Point-by-point response to the issues raised by Referee 1 (Lukas Kohl)

We thank Lukas Kohl for the positive evaluation of our work and for the helpful comments to improve the manuscript. All comments and requested changes were taken into account. Please note that comments by the referee are in italics and that in the authors' answer the mentioned line numbers refer to the version of the revised manuscript including track changes.

#### Referee 1:

Schroll and co-authors studied the stable carbon isotope values of methane emitted during the aerobic decomposition of organic matter by two fungal species. Methane production by fungi during plant litter decomposition is a novel pathway of methane formation, that was recently documented by the authors and others. This manuscript, however, is the first study of the stable carbon isotope ( $d^{13}C$ ) values associated with this novel pathway and their relationship substrate  $d^{13}C$  values.

This study addresses/closes a knowledge gap in the isotope systematics of atmospheric methane that is relevant to the Biogeosciences readership. The authors used state of the art methods, and their conclusions are well supported by their results. The manuscript is well structured and easy to follow.

The study's strength is that this is the first study of its kind and provides unique stable isotope fractionation factors between biomass and methane produced by fungi. The study also used very robust measurement methods (GC/IRMS with preconcentration) that exceeds the precision, accuracy, and specificity of laser-based analysers. The main limitations of the study are that the authors did not test for contaminations by other microbial species during this study (this was, however, tested by the authors in similar incubations in a previous study). Another limitation is that the authors were not able to identify controls over relatively large variations in methane isotope values beside differences between C3 and C4 plants. This, however, is understandable given that the biochemistry of aerobic methane production in fungi remains poorly understood, and the authors contribution will surely help elucidate these pathways in the future.

<u>Authors</u>: We thank the referee for the positive evaluation of our manuscript. The reviewer's concerns are addressed below.

# Main comment:

The authors used two distinct fungal species, and state that these species include both white rot and brown rot fungi. However, I was unable to find where in the manuscript the authors identify which fungal species belongs to which group.

<u>Authors</u>: A description of which fungal species belongs to white and brown rot fungi was added to section '2.1 Selected fungi' (L77-78).

#### Minor comments:

1) L56. remove 'applications of' for easier sentence structure

Authors: Change applied.

2) L57-58. clarify what 'they' refers to in 'they might be used..', also, avoid 'fingerprints' ('characteristic d<sup>13</sup>C values?)

Authors: 'they' was clarified as ' $\delta^{13}$ C-CH<sub>4</sub> values' and 'fingerprints' was changed to 'characteristic  $\delta^{13}$ C values'.

- 3) L61. 'global isotopic patters': Do you mean the d<sup>13</sup>C values of atmospheric CH<sub>4</sub>?' Authors: Correct. For clarification purposes 'δ<sup>13</sup>C-CH<sub>4</sub>' was added.
- 4) L66. 'isotope patterns': stable isotope values?

Authors: Change applied.

5) L77-78. clarify which fungi is the white rot and which one is the brown rot one.

<u>Authors</u>: A specification of which fungal species belongs to white and brown rot fungi was added to section '2.1 Selected fungi'.

6) L148. is 0.06mg correct? This seems a very low sample inweight for EA/IRMS, although not impossible. Also, did you analyse analytical replicates? A single 0.06mg inweight is likely associated with a significant subsampling error.

<u>Authors</u>: Yes, the sample weight is correct. Around 0.06 mg of sample was used for the EA/IRMS measurements. Three replicates of each substrate were measured (n=3). Standard deviations for  $\delta^{13}$ C of the substrates were 0.5 ‰ for pine wood, 0.6 ‰ for grass and 0.1 ‰ for corn.

7) L170-173. You could add a note that the low R2 resulted from the lack of a change in d<sup>13</sup>C values (emission d<sup>13</sup>C was similar to background d<sup>13</sup>C). In this case, a low R2 does not indicate a poor relation between concentration and d<sup>13</sup>C value.

Authors: Thank you for the very helpful comment. We added a note according to the reviewer's suggestion.

8) L176-177. 'The SDs are given with a confidence interval of 1 σ': sentence not needed and meaningless.

Authors: Change applied.

9) L182-188. not needed, can be removed.

<u>Authors</u>: Please note, that for better readability we would like to keep this paragraph as it clearly explains the structure of section '3 Results and Discussion' and makes this section easy to follow for the reader.

10) L194. 'where': use 'in which' instead

Authors: Change applied.

11) L201-239 and Table 1. stating CH<sub>4</sub>:CO<sub>2</sub> ratios in µmol/mol instead of nmol/mmol would improve clarity.

<u>Authors</u>: Thank you for your suggestion. The units of the  $CH_4$ :  $CO_2$  ratios were changed accordingly throughout the whole manuscript.

12) L314-315. 'distinct differences in the patterns': redundant structure, could be simplified.

Authors: We reworded the sentence.

13) L318-324. This section could use some language editing for better flow. e.g. L306: 'the used growth substrates': The growth substrates used for this study... or similar.

Authors: This section was revised for a better flow.

14) L319-320. 'consist of various amounts': contain distinct amount of cellulose, [..], and other compounds.

Authors: Changes applied.

15) L320-321. structure in parenthesis: grammar

Authors: Change applied.

16) L321-322. ... source signatures might \_depend\_ on the metabolic pathways \_used by\_ the fungal species \_as well as\_ the chemical composition of the substrate (or similar)

Authors: Change applied.

17) L323. Therefore, we suggest: remove this phrase. "The selected ..."

Authors: Change applied.

18) L326. Figure 5  $\_compares\_\_the\_\_d^{13}C\text{-}CH_4$  values.

Authors: Change applied.

19) L334. 'depending on the photosynthetic pathway (C3, C4, or CAM)'

Authors: Change applied.

Point-by-point response to the issues raised by Referee 2

We thank Referee 2 for the positive evaluation of our work and for the helpful comments to improve the manuscript. All comments and requested changes were taken into account. Please note that comments by the referee are in italics and that in the authors' answer the mentioned line numbers refer to the version of the revised manuscript including track changes.

## Referee 2:

General comments: Methane is the second important anthropogenic greenhouse gas after carbon dioxide. Recent studies have shown that this gas can be produced under aerobic conditions by plants, algae, fungi and animals. In this manuscript, Schroll et al. cultivated two saprotrophic fungi on three different substrates and measured the stable carbon isotope values of methane. This study is the first to report the analysis of stable carbon isotope values of methane emitted from saprotrophic fungi. The authors found that the source values of  $\delta 13$ CH4, emitted by the fungi, were dependent on the fungal species and the metabolized substrate. Although this paper has some limitations in terms fungal species and substrates, it certainly opens the door for new and exciting work in the area of aerobic methane emissions. Overall, this is a well-written manuscript and deserves to be published in Biogeosciences after minor revisions.

<u>Authors</u>: We thank the referee for the positive evaluation of our manuscript. The reviewer's concerns are addressed below.

# Specific comments:

1) L16. eukaryotes,

Authors: Change applied.

2) L17-18. ecosystems via decomposition of plant litter

Authors: Change applied.

3) L18. Although the methane

Authors: Change applied.

4) L19. In this study,

Authors: Change applied.

5) L20-21. The common names of fungi must be mentioned here

Authors: The common names of the fungi have been added to the revised manuscript.

6) L21., cultivated... (pine...), reflecting

Authors: Change applied.

7) L21-22. Which grass? It is better to provide the Latin names of pine, grass (species name) and corn

Authors: The Latin names of the pine, grass and corn species have been added to the revised manuscript.

8) L23. Keeling; K must be uppercase here and in other places

Authors: Change applied.

9) L27. 'Whilst' should be replaced; it is mentioned in the previous sentence

Authors: Change applied.

10) L29. We found that the values of  $\delta$ 13CH4 emitted

Authors: Change applied.

11) L30. What is 'They' in 'They cover'?

Authors: Change applied.

12) L34. Fossil fuel burning indicates a process but not source; source is fossil fuel, biomass, and...

Authors: Change applied.

13) L37. microorganisms,

Authors: Change applied.

14) L40. discovered,

Authors: Change applied.

15) L45. It is better to delete 'therefore'

Authors: Change applied.

16) L46-47. White rot fungi (e.g., Latin name)... brown rot fungi (e.g., Latin name)

Authors: Examples for white rot fungi and brown rot fungi are now included in the manuscript.

17) L49. in the synthesis of CH4

Authors: Change applied.

18) L51. archaea with essential substrate... in fungus-infected wood stem

Authors: Change applied.

19) L55. might be an underestimated

Authors: Change applied.

20) L56. It is better to delete 'Applications of'; It is better to start the sentence with Stable isotope procedures <u>Authors</u>: 'Applications of' has been deleted. Please note, that we would like to write 'Stable carbon isotopes', as in this context it refers to stable isotopes in a general meaning.

21) L57-58. 'they' is referred to what?

Authors: Change applied.

22) L64. have been identified

Authors: Change applied.

23) L67-68. plant-derived CH4..., and UV-induced CH4...

Authors: Changes applied.

24) L69. In this study, we...

Authors: Change applied.

25) L76. Pleurotaceae and Polyporacaeae are the family names and should not be italicized.

Authors: Changes applied.

26) L81. Both common and Latin names should be provided for pine, grass (specific plant species) and corn Authors: Both names have been added to the revised manuscript.

27) L97-98. It is better to provide the temperature for autoclave

Authors: A more detailed description of the autoclave method was added to this section.

28) L114. What are those five different gases?

<u>Authors</u>: The five reference gases were certified gas mixtures of CH<sub>4</sub> and CO<sub>2</sub> with five different concentrations by Deuste Steininger GmbH. The name of the company was added to the manuscript to clarify the origin of the reference gases.

29) L141-143. Is 'the working reference gas' the standard reference gas?

Authors: We modified 'working reference gas' to read "working standard". We also corrected an error (L142) were the two reference standards are CH<sub>4</sub> and not CO<sub>2</sub>. Those two CH<sub>4</sub> reference standards are calibrated and certified and are used for the normalization of the samples. According to the 'Principle of identical treatment' the CH<sub>4</sub> reference gases were measured exactly in the same way as the samples.

30) L149. substrate was put... the resulting gases were separated...

Authors: Change applied.

31) L151. 27.5 m ... then reached

Authors: Change applied.

32) L153. Keeling

Authors: Change applied.

33) L159. Keeling

Authors: Change applied.

34) L161. Keeling...Keeling

Authors: Change applied.

35) L163. It is better to delete the first 'grown on pine'

Authors: Change applied.

36) L167. Keeling

Authors: Change applied.

37) L178. Was there a reason for using Fisher test instead of a robust test, such as Tukey's test?

Authors: The statistical evaluation with two way ANOVAs was chosen to conclude if there is a general effect of the fungi and substrates on  $CH_4$  and  $CO_2$  mixing-ratios,  $\delta^{13}CH_4$  and  $\delta^{13}CO_2$  values and the  $CH_4$ :  $CO_2$  emission ratios. The results of the post-hoc tests (Fisher least significance difference and Tukey) are attached in the supplement to this comment. Please note, that the post-hoc tests only have a limited value as there are only three repeated measurements for each parameter (n=3) and post-hoc tests are generally designed for a greater number of repeated measurements. Therefore, we prefer not to show the post-hoc tests in this manuscript and keep the general effects that are expressed by the two-way ANOVAs. Nevertheless, for the  $\delta^{13}C$ - $CH_4$  and  $\delta^{13}C$ - $CO_2$  isotope values p-values calculated with the Fisher LSD and Tukey test are similar and produce only minor differences. Please note that p-values (> 0.05) for  $CH_4$  and  $CO_2$  mixing-ratios might occur because either the quantity of emitted  $CH_4/CO_2$  by the fungi is similar and/or the biomass of the fungi within the flasks varies. The manuscript was changed accordingly (L177-178) to clarify that the statistical methods applied in the manuscript refer to the results of two-way ANOVAs.

38) L185. Keeling

Authors: Change applied.

39) L187. The second 'source' can be deleted.

Authors: Change applied.

40) L193. 'the' should be deleted.

Authors: Change applied.

41) L197. The second 'grown' should not be italicized.

Authors: Change applied.

42) L203-205. Most of the controls? It is better to be specific.

Authors: The sentence was modified to be more specific.

43) L205. ... . respectively were observed

Authors: This part of the sentence was replaced because of the changes made to the previous comment 42).

44) L215. was present

Authors: Change applied.

45) L229. thereby. both...; the 'both' after substrate should be deleted.

Authors: Change applied.

46) L230. Is it P < 0.001?; a comma should be added after sapidus

Authors: Yes. it is p<0.001! The comma was added after *P. sapidus*.

47) L237. 'in a good accordance' is not clear. it needs to be rewritten.

<u>Authors</u>: 'in a good accordance' was replaced by 'in the same order of magnitude' to make this sentence clearer.

48) L238. It should be noted that CH4

Authors: Change applied.

49) L272-274. It is better to rewrite this sentence. like: ...Keeling plot analysis that range ... are presented.

Authors: Change applied.

50) L276 and L280. P < 0.001 (number should not be italicized)

Authors: Change applied.

51) L289. Keeling; one of the 'values' should be deleted.

Authors: Change applied.

52) L295. 'as so far' is not clear

Authors: 'As so far' was replaced by 'as up to the present' to make the sentence clearer.

53) L297-298. The values of... that range from... are presented in Table 2

Authors: Change applied.

54) L300. 'more' should be deleted.

Authors: Change applied.

55) L304-305. 'Although... substrate' is not a sentence and should be rewritten.

Authors: The sentence was rewritten.

56) L309. 'usually' should be deleted from here and added after 'are'

Authors: Change applied.

57) L311. 'slightly more' should be reworded.

Authors: Change applied.

58) L318. CH4 and CO2 are derived

Authors: Change applied.

59) L331. a wide range

Authors: Change applied.

60) L351. sources. such as methanogenic archaea and eukaryotes.

<u>Authors</u>: Thanks for the note. We changed 'abiotic processes' to 'abiotic CH<sub>4</sub> sources' because the term 'abiotic processes' might be misleading. Nevertheless, we would like to keep the 'abiotic CH<sub>4</sub> sources' in this sentence.

61) L351. 'and from abiotic processes' should be deleted or modified in such a way to show sources

Authors: Please see response to previous comment 60).

62) L353. processes. resulting

Authors: Change applied.

63) L354-357. The sentence that starts with 'Thus. studying' is not clear and should be rewritten.

Authors: Change applied.

64) L358. research. stable

Authors: Change applied.

65) L376. Grant Numbers

Authors: Change applied.

66) L401. In CO2. 2 should be subscript.

Authors: Change applied.

67) L408. The title of this paper should be written in correct format.

Authors: Change applied.

68) L458. The Latin name should be italicized.

Authors: Change applied.

69) L464. In CH4. 4 should be subscript.

Authors: Change applied.

70) L465-L466. CH4 and 13C/12C should be written in correct format.

Authors: Changes applied.

71) L468-470. In CH4. 4 should be subscript; In 13C/12C. 13 and 12 should be superscript.

Authors: Changes applied.

72) L529. The Latin name should be italicized.

Authors: Change applied.

73) L550. Plant Cell Environ.

Authors: Change applied.

# The stable carbon isotope signature of methane produced by saprotrophic fungi

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**Abstract.** Methane (CH<sub>4</sub>) is the most abundant organic compound in the atmosphere with emissions from many biotic and abiotic sources. Recent studies have shown that CH<sub>4</sub> production occurs under aerobic conditions in eukaryotes, such as plants, animals, algae and saprotrophic fungi. Saprotrophic fungi play an important role in nutrient recycling in terrestrial ecosystems by their ability tovia decompositione of plant litter. Even Although the CH<sub>4</sub> production by saprotrophic fungi has been reported, so far, no data for stable carbon isotope values of the emitted  $CH_4(\delta^{13}C-CH_4 \text{ values})$  is available. In this study, we measured the  $\delta^{13}$ C values of CH<sub>4</sub> and carbon dioxide ( $\delta^{13}$ C-CO<sub>2</sub> values) emitted by the two saprotrophic fungi *Pleurotus sapidus* (oyster mushroom) and Laetiporus sulphureus (sulphur shelf) cultivated, on three different substrates pine wood (Pinus sylvestris), grass (mixture of Lolium perenne, Poa pratensis, Festuca rubra) and corn (Zea mays), reflecting both C3 and C4 plants with distinguished bulk  $\delta^{13}$ C values. Applying keeling Keeling plots, we found that the  $\delta^{13}$ C source values of CH<sub>4</sub> emitted from fungi cover a wide range from -40 mUr to -69 mUr depending on the growth substrate and fungal species. Whilst little apparent carbon isotopic fractionation (in the range of -0.3 mUr to 4.6 mUr) was calculated for  $\delta^{13}$ C values of CO<sub>2</sub> released from P. sapidus and L. sulphureus relative to the bulk  $\delta^{13}$ C values of the growth substrates, much larger carbon isotopic fractionations (ranging from -22 mUr to -42 mUr) were observed for the formation of CH<sub>4</sub>. WhilstAlthough the two fungal species showed similar δ<sup>13</sup>CH<sub>4</sub> source values when grown on pine wood, δ<sup>13</sup>CH<sub>4</sub> source values differed substantially between the two fungal species when grown on grass or corn. We found that the source values of  $\delta^{13}$ CH<sub>4</sub>-source values emitted by saprotrophic fungi are highly dependent on the fungal species and the metabolized substrate. The source values of  $\delta^{13}$ CH<sub>4</sub>They cover a broad range of 8<sup>13</sup>CH<sub>4</sub> values and overlap with values reported for methanogenic archaea, thermogenic degradation of organic matter and other eukaryotