficiently on  $\beta$ -  
218 cyclodextrin, with CO<sub>2</sub> levels increasing to a maximum of 3%, while CO<sub>2</sub> remained < 0.6% in PI incubations.



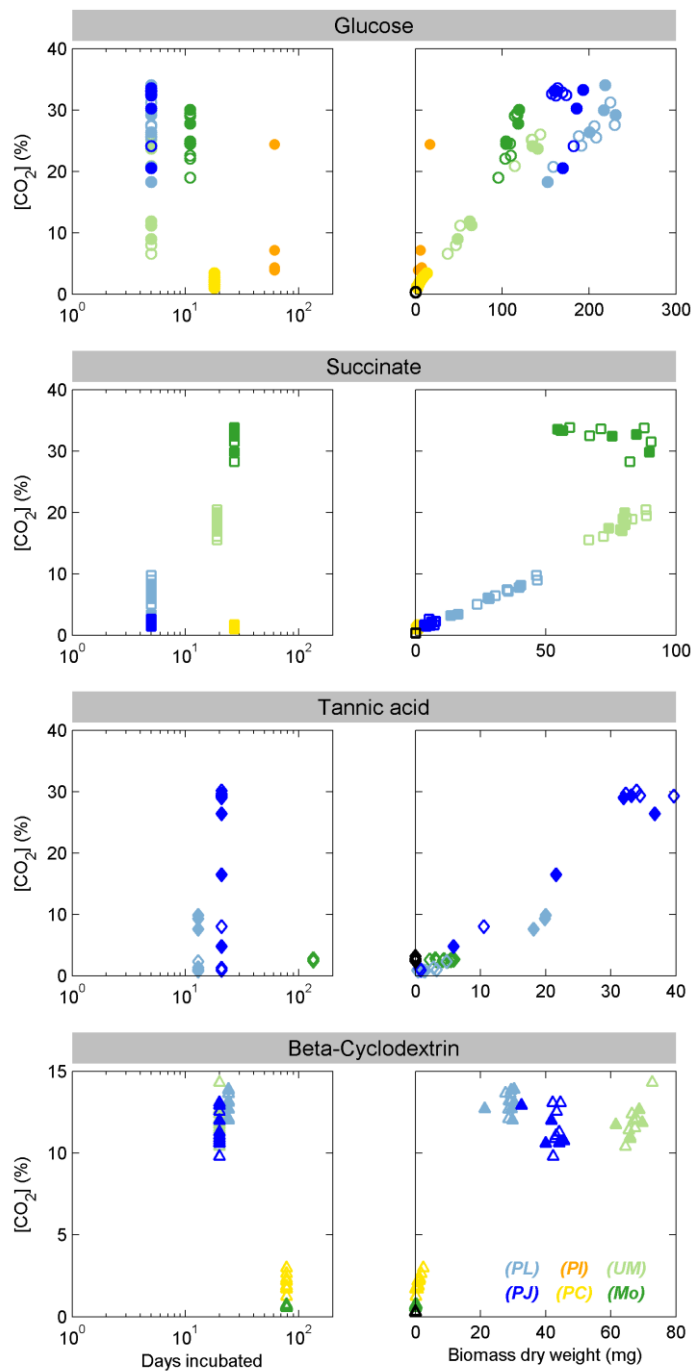




Figure 1. Growth of fungal species on each substrate as indicated by production of CO<sub>2</sub> versus days of incubation (left panels) or dry biomass (right panels). Filled symbols indicate samples for which the C<sub>18:2</sub> biomarker was measured by Pyr-GC-IRMS. Colors represent the Ascomycetes species *Penicillium janczewskii* (PJ<sub>2</sub> (dark blue) and *Paecilomyces lilacinus* (PL<sub>2</sub> (light blue), Zygomycetes *Mortierella* sp. (species-MO<sub>2</sub> (dark green) and *Umbelopsis* sp. (UM<sub>2</sub> (light green), and Basidiomycetes species *Phanerochaete chrysosporium* (PC<sub>2</sub> (orange) and *Paxillus involutus* (PI<sub>2</sub> (yellow). The symbols denote incubations with glucose (circles), succinate (squares), tannic acid (diamonds), or  $\beta$ -cyclodextrin (triangles).

The growth substrates induced a wide range in CUE values, ranging from 0.1 to 0.6 (Fig. 2). Higher CUE values were typically observed for Ascomycetes and Zygomycetes species growing on glucose, and lower values for their growth on succinate and tannic acid. CUE estimated for Basidiomycetes species was always low (< 0.15). The CUE range for growth on glucose (0.1-0.6),  $\beta$ -cyclodextrin (0.1-0.6), and succinate (0.2-0.5) were larger than observed for tannic acid (0.15-0.3).

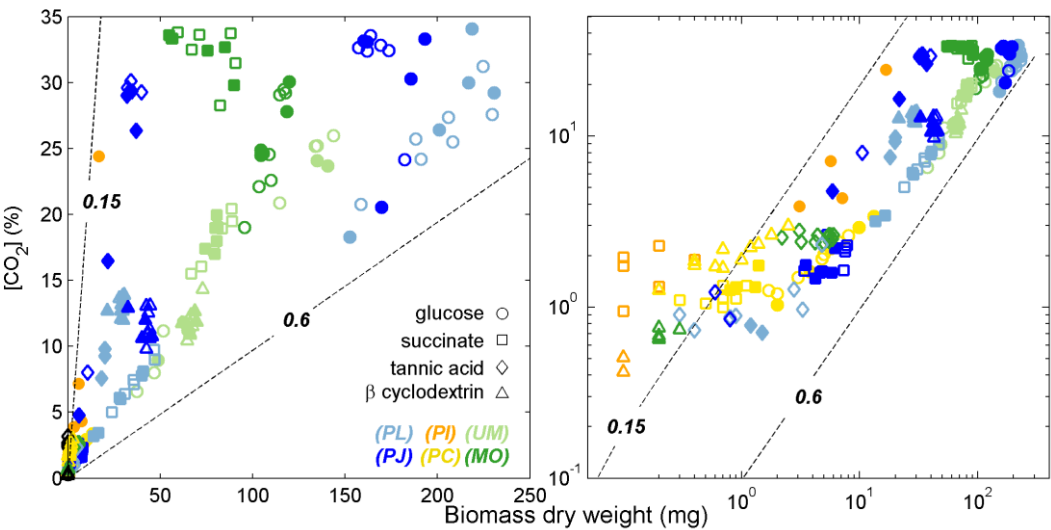


Figure 2. Accumulation of biomass and headspace CO<sub>2</sub> for fungal species from three phyla growing on monomers or complex substrates. The right panel includes the same data on a log-log scale to depict the trends of fungi exhibiting minimal growth. Lines indicate CUE trajectories of 0.15 or 0.6, and were calculated assuming that fungal biomass was 44% C (w/w). Colors and symbols are redundant with Fig. 1. Filled symbols indicate samples that were analyzed by Pyr-GC-IRMS.

Fungal respiration of the growth substrates led to decreasing  $\delta^{13}\text{C}$ -CO<sub>2</sub> values as fungal biomass was produced, which followed a hyperbolic trend expected for the mixing of CO<sub>2</sub> from two different sources (Text S1; e.f., Kendall and Caldwell, 1998).



The atom %  $^{13}\text{C}$  in control incubations with no fungal inoculum was measured at the latest time of harvest of inoculated incubations and stayed below 0.4% except tannic acid which ranged from 2–3%; the  $\delta^{13}\text{C}$  values of the substrates were glucose = -26.5‰; 194 succinate = -28.3‰, tannic acid = -27.4‰,  $\beta$ -cyclodextrin = -10.6‰. The mixing relationship was modeled using all  $\text{CO}_2$  data, 195 across all incubations, and integrated to approximate the mixing-weighted average  $F^{13}\text{C}$  value of inorganic C for each incubation (cf., Text S1, Fig. S2), which was finally applied in the denominator of Eq. 1 to estimate the fraction of lipid C derived from inorganic C. For incubations that produced sufficient fungal biomass for stable C isotopic analysis, the weighted 198 average  $\delta^{13}\text{C}$  values of inorganic C that were applied in Eq. 1 ranged from 200 to 1400 ‰ (i.e., ~1.3 to 2.6 AT%  $^{13}\text{C}$ ).

### 3.2 Stable isotopic composition of fungal lipids

Fungal respiration of the different (unlabeled) growth substrates led to decreasing  $\delta^{13}\text{C}$ - $\text{CO}_2$  values as fungal biomass was produced, which followed a hyperbolic trend expected for the mixing of  $\text{CO}_2$  from two different sources (Text S1; c.f., Kendall and Caldwell, 1998). The atom %  $^{13}\text{C}$  in control incubations with no fungal inoculum was measured at the latest time of harvest of inoculated incubations and stayed below 0.4%, except for incubations with tannic acid, where it ~~ie~~ ranged between from 22% and -3%; the  $\delta^{13}\text{C}$  values of the substrates were glucose = -26.5‰; succinate = -28.3‰, tannic acid = -27.4‰,  $\beta$ -cyclodextrin = -10.6‰. The mixing relationship was modeled using all  $\text{CO}_2$  data, across all incubations, and integrated to approximate the mixing-weighted average  $^{13}\text{F}$  value of ~~inorganic-IC~~ for each incubation (cf., Text S1, Fig. S2), which was finally applied in the denominator of Eq. 1 to estimate the fraction of lipid-C derived from ~~inorganic-IC~~. For incubations that produced sufficient fungal biomass for stable C isotopic analysis, the weighted average  $\delta^{13}\text{C}$  values of ~~inorganic-IC~~ that were applied in Eq. 1 ranged from 200 to 1400 ‰ (i.e., ~1.3 to 2.6 AT%  $^{13}\text{C}$ ), and was largely dependent on how much of the substrate was respired to  $\text{CO}_2$ .

#### 3.2.1 Carbon isotopes

The  $\delta^{13}\text{C}$  values of fungal biomarker  $\text{C}_{18:2}$  was determined as described in section 2.2.4 and is reported in this section as standard 202 delta values (‰). The inorganic C incorporation into the biomarker was calculated based on the following equation:

$F^{13}\text{C}$

$$\% \text{IC}_{(\text{assimilation})} = \frac{F^{13}\text{C}_{\text{lipidsCDIC}} - (\text{labelingmedium}) - F^{13}\text{C}_{\text{lipidssubstrate}} - \text{control}}{F^{13}\text{C}_{\text{lipidsCDIC}} - (\text{labelingmedium}) - F^{13}\text{C}_{\text{lipidssubstrate}} - \text{control}} \times 100\%$$

(Eq. 1)

**Equation 1:** Inorganic carbon (IC) assimilation was calculated as the difference in the  $^{13}\text{C}$  atom fraction ( $F^{13}\text{C}$ ) of the lipids from the 205 labeling experiment compared to the natural (control), relative to the difference between the mixing-weighted average  $F^{13}\text{C}$  of 206 dissolved inorganic C (DIC, cf. Text S1) and the  $F^{13}\text{C}$  of the substrate. F was calculated as  $F^{13}\text{C} = (R^{13}\text{C}/42\text{C}) / (R^{13}\text{C}/42\text{C} + 1)$ , where



$R$  is calculated from the  $\delta^{13}\text{C}$  ratios as measured with the IRMS equipment using the reverse of the  $\delta$  notations ( $\delta^{13}\text{C} = 208$   
 $(\frac{^{13}\text{C}/^{12}\text{C}_{\text{sample}}}{^{13}\text{C}/^{12}\text{C}_{\text{ref}}} - 1) * 1000$  (modified after Boshker & Middelburg 2002; Wegener et al., 2012).

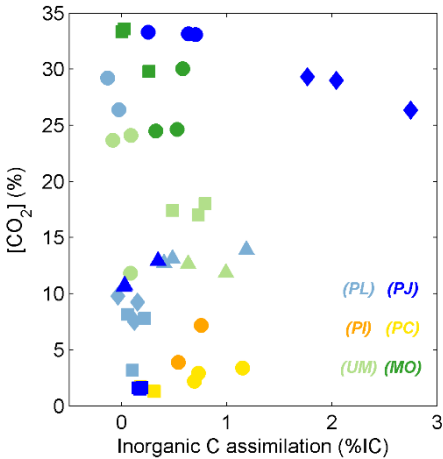
The  $\delta^{13}\text{C}$  values of fungal biomarkers fatty acid  $\text{C}_{18:2}$  (Table 1) produced under natural cultivation conditions with glucose (i.e., non-labeled;  $\text{AT}\%_{\text{DIC}} \sim 1\%$ ) ranged from  $-24.1\%$  to  $-21.2\%$  across all strains ( $n = 6$  species). As expected,  $\text{C}_{18:2}$  harvested from the labeled incubations exhibited slightly higher  $\delta^{13}\text{C}$  values (up to  $+11\%$ ; PC grown on glucose) than the corresponding experiment amended with natural bicarbonate, likely owing to the incorporation of labeled inorganic  $\text{C}$  into the  $\text{C}_{18:2}$  fatty acid.

**Table 1:  $\delta^{13}\text{C}$  values of fungal biomarker  $\text{C}_{18:2}$  harvested from incubations with non-labeled substrates (nat) or those amended with**

$^{13}\text{C}$ -labeled bicarbonate. Incorporation of inorganic  $\text{C}$  (%IC) was calculated based on Eq.1. Errors represent the standard deviation of replicate incubations. Not all fungal species grew sufficiently on all substrates, and thus some did not give enough biomass for analysis (n.d.) and therefore no inorganic  $\text{C}$  incorporation was calculated. %IC could not have been was not determined (n.d.). ~~Errors represent the standard deviation of replicate incubations.~~

Species	Glucose $\delta^{13}\text{C}$ (‰)		IC (%)	Succinate $\delta^{13}\text{C}$ (‰)		IC (%)	Tannic acid $\delta^{13}\text{C}$ (‰)		IC (%)	$\beta$ -cyclodextrin $\delta^{13}\text{C}$ (‰)		IC (%)
	nat	+		nat	+		nat	+		nat	+	
<i>Paxillus involutus</i> (PI)	-21.5	-16.7	0.6 ( $\pm 0.2$ )	n.d	n.d	n.d	n.d	n.d	n.d	n.d	n.d	n.d
<i>Phanerodontia chrysosporium</i> (PC)	-24.1	-15.9	0.9 ( $\pm 0.3$ )	-27.3	-23.5	0.3 ( $\pm 0.1$ )	n.d	n.d	n.d	n.d	n.d	n.d
<i>Mortierella</i> (MO)	-21.9	-20.7	0.5 ( $\pm 0.1$ )	-31.7	-31.5	0.1 ( $\pm 0.1$ )	n.d	n.d	n.d	n.d	n.d	n.d
<i>Umbelopsis</i> (UM)	-21.2	-21.1	0.1 ( $\pm 0.0$ )	-30.6	-28.2	0.7 ( $\pm 0.2$ )	n.d	n.d	n.d	-21.7	-18.1	0.8 ( $\pm 0.3$ )
<i>Penicillium janczewskii</i> (PJ)	-23.2	-22.1	0.5 ( $\pm 0.2$ )	-30.1	-27.8	0.2 ( $\pm 0.0$ )	-25.9	-20.5	2.2 ( $\pm 0.5$ )	-20.7	-20.1	0.1 ( $\pm 0.2$ )
<i>Paecilomyces lilacinus</i> (PL)	-23.0	-23.5	n.d	-30.8	-29.9	0.1 ( $\pm 0.0$ )	-25.4	-25.0	0.1 ( $\pm 0.0$ )	-19.1	-17.6	0.7 ( $\pm 0.4$ )

The estimated incorporation of IC into  $\text{C}_{18:2}$  (%IC) typically ranged up to 1%; only PJ grown on tannic acid exhibited higher %IC values, which ranged up to 2.2% (Table 1, Fig. 23). There were no general trends observed in %IC with other measured or estimated parameters, including CUE; however, for the two species that were able to grow on tannic acid, %IC was positively correlated with the amount of  $\text{CO}_2$  and biomass produced during the incubation ( $R^2 > 0.85$ ,  $n = 5$ ,  $p < 0.01$ ).



**Figure 32.** %IC values for fungal species respiring glucose (circles), succinate (squares), tannic acid (diamonds), or  $\beta$ -cyclodextrin (triangles). Colors represent individual fungal phyla isolates as described in Fig. 1.

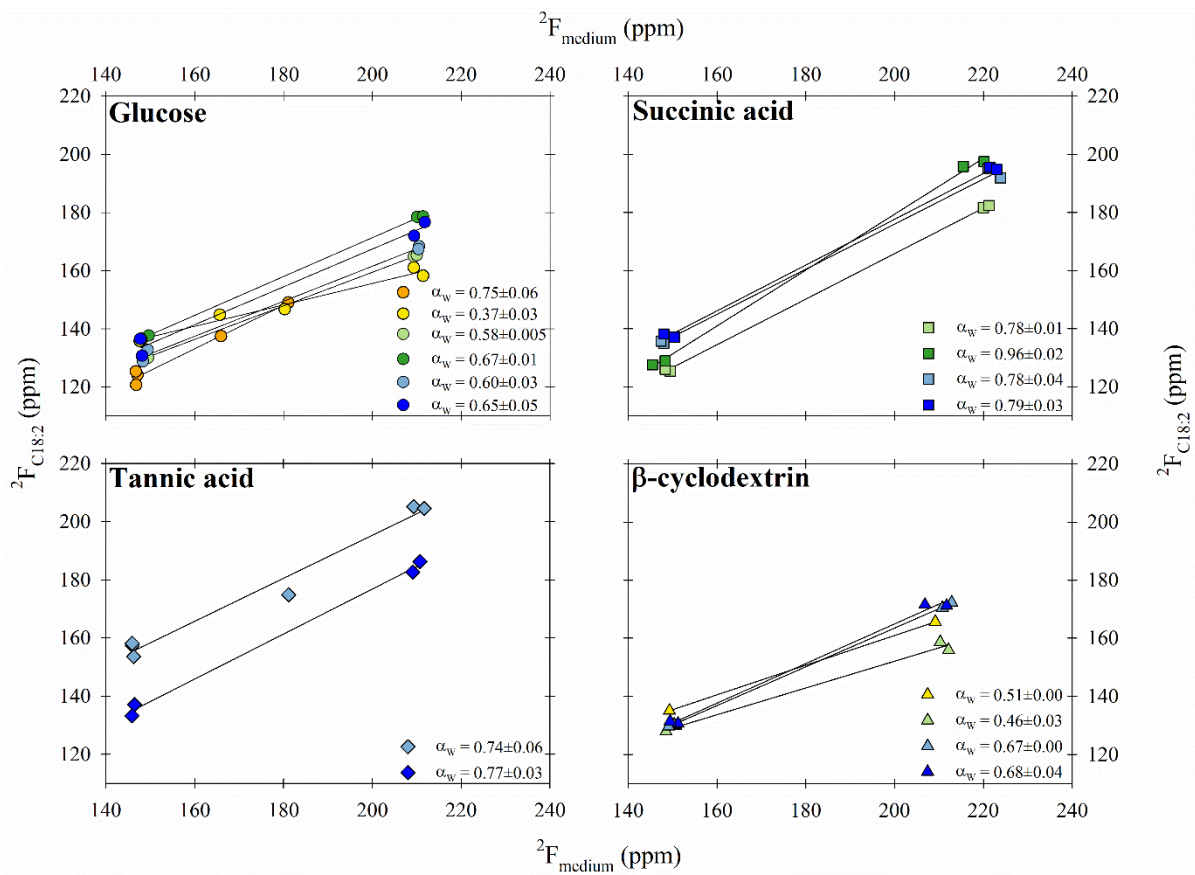
### 3.2.2 Water assimilation factor ( $\alpha_w$ )

The “net” contribution of water hydrogen to lipid H is reported as the water hydrogen assimilation factor  $\alpha_w$  (Kopf et al., 2015), and was estimated based on the slope of the linear regression line between H isotopic composition of lipid versus growth medium water (Fig. 43), which ranged from natural MilliQ ( $\delta^2\text{H} = -45\text{‰} \pm 10\text{‰}$ ) to the labeled treatments ( $65\text{‰} \pm 4\text{‰}$ ;  $166\text{‰} \pm 10\text{‰}$ ;  $368\text{‰} \pm 27\text{‰}$ ). The  $\alpha_w$  values for the fungal biomarker  $\text{C}_{18:2}$  grown on glucose ranged from  $0.37 \pm 0.03$  to  $0.75 \pm 0.06$  with an average value of  $0.60 \pm 0.05$  ( $n = 6$  species;  $\pm\text{SEM}$ ). When grown on succinic acid, the  $\alpha_w$  values for  $\text{C}_{18:2}$  harvested from individual species ranged from  $0.78 \pm 0.01$  to  $0.96 \pm 0.02$  with an average value of  $0.83 \pm 0.04$  ( $n = 4$  species;  $\pm\text{SEM}$ ). When grown on tannic acid, the  $\alpha_w$  values for  $\text{C}_{18:2}$  harvested from individual species ranged from  $0.74 \pm 0.06$  to  $0.77 \pm 0.03$ , and when grown on  $\beta$ -cyclodextrin the  $\alpha_w$  values for  $\text{C}_{18:2}$  ranged from  $0.46 \pm 0.03$  to  $0.68 \pm 0.04$  with an average value of  $0.58 \pm 0.06$  ( $n = 4$  species;  $\pm\text{SEM}$ ). The average  $\alpha_w$  values for  $\text{C}_{18:2}$  for all substrates and species was  $0.67 \pm 0.04$  ( $\pm\text{SEM}$ ).

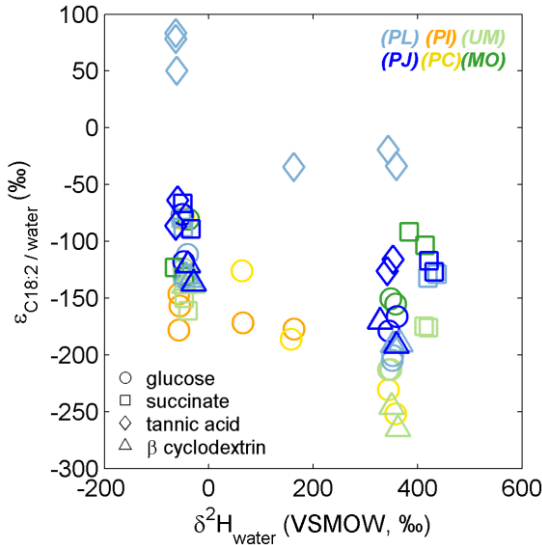
The range of traditionally reported isotope effects  $\alpha_{\text{C}_{18:2}/\text{water}}$  and  $\varepsilon_{\text{C}_{18:2}/\text{water}}$  (Sessions and Hayes, 2005) for all natural and  $^2\text{H}$ -labeled fungal growth experiments was 0.73 to 1.08 and -265 to +83 ‰, respectively. PL growth on tannic acid exhibited the highest values (0.97 to 1.08 and -35 to +83 ‰, respectively; Fig. 5, Table S1), while all other growth experiments  $\alpha_{\text{C}_{18:2}/\text{water}}$  and  $\varepsilon_{\text{C}_{18:2}/\text{water}}$  remained  $< 0.94$  and  $-65$  ‰, respectively. The average ( $\pm\text{SD}$ )  $\varepsilon_{\text{C}_{18:2}/\text{water}}$  values were not significantly different for



fungi growing on glucose ( $-151 \pm 121$  ‰,  $n = 6$ ), succinate ( $-121 \pm 44$  ‰,  $n = 4$ ), tannic acid ( $-39 \pm 63$  ‰,  $n = 2$ ), or  $\beta$ -cyclodextrin ( $-171 \pm 82$  ‰,  $n = 3$ ).



**Figure 43.** The water hydrogen assimilation factor ( $\alpha_w$  values) estimated as the slope of the fractional  $^{21}\text{H}$  abundance ( $2F$ ) in lipids ( $^{21}\text{F}_{\text{C18:2}}$ ) versus medium water ( $^{21}\text{F}_{\text{medium}}$ ). Data are shown for fungal biomarker  $\text{C}_{18:2}$  produced during growth on the different substrates (glucose, succinic acid, tannic acid and  $\beta$ -cyclodextrin) and harvested from the different fungal isolates/species [*Paxillus involutus* (PI), *Phanerochaete chrysosporium* (PC), *Mortierella* sp. (MO), *Umbelopsis* sp. (UM), *Penicillium janczewskii* (PJ), and *Paecilomyces lilacinus* (PL)].  $R^2$  values for all slopes were  $> 0.97$ .



**Figure 5. The apparent isotope effect ( $\epsilon_{C18:2/water}$ ) for fungi grown in medium having variable  $\delta^2H_{water}$  composition. The colors and symbols are redundant with Fig. 1. For each species-substrate pair, the large range and/or decrease in  $\epsilon_{C18:2/water}$  in  $^2H$ -enriched medium are expected, given the additional contributions of substrate-H and metabolic water-H to lipid-H during biosynthesis.**

## 4 Discussion

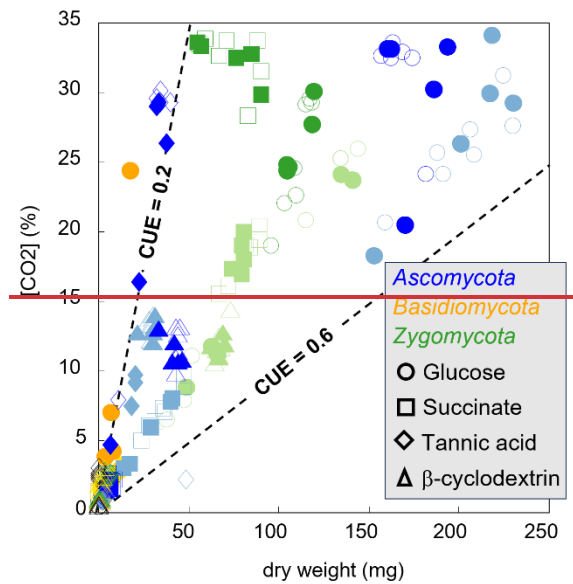
### 4.1 Fungal growth dynamics

Collectively, the fungal incubation experiments included ~~a total of~~ six species representing three different phyla growing on one of four substrates, and exhibited a large range in the relative amounts of  $CO_2$  (0.2-34% v/v) and biomass produced (0-230 mg dry weight; Fig. 1), with ~~the estimated carbon-use efficiency~~  $[CUE = \text{biomassC} / (\text{CO}_2 + \text{biomassC})]$  CUE ranging from 0.2-15 to 0.6 (Fig. 24). ~~The incubations were initiated under atmospheric, oxic conditions, such that fungi were able to respire the substrate aerobically.~~ While atmospheric, oxic conditions likely prevailed during most of the incubation period, it is probable that some incubations turned anoxic when  $CO_2$  levels exceeded 21%, which occurred ~~primarily~~ in incubations of Ascomycota ~~et~~ et ~~Zygomycetes~~ mycota ~~growing on with~~ growing on ~~glucose, Ascomycota~~ growing on ~~tannic acid, and/or Zygomycetes~~ growing on succinate ~~Mortierella~~. ~~The accumulation of  $CO_2$  is~~ an unintended ~~necessary~~ consequence of performing the incubations in closed bottles, which was required to prevent the escape of  $^{13}C$ -labeled Inorganic-C and also to prevent microbial contaminations. Nevertheless, such alteration between oxic and anoxic conditions is common in natural environments, and the measured inorganic-IC assimilation into fungal lipids was consistently low ( $< 3\%$ ; Fig. 23), regardless the implied anoxia.





Furthermore, the observed variability in CUE, %IC, and  $a_w$  between growth experiments were not correlated with large increases in headspace  $\text{CO}_2$ .



**Figure 4. Carbon use efficiency of fungal species from three phyla growing on monomers or complex substrates. Lines were calculated assuming that fungal biomass was 44% C (w/w). Colors and symbols are redundant with Fig. 1.**

#### 4.2 Fungal IC assimilation into lipids

A fundamental process in nature and basis for ecological food webs is the fixation of inorganic-IC via photosynthesis and/or chemosynthesis by autotrophic organisms. Inorganic-The IC assimilation by heterotrophic organisms also plays an important role in ensuring the provision of energy and to replenish intermediates in the TCA cycle that have been released for biosynthesis (Kornberg 1965). Therefore, inorganic-IC assimilation is a measure-proxy for of both anabolic processes and the catabolic status of the cell, influenced by assimilation, biosynthesis, anaplerotic reactions, and redox balancing reactions (Braun et al., 2021; Erb 2011). Previous reports on the by-fixation of inorganic-IC (%IC) via anaplerotic pathways into heterotrophic biomass varied between 1% and —8% (Dijkhuizen & Harder, 1985; Feisthauer et al., 2008; Romanenko 1964; Roslev et al., 2004), whereas for fungi it was previously reported to amount to roughly 1% (Sorokin 1961; Schinner & Concini, 1981; Schinner et al., 1982), and was recently shown to vary between 2% and —12% for Ascomycetesmycota when grown on glucose or glutamic acid (Jabinski et al., 2024). Our results, focusing on a specific fatty acid biomarker, demonstrate a low range in %IC for all different substrates and species tested in this study (0 - 3%), with the Ascomycetesmycota (0 - 2%) assimilating relatively less inorganic-C than previously reported species ( $4.6\% \pm 1.6\%$ ; Jabinski et al.,



2024). The highest observed incorporation was  $2.2 \pm 0.58\%$  by *Penicillium janczewskii* (PJ,  $n = 3$ ) when grown for 21 days on tannic acid (Table 1; Fig. 343). Only, not the other Ascomycota species, *Paecilomyces lilacinus* (PL), grew sufficiently on tannic acid; notably, this experiment yielded high production of CO<sub>2</sub> and biomass, (up to 10% CO<sub>2</sub> and 20 mg dry weight after 13 days; %IC =  $0.14 \pm 0.02\%$ ,  $n = 3$ ; Figs. 1, 3), suggesting that increased assimilation of inorganic-IC by PJ may promote the ability to respire the complex substrate have occurred during the extra week of incubation and promoted higher biomass production. The high CO<sub>2</sub> levels also suggest that the incubations of PJ with tannic acid may have turned anoxic, which may also explain the higher inorganic-IC incorporation in these incubations. Overall, heterotrophic IC assimilation does not appear to be a hallmark of any of the variety of fungal taxa or catabolic pathways probed in this study.

### 4.3 Water hydrogen derived incorporation into fungal lipid biomarker C18:2 Lipid synthesis

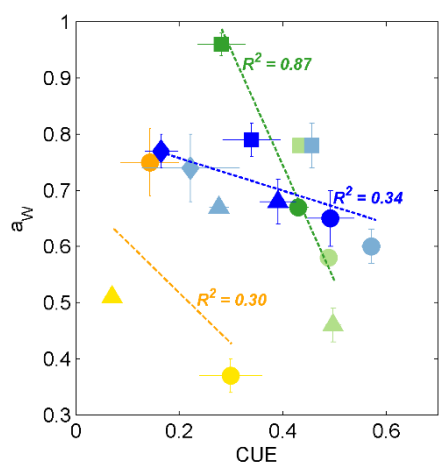
As demonstrated previously, the regression slope between hydrogen isotopic composition of water medium and microbial lipids (i.e.,  $\epsilon_{aW}$ ) varies with the type of metabolism (Zhang et al., 2009; Valentine, 2009; Wijker et al., 2019; Jabinski et al., 2024). For fatty acid biosynthesis, H incorporation is suggested to be a function of transporters and electron acceptors (NADPH and NADH), with contributions accounting for around half of all lipid hydrogen (Maloney et al., 2024). The remaining comprises equal contributions of H obtained directly from environmental water or acetyl-CoA (Valentine, 2009; Zhang et al., 2009; Caro et al., 2023). The consensus from previous studies that investigated the lipids of heterotrophic bacteria is that microbial heterotrophs exhibit  $\epsilon_{aW}$  values ranging from 0 to 1.2, with a mean of  $0.71 \pm 0.17$  (e.g., Caro et al., 2023), though some organisms have exhibited  $a_W$  values exceeding 1 (Dirghangi et al., 2013; Jabinski et al., 2024). Jabinski et al. (2024) demonstrated that five species of heterotrophic Ascomycetes mycota exhibit similar  $\epsilon_{aW}$  values ( $0.62 \pm 0.04$ ) for the fungal biomarker C<sub>18:2</sub> during growth on glucose. Zhang et al. (2009) demonstrated reported similar  $a_W$  values for with *E. coli* grown on glucose  $a_W$  values of a similar range ( $0.63 \pm 0.03$ ). In the current study,  $\epsilon_{aW}$  values for the fungal biomarker C<sub>18:2</sub> during growth on glucose ( $0.60 \pm 0.05$ ) were agreeable with Jabinski et al. (2024), but more variable, likely owing to the broader phylogenetic coverage of the current study. The similarly large variability in  $\epsilon_{C18:2/water}$  values can be partly attributed to the large range in  $\delta^2H$  of medium water, which contributes H together with the substrate and metabolic water to determine  $\delta^2H_{C18:2}$ , even though the net isotope effect ( $\alpha_{C18:2/water}$ ) may be consistent for a specific species-substrate pair (Fig. 5; Session and Hayes, 2005; Kopf et al., 2015). In other words,  $\delta^2H_{C18:2}$  of heterotrophic fungi exhibits more inertia than  $\delta^2H_{H_2O}$  in highly labeled incubations, yielding lower  $\epsilon_{C18:2/water}$  values compared to natural abundance incubations. However, More robust, significant differences in  $\epsilon_{aW}$  values were observed in  $a_W$  values both between and within the different phyla and substrates tested.

#### 4.3.1 Trends across fungal phyla

Ascomycetes mycota species exhibited the most consistent  $\epsilon_{aW}$  values among phyla when grown on each of the four different substrates [ $0.63 \pm 0.03$  (glucose Glu);  $0.78 \pm 0.01$  (succinate SA);  $0.76 \pm 0.02$  (tannic acid TA);  $0.67 \pm 0.01$  ( $\beta$ -cyclodextrin BC)], but also the largest variability in CUE (0.08-0.59; Fig. 2). CUE and  $a_W$  were not significantly correlated



across incubations of Ascomycetesmycota, suggesting that drastic changes in the central metabolic pathways that convert C substrates to either biomass or energy reserves did not systematically alter the net water-H incorporation into lipids. This in contrast to the prevailing notion that  $a_w$  values respond to changes in NADPH production and turnover within the cell (e.g. Wijker et al., 2019). Basidiomycetesmycota only produced sufficient biomass when fed Basidiomycetesmycota-only produced enough biomass for isotopic analysis when grown on substrates that activated the glycolytic pathway (-glucose and/or  $\beta$ -cyclodextrin;  $CUE < 0.3$ ), yet and showed high variability in  $a_w$  between species ( $0.37 \pm 0.03 < a_w < 0.75 \pm 0.06$ ; Fig. 4). For example, during growth on glucose, *P. involutus* exhibited much higher  $a_w$  values than *P. chrysosporium* ( $0.75 \pm 0.06$  versus  $0.37 \pm 0.03$ , respectively), and both of these values which were beyond the more confined range of  $a_w$  values determined for isolates belonging to Zygomycetesmycota and Ascomycetesmycota species. For incubations in which the investigated Zygomycetesmycota strains produced enough biomass to determine  $a_w$  (i.e., glucose, succinate,  $\beta$ -cyclodextrin;  $n = 5$ ; Fig. 4), we observed a highly significant inverse correlation with CUE ( $R^2 = 0.87$ ,  $p < 0.01$ ; Fig. 6), suggesting the potential for  $a_w$  to serve as a proxy for growth efficiency.



**Figure 6. Biplot of CUE and  $a_w$  values for fungi growing on glucose (circles), succinate (squares), tannic acid (diamonds), or  $\beta$ -cyclodextrin (triangles). The colors are redundant with Fig. 1, representing the phyla Ascomycetesmycota (blue shades,  $n = 8$ ), Zygomycetesmycota (green shades,  $n = 5$ ), and Basidiomycetesmycota (yellow and orange symbols,  $n = 3$ ). The dashed lines and  $R^2$  values indicate the linear regression for the corresponding phylum across all substrates that yielded sufficient biomass. Only the regression for Zygomycetesmycota was significant ( $p < 0.01$ ).**

Growth on  $\beta$ -cyclodextrin, which consists of seven glucanopyranose units ( $C_6H_{12}O_6$ ), exhibited similar  $a_w$  values ( $0.58 \pm 0.06$ ) as growth on glucose ( $0.60 \pm 0.05$ ), suggesting that the catabolism of glucose subunits via glycolysis overprints signals of water-H incorporation that may derive during degradation of the  $\beta$ -cyclodextrin oligomer. Succinate yielded significantly higher  $a_w$  values ( $0.83 \pm 0.05$ ), which was in the same range as reported for *E. coli* when grown on succinate ( $a_w$   $0.80 \pm 0.05$ ;



Zhang et al., 2009) and was more similar to that reported previously for fungal growth on glutamic acid ( $0.90 \pm 0.07$ ; Jabinski et al., 2024).

Considering all fungal incubations, a one-way analysis of variance (ANOVA; Holm-Sidak method; SigmaPlot v11) confirmed the significant difference in  $a_w$  values between growth on glucose and glutamic acid ( $p < 0.001$ ), glutamic acid and  $\beta$ -cyclodextrin ( $p < 0.001$ ), succinate and glucose ( $p < 0.003$ ), and succinate and  $\beta$ -cyclodextrin ( $p < 0.005$ ). It also confirmed that there was no significant difference between the other substrate combinations ( $p > 0.005$ ). Notably,

#### 4.3.1 Trends across C substrates

Across all incubations, the similar  $a_w$  values determined for growth on glucose ( $0.60 \pm 0.05$ ) versus  $\beta$ -cyclodextrin ( $0.58 \pm 0.06$ ), of which the latter which consists of seven glucanopyranose units ( $C_6H_{12}O_6$ ), suggests that the catabolism of glucose subunits via glycolysis overprints signals of water-H incorporation that may derive during degradation of the  $\beta$ -cyclodextrin oligomer. Alternatively to glycolysis, succinate yielded significantly higher  $a_w$  values ( $0.83 \pm 0.05$ ), which was in the same range as reported for *E. coli* when grown on succinate ( $a_w 0.80 \pm 0.05$ ; Zhang et al., 2009), and was more similar to that reported previously for fungal growth on glutamic acid ( $0.90 \pm 0.07$ ; Jabinski et al., 2024). Considering all fungal incubations, a one-way analysis of variance (ANOVA; Holm-Sidak method; SigmaPlot v11) confirmed the significant difference in  $a_w$  values between growth on glucose and glutamic acid ( $p < 0.001$ ), glutamic acid and  $\beta$ -cyclodextrin ( $p < 0.001$ ), succinate and glucose ( $p < 0.003$ ), and succinate and  $\beta$ -cyclodextrin ( $p < 0.005$ ). It also confirmed that there was no significant difference between the other substrate combinations ( $p > 0.05$ ).

Glutamic acid and succinate are thought to be introduced into the TCA cycle through coupled metabolites, where succinate is a direct metabolite inside-of the TCA cycle and glutamic acid is converted to  $\alpha$ -ketoglutarate intermediate by transamination before entering the TCA cycle, which is only 2 steps from succinate (Cooper et al., 2014). Also, being acids, these substrates may have a greater capacity than saccharides to exchange H with ambient water at experimental pH (typically  $2 < \text{pH} < 5.2$ ), especially glutamic acid, which also comprises an amino moiety. Tannic acid ( $0.76 \pm 0.02$ ) yielded no significant differences ( $p > 0.005$ ) from the other substrates, and is reported to be degraded to different subunits including gallic acid and glucose (Banerjee and Mahapatra, 2012; Lekha and Lonsane, 1997 and references within). Aromatic degradation pathways employed by fungi generate intermediates that go through the  $\beta$ -ketoadipate pathway (Mäkelä et al., 2015) before entering the TCA cycle as a succinyl-CoA metabolite (Lekha and Lonsane, 1997). The  $a_w$  values induced by degradation of TA-tannic acid suggest that it integrates both the lower  $a_w$  signature of glycolysis and higher  $a_w$  signature of the TCA cycle (Fig. 57). Similarly to trends in  $a_w$ , the  $\epsilon_{C18:2/\text{water}}$  values estimated for fungal growth on succinate or tannic acid were typically higher than glucose or  $\beta$ -cyclodextrin, for a given  $\delta^2\text{H}_{\text{H}_2\text{O}}$  treatment (Fig. 5; Table S1).

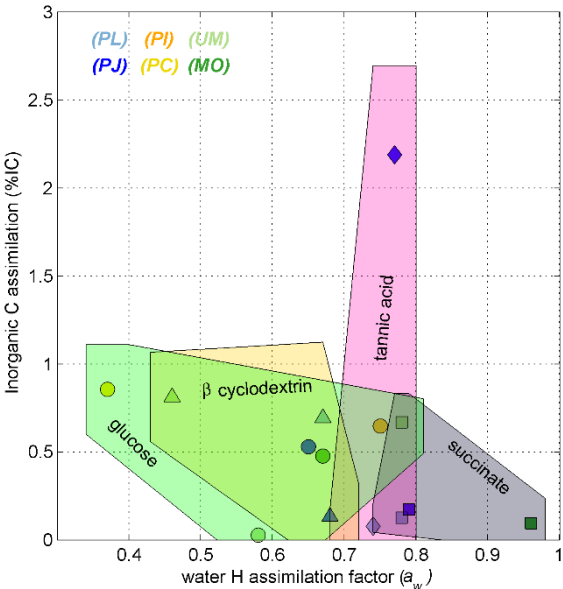
Together, our incubation experiments suggest that  $a_w$  values determined for the fungal biomarker  $C_{18:2}$  could not distinguish between fungal growth on relatively labile monomers (i.e., glucose and succinate; requiring as few as 5 days of cultivation) versus larger, less-labile substrates (i.e.,  $\beta$ -cyclodextrin and tannic acid; requiring 20 to 183 days of cultivation). However, with the exception of Basidiomycetes mycota,  $a_w$  values of fungal lipid biomarkers may be indicative of fungi employing



primarily glycolytic or TCA pathways. Environmental assays that quantify fungal lipid production via the incorporation of ambient water-H (i.e., the lipid-SIP approach) may upscale to total production estimates by applying our calculated mean  $a_w$  value of  $0.69 \pm 0.03$  [ $n = 27$ ;  $\pm$  (SEM)], which is consistent with the  $a_w$  value of 0.71 recommended for soil microbial communities ~~previously~~ (Caro et al., 2023). For  $^2\text{H}$ -SIP investigation of fungal ecotypes supplied with TCA metabolites, such as mycorrhiza, the  $a_w$  value of  $\text{C}_{18:2}$  may range up to  $0.83 \pm 0.05$ . Similarly, such analyses Similar approaches could be applied to environmental samples, such that the  $a_w$  values of phospholipids containing  $\text{C}_{18:2}$  fatty acids ~~could be used to reconstruct~~ inform the distribution of predominant metabolic ecotypes across a soil profile. ~~preferred C sources of saprotrophic microbes under variable environmental conditions (ample water vs. drought) — either at geological time scales, or (coupled with widespread environmental sequencing analyses of microbial communities) addressing metabolic processes in recent biological samples.~~

#### 4.4 Dual-SIP approach

Dual-SIP experiments with  $^2\text{H}_2\text{O}$  and  $^{13}\text{C}$ -dissolved IC previously highlighted the potential to track microbial activity and distinguish heterotrophic vs autotrophic metabolic modes within environmental settings and pure cultures (Kellerman et al., 2012, 2016; Wegener et al., 2012; Huguet et al., 2017; Wu et al., 2018, 2020). This approach was also previously applied to investigate fungal pure cultures (Jabinski et al., 2024), in which the plot of assimilation of ~~inorganic~~-IC versus water-H into the fungal biomarker  $\text{C}_{18:2}$  could distinguish five Ascomycetes ~~mycota~~ species growing on glucose or glutamic acid, with  $a_w$  values explaining most of the variability. While calculated IC: $a_w$  are useful to distinguish autotrophic from heterotrophic growth (cf. Wegener et al., 2016), all calculated values in this study remained near zero, with %IC ranging up to 3% and  $a_w$  values ranging from 0.637 to 0.896 (Fig. 4). This pure culture study therefore suggests that fungal assimilation of ~~inorganic~~ IC is low and less insightful than the more distinguishable  $a_w$  values for identifying the relative contributions of fungal phylotypes or ecotypes in environmental assays.



**Figure S7. Scatterplot of %IC and  $a_w$  values** scatter-plot for  $C_{18:2}$  harvested from incubations of Ascomycetesmycota (blue symbols), Basidiomycetesmycota (yellow and orange symbols), and Zygomycetesmycota (green symbols). The shaded polygons span the range of %IC and  $a_w$  values ( $\pm$  propagated error) for fungal growth on glucose (circles, green shape), succinate (squares, grey shape), tannic acid (diamonds, pink shape), or  $\beta$ -cyclodextrin (triangles, yellow shape). Each data point represents one species-substrate pair determined for  $n > 4$  growth experiments. identifying the grouping of glucose (circles, green shape), succinic acid (squares, grey shape), tannic acid (diamonds, pink shape) and  $\beta$ -cyclodextrin (triangles, yellow shape) incubations with Ascomycetes, Basidiomycetes, and Zygomycetes. Refer to Fig. 1, Caption for further details.

## 5 Conclusion

The purpose of this work was to apply the dual-SIP assay on pure fungal cultures to define the effect of different carbon-organic C substrates on the incorporation of water-H and inorganic-C into their membrane lipids. Although heterotrophic  $CO_2$  fixation by microbes may range up to 8% of biomass C, the inorganic-IC assimilation into the fungal biomarker  $C_{18:2}$  harvested from six species representing Ascomycetesmycota, Basidiomycetesmycota, and Zygomycetesmycota did not vary consistently between species or substrate, and remained below 3%. Our findings suggest that the fungal catabolic pathways activated by the variety of substrates tested in this study cannot fully account for the higher levels of heterotrophic  $CO_2$  incorporation reported in natural systems. However, *Penicillium janczewskii*, the species that was most successful at respiring tannic acid, also exhibited the highest %IC value of all incubations (Fig. 3+; Fig. 7S), suggesting that fungal degradation of similarly



complex substrates may rely in part on the assimilation of inorganic-IC (e.g., via anaplerotic reactions). The use of SIP to estimate %IC of heterotrophs required (i) a closed system to prevent loss of  $^{13}\text{C}$  label to the atmosphere, and (ii) a high label dose, to contend with the accumulation of  $\text{CO}_2$  respired from growth substrate. These conditions intensified upon decreases in pH during the growth experiments, thereby shifting the speciation of Inorganic-C toward  $\text{CO}_2$ , which may have further, yet unknown consequences for anaplerotic incorporation of the IC. Future applications to determine %IC for microbial heterotrophs should consider repeated spiking or continuous cultivation practices to better stabilize pH and  $\delta^{13}\text{C}_{\text{DIC}}$ . Likewise, it would be worth considering isotopic analyses of other potential bioindicator compounds (e.g., sterols, peptides, aminosugars) to see whether incorporation of the stable isotopes IC into such other compounds would render higher levels than the fatty acids.

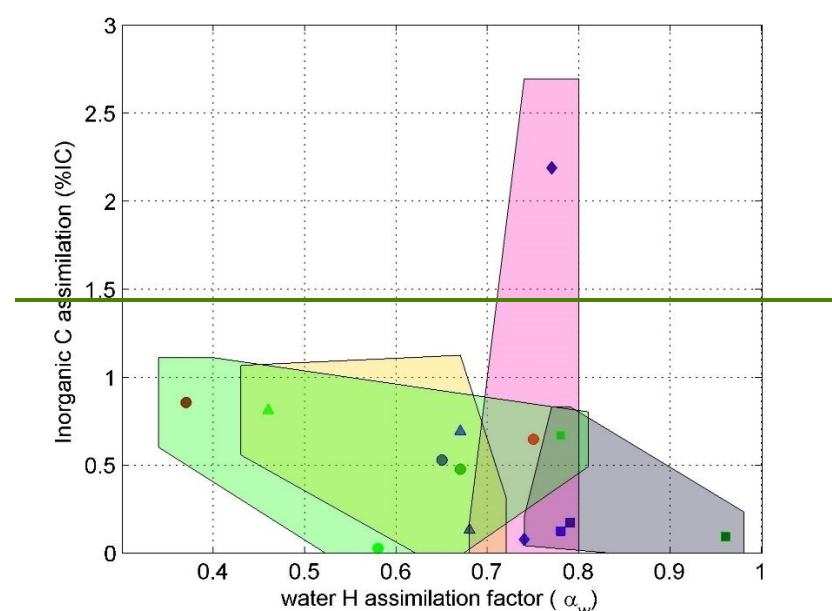


Figure 5. %IC and  $\alpha_w$  value scatter plot for  $\text{C}_{18:2}$  identifying the grouping of glucose (circles, green shape), succinic acid (squares, grey shape), tannic acid (diamonds, pink shape) and  $\square$  cyclodextrin (triangles, yellow shape) incubations with *Ascomycetes*, *Basidiomycetes*, and *Zygomycetes*. Refer to Fig. 1. Caption for further details.

In contrast to %IC, we conclude that substrates that activated the glycolysis pathway yielded significantly lower  $\alpha_w$  values than those catabolized as TCA intermediates. The expanded dataset provided by reported in this study suggests that the accuracy of indicates that inorganic-IC assimilation by heterotrophic fungi accounts for  $< 3\%$  of lipid carbon, and fungal production estimated by  $^2\text{H}$ -lipid SIP experiments can be adjusted-improved by applying the an-average  $\alpha_w$  value of 0.69 for saprotrophic phyte fungi or up to 0.83 for mycorrhizal mycorrhizal fungi, to provide a more accurate estimate of total lipid production. Furthermore, determination of  $\alpha_w$  values in environmental  $^2\text{H}$ -SIP assays may be useful to identify the prevalence





relative contributions of fungal ecotypes that rely on C substrates fueling glycolysis (e.g., leaf litter) versus those that are fed primarily by TCA intermediates (e.g., root or microbial exudates). Lastly, to our knowledge, the two Zygomycetesmycota strains investigated in this study provide the first evidence of a potential correlation between  $a_w$  and CUE (Fig. 6), encouraging further exploration of the link between these two parameters, both of which are coupled to microbial central metabolic pathways.

## Data availability

Data presented in the figures and tables are available in supplementary material (Table S1) and can be obtained by contacting the corresponding author and will be made available on the Fractome Database (<https://fractome.caltech.edu/>).

## Author contribution

Stanislav Jabinski, Conceptualization, Data curation, Formal analysis, Investigation, Methodology, Software, Validation, Visualization, Writing – original draft, Writing – review and editing

Vítězslav Kučera, Investigation, Methodology, Resources, Writing – review and editing

Marek Kopáček, Formal analysis, Methodology, Resources, Validation,

Jan Jansa, Conceptualization, Formal analysis, Methodology, Resources, Validation, Writing – review and editing

Travis B. Meador, Conceptualization, Data curation, Formal analysis, Funding acquisition, Investigation, Methodology, Project administration, Resources, Software, Supervision, Validation, Visualization, Writing – original draft, Writing – review and editing

## Competing interests

The authors declare that they have no conflict of interest.

## Disclaimer Acknowledgements

We thank Ljubov Poláková for the support of the stable isotope measurements and laboratory protocols; the Collection of Microscopic Fungi of the Institute of Soil Biology BC CAS for providing the fungal species *Penicillium janczewskii* strain BCCO20\_0265 and the Institute of Microbiology CAS for providing the fungal species *Paxillus involutus* strain SB-22; *Phanerodontia chrysosporium* strain CCM8074, *Mortierella* strain RK-38; *Umbelopsis* strain RK-43 and *Paecilomyces lilacinus* strain DP-23. This Project was funded by the Czech Science Foundation GACR nr. 20-223805 (FUNSIF) and supported by MEYS CZ grant LM2015075 Projects of Large Infrastructure for Research, Development and Innovations as



548 well as the European Regional Development Fund-Project: research of key soil-water ecosystem interactions at the SoWa  
549 Research Infrastructure (No. CZ.02.1.01/0.0/0.0/16\_013/0001782).

550

551

552

553

554



## References

- Baldrian, P., Voříšková, J., Dobiášová, P., Merhautová, V., Lisá, L., & Valášková, V. ~~(2011)~~. Production of extracellular enzymes and degradation of biopolymers by saprotrophic microfungi from the upper layers of forest soil. *Plant and Soil*, 338(1-2), 111-125. doi:10.1007/s11104-010-0324-3, 2011.
- Banerjee, D., & Mahapatra, S. ~~(2012)~~. Fungal tannase: a journey from strain isolation to enzyme applications. *Dyn Biochem Process Biotechnol Mol Biol*, 6(2), 49-60, 2012.
- Boer, W. D., Folman, L. B., Summerbell, R. C., & Boddy, L. ~~(2005)~~ Living in a fungal world: impact of fungi on soil bacterial niche development. *FEMS microbiology reviews*, 29(4), 795-811, (2005).
- Boschker HTS, Middelburg JJ. ~~2002~~ Stable isotopes and biomarkers in microbial ecology. *FEMS Microbiol Ecol* 40:85–95. <https://doi.org/10.1111/j.1574-6941.2002.tb00940.x> 2002.
- Boschker, H. T. S., Nold, S. C., Wellsbury, P., Bos, D., De Graaf, W., Pel, R., ... & Cappenberg, T. E. ~~(1998)~~. Direct linking of microbial populations to specific biogeochemical processes by <sup>13</sup>C-labelling of biomarkers. *Nature*, 392(6678), 801-805, 1998.
- Braun A, Spona-Friedl M, Avramov M, Elsner M, Baltar F, Reinthaler T, Herndl GJ, Griebler C. ~~2021~~ Reviews and syntheses: heterotrophic fixation of inorganic carbon—significant but invisible flux in environmental carbon cycling. *Biogeosciences* 18:3689–3700. <https://doi.org/10.5194/bg-18-3689-2021>, 2021.
- Bukovská P, Bonkowski M, Konvalinková T, Beskid O, Hujšlová M, Püschel D, ~~et al.~~ Utilization of organic nitrogen by arbuscular mycorrhizal fungi—is there a specific role for protists and ammonia oxidizers? *Mycorrhiza*. 2018;28(3):269–83. <https://doi.org/10.1007/s00572-018-0825-0>, 2018.
- Canarini, A., Fuchslueger, L., Schnecker, J., Metze, D., Nelson, D. B., Kahmen, A., ... & Richter, A. Soil fungi remain active and invest in storage compounds during drought independent of future climate conditions. *Nature Communications*, 15(1), 10410, (2024).
- Carlson C.A., N.R. Bates, D.A. Hansell, D.K. Steinberg. ~~2001~~ Carbon Cycle. In: J. Steele, S. Thorpe, K. Turekian (Eds.) *Encyclopedia of Ocean Science*, 2nd Edition. Academic Press, 477-486, 2001.
- Caro, T. A., McFarlin, J., Jech, S., Fierer, N., & Kopf, S. ~~(2023)~~ Hydrogen stable isotope probing of lipids demonstrates slow rates of microbial growth in soil. *Proceedings of the National Academy of Sciences*, 120(16), e2211625120, (2023).
- Ciais, P., Dolman, A. J., Bombelli, A., Duren, R., Peregon, A., Rayner, P. J., . . . Marland, G. ~~(2014)~~ Current systematic carbon-cycle observations and the need for implementing a policy-relevant carbon observing system. *Biogeosciences*, 11(13), 3547-3602. doi:10.5194/bg-11-3547-2014, (2014).
- Cooper AJL, Kuhara T. ~~2014~~  $\alpha$ -Ketoglutaramate: an overlooked metabolite of glutamine and a biomarker for hepatic encephalopathy and inborn errors of the urea cycle. *Metab Brain Dis* 29:991–1006. <https://doi.org/10.1007/s11011-013-9444-9>, 2014.



- 587 Dijkhuizen, L., & Harder, W. ~~(1985)~~ Microbial metabolism of carbon dioxide. Comprehensive biotechnology: the principles,  
 588 applications, and regulations of biotechnology in industry, agriculture, and medicine/editor-in-chief, Murray Moo-Young,  
 589 ~~(1985)~~.
- 590 Dumont, M. G., & Murrell, J. C. ~~(2005)~~ Stable isotope probing—linking microbial identity to function. Nature Reviews  
 591 Microbiology, 3(6), 499-504. ~~(2005)~~.
- 592 Erb T.J. ~~2011~~ Carboxylases in natural and synthetic microbial pathways. Appl Environ Microbiol 77:8466–8477.  
 593 <https://doi.org/10.1128/AEM.05702-11>. ~~(2011)~~.
- 594 Feisthauer, S., Wick, L. Y., Kästner, M., Kaschabek, S. R., Schlömann, M., & Richnow, H. H. ~~(2008)~~ Differences of  
 595 heterotrophic <sup>13</sup>CO<sub>2</sub> assimilation by Pseudomonas knackmussii strain B13 and Rhodococcus opacus 1CP and potential impact  
 596 on biomarker stable isotope probing. Environmental Microbiology, 10(6), 1641-1651. ~~(2008)~~.
- 597 Fioretto, A., Di Nardo, C., Papa, S., & Fuggi, A. ~~(2005)~~ Lignin and cellulose degradation and nitrogen dynamics during  
 598 decomposition of three leaf litter species in a Mediterranean ecosystem. Soil Biology and Biochemistry, 37(6), 1083-1091.  
 599 doi:10.1016/j.soilbio.2004.11.007. ~~(2005)~~.
- 600 Fischer, C. R., Bowen, B. P., Pan, C., Northen, T. R., & Banfield, J. F. ~~(2013)~~ Stable-isotope probing reveals that hydrogen  
 601 isotope fractionation in proteins and lipids in a microbial community are different and species-specific. ACS chemical biology,  
 602 8(8), 1755-1763. ~~(2013)~~.
- 603 Frey, S. D. ~~(2019)~~ Mycorrhizal fungi as mediators of soil organic matter dynamics. Annual review of ecology, evolution, and  
 604 systematics, 50, 237-259. doi:10.1146/annurev-ecolsys-110617-062331. ~~(2019)~~.
- 605 Grinhut, T., Hadar, Y., & Chen, Y. ~~(2007)~~ Degradation and transformation of humic substances by saprotrophic fungi: processes  
 606 and mechanisms. Fungal biology reviews, 21(4), 179-189. ~~(2007)~~.
- 607 [Hayes, J. M. An introduction to isotopic calculations. Woods Hole Oceanographic Institution, Woods Hole, MA, 2543, 2004.](#)
- 608 Hoefs, J. ~~(2018)~~ Stable isotope geochemistry. Springer International Publishing AG, part of Springer Nature.  
 609 doi:10.1007/9783-319-78527-1. ~~(2018)~~.
- 610 Hoëgberg, P., Nordgren, A., Buchmann, N., Taylor, A. F., Ekblad, A., Hoëgberg, M. N., . . . Read, D. J. ~~(2001)~~ Large-scale  
 611 forest girdling shows that current photosynthesis drives soil respiration. Nature, 411(6839), 789-792. ~~(2001)~~.
- 612 Huguet, A., Meador, T. B., Laggoun-Défarge, F., Köneke, M., Wu, W., Derenne, S., & Hinrichs, K.-U. ~~(2017)~~ Production  
 613 rates of bacterial tetraether lipids and fatty acids in peatland under varying oxygen concentrations. Geochimica et  
 614 Cosmochimica Acta, 203, 103-116. doi:10.1016/j.gca.2017.01.012. ~~(2017)~~.
- 615 Jabinski, S., d. M. Rangel, W., Kopáček, M., Jílková, V., Jansa, J., & Meador, T. B. ~~(2024)~~ Constraining activity and growth  
 616 substrate of fungal decomposers via assimilation patterns of inorganic carbon and water into lipid biomarkers. Applied and  
 617 Environmental Microbiology, 90(4), e02065-23. ~~(2024)~~.



- 618 Kellermann, M. Y., Wegener, G., Elvert, M., Yoshinaga, M. Y., Lin, Y.-S., Holler, T., . . . Hinrichs, K.-U. ~~(2012)~~ Autotrophy as  
619 a predominant mode of carbon fixation in anaerobic methane-oxidizing microbial communities. *Proceedings of the National*  
620 *Academy of Sciences*, 109(47), 19321-19326. doi:10.1073/pnas.1208795109. ~~(2012)~~.
- 621 Kellermann, M. Y., Yoshinaga, M. Y., Wegener, G., Krukenberg, V., & Hinrichs, K.-U. ~~(2016)~~ Tracing the production and fate  
622 of individual archaeal intact polar lipids using stable isotope probing. *Organic Geochemistry*, 95, 13-20.  
623 doi:10.1016/j.orggeochem.2016.02.004. ~~(2016)~~.
- 624 Kirk, T. K., & Farrell, R. L. ~~(1987)~~ "Enzymatic" combustion": the microbial degradation of lignin. *Annual Reviews in*  
625 *Microbiology*, 41(1), 465-501. doi:10.1146/annurev.mi.41.100187.002341. ~~(1987)~~.
- 626 Kopf, S. H., McGlynn, S. E., Green-Saxena, A., Guan, Y., Newman, D. K., & Orphan, V. J. ~~(2015)~~ Heavy water and (15) N  
627 labelling with NanoSIMS analysis reveals growth rate-dependent metabolic heterogeneity in chemostats. *Environ Microbiol*,  
628 17(7), 2542-2556. doi:10.1111/1462-2920.12752. ~~(2015)~~.
- 629 Kornberg HL. ~~1965~~ Anaplerotic sequences in microbial metabolism. *Angew Chem Int Ed Engl* 4:558-565.  
630 https://doi.org/10.1002/anie.196505581. ~~1965~~.
- 631 Kreuzer-Martin, H. W. ~~(2007)~~ Stable isotope probing: linking functional activity to specific members of microbial  
632 communities. *Soil Science Society of America Journal*, 71(2), 611-619. ~~(2007)~~.
- 633 Lekha, P. K., & Lonsane, B. K. ~~(1997)~~ Production and application of tannin acyl hydrolase: state of the art. *Advances in applied*  
634 *microbiology*, 44, 215-260. ~~(1997)~~.
- 635 Lindahl, B. D., & Tunlid, A. ~~(2015)~~ Ectomycorrhizal fungi—potential organic matter decomposers, yet not saprotrophs. *New*  
636 *Phytologist*, 205(4), 1443-1447. doi:10.1111/nph.13201. ~~(2015)~~.
- 637 Maloney, A. E., Kopf, S. H., Zhang, Z., McFarlin, J., Nelson, D. B., Masterson, A. L., & Zhang, X. (2024). Large enrichments  
638 in fatty acid 2H/1H ratios distinguish respiration from aerobic fermentation in yeast *Saccharomyces cerevisiae*. *Proceedings*  
639 *of the National Academy of Sciences*, 121(20), e2310771121, (2024).
- 640 Osburn, M. R., Sessions, A. L., Pepe-Ranney, C., & Spear, J. R. ~~(2011)~~ Hydrogen-isotopic variability in fatty acids from  
641 Yellowstone National Park hot spring microbial communities. *Geochimica et Cosmochimica Acta*, 75(17), 4830-4845. ~~(2011)~~.
- 642 Romanenko VI. ~~1964~~ Heterotrophic assimilation of CO<sub>2</sub> by bacterial flora of water. *Mikrobiologiya* 33:679-683. ~~1964~~.
- 643 Roslev P, Larsen MB, Jørgensen D, Hesselsoe M. ~~2004~~ Use of heterotrophic CO<sub>2</sub> assimilation as a measure of metabolic  
644 activity in planktonic and sessile bacteria. *J Microbiol Methods* 59:381-393. https://doi.org/10.1016/j.mimet.2004.08.002.  
645 ~~2004~~.
- 646 Sachse, D., Billault, I., Bowen, G. J., Chikaraishi, Y., Dawson, T. E., Feakins, S. J., ... & Kahmen, A. ~~(2012)~~ Molecular  
647 paleohydrology: interpreting the hydrogen-isotopic composition of lipid biomarkers from photosynthesizing organisms.  
648 *Annual Review of Earth and Planetary Sciences*, 40, 221-249. ~~(2012)~~.
- 649 Schinner F, Concin R, Binder H. ~~1982~~ Heterotrophic CO<sub>2</sub> fixation by fungi in dependence on the concentration of the carbon  
650 source. Paper presented at the Phyton; annales rei botanicae. ~~1982~~.



- 651 Schinner F, Concin R. ~~1981~~ Carbon dioxide fixation by wood-rotting fungi. Eur J For Pathol 11:120–123.  
652 <https://doi.org/10.1111/j.1439-0329.1981.tb00077.x>. ~~1981~~.
- 653 [Sessions, A.L., Hayes, J.M., 2005. Calculation of hydrogen isotopic fractionations in biogeochemical systems. \*Geochim.\*](#)  
654 [Cosmochim. Acta](#) 69, 593–597.
- 655 Smith, S. E., & Read, D. (~~2008~~) Mineral nutrition, toxic element accumulation and water relations of arbuscular mycorrhizal  
656 plants. Mycorrhizal symbiosis, 3, 145–148. doi:10.1016/B978-012370526-6.50007-6. ~~(2008)~~.
- 657 Šnajdr, J., Cajthaml, T., Valášková, V., Merhautová, V., Petránková, M., Spetz, P., . . . Baldrian, P. (~~2011~~) Transformation of  
658 Quercus petraea litter: successive changes in litter chemistry are reflected in differential enzyme activity and changes in the  
659 microbial community composition. FEMS Microbiology Ecology, 75(2), 291–303. doi:10.1111/j.1574-6941.2010.00999.x.  
660 ~~(2011)~~.
- 661 Sorokin, J. I. (~~1966~~) On the carbon dioxide uptake during the cell synthesis by microorganisms. Zeitschrift für allgemeine  
662 Mikrobiologie, 6(1), 69–73. doi:10.1002/jobm.3630060107. ~~(1966)~~.
- 663 Sorokin, Y. I. (~~1961~~) Heterotrophic carbon dioxide assimilation by microorganisms. Zhurnal Obshchei Biologii, 22(4), 265272.  
664 ~~(1961)~~.
- 665 Treonis, A. M., Ostle, N. J., Stott, A. W., Primrose, R., Grayston, S. J., & Ineson, P. (~~2004~~) Identification of groups of  
666 metabolically-active rhizosphere microorganisms by stable isotope probing of PLFAs. Soil Biology and Biochemistry, 36(3),  
667 533–537. ~~(2004)~~.
- 668 Valentine, D. L. (~~2009~~) Isotopic remembrance of metabolism past. Proceedings of the National Academy of Sciences, 106(31),  
669 12565–12566. doi:10.1073/pnas.0906428106. ~~(2009)~~.
- 670 [Warren, C. R. D2O labelling reveals synthesis of small, water-soluble metabolites in soil. \*Soil Biology and Biochemistry\*, 165,](#)  
671 [108543, 2022.](#)
- 672 Wegener, G., Bausch, M., Holler, T., Thang, N. M., Prieto Mollar, X., Kellermann, M. Y., . . . Boetius, A. (~~2012~~) Assessing  
673 sub-seafloor microbial activity by combined stable isotope probing with deuterated water and <sup>13</sup>C-bicarbonate. Environmental  
674 Microbiology, 14(6), 1517–1527. doi:10.1111/j.1462-2920.2012.02739.x. ~~(2012)~~.
- 675 Wegener, G., Kellermann, M. Y., & Elvert, M. (~~2016~~) Tracking activity and function of microorganisms by stable isotope  
676 probing of membrane lipids. Current Opinion in Biotechnology, 41, 43–52. doi:10.1016/j.copbio.2016.04.022. ~~(2016)~~.
- 677 Wijker, R. S., Sessions, A. L., Fuhrer, T., & Phan, M. (~~2019~~) <sup>2</sup>H/<sup>1</sup>H variation in microbial lipids is controlled by NADPH  
678 metabolism. Proceedings of the National Academy of Sciences, 116(25), 12173–12182. doi:10.1073/pnas.1818372116. ~~(2019)~~.
- 679 Willers, C., Jansen van Rensburg, P. J., & Claassens, S. (~~2015~~) Phospholipid fatty acid profiling of microbial communities—a  
680 review of interpretations and recent applications. Journal of applied microbiology, 119(5), 1207–1218. ~~(2015)~~.
- 681 Wu W, Meador T, Hinrichs K-U. ~~2018~~ Production and turnover of microbial organic matter in surface intertidal sediments.  
682 Org Geochem 121:104–113. <https://doi.org/10.1016/j.orggeochem.2018.04.006>. ~~2018~~.



~~Wu, W., Meador, T. B., Könneke, M., Elvert, M., Wegener, G., & Hinrichs, K. U. (2020).~~ Substrate-dependent incorporation  
of carbon and hydrogen for lipid biosynthesis by *Methanosarcina barkeri*. *Environmental Microbiology Reports*, 12(5),  
555567. doi:10.1111/1758-2229.12876. ~~(2020).~~  
Zhang, X., Gillespie, A. L., & Sessions, A. L. ~~(2009).~~ Large D/H variations in bacterial lipids reflect central metabolic  
pathways. *Proceedings of the National Academy of Sciences*, 106(31), 12580-12586. doi:10.1073/pnas.0903030106. ~~(2009).~~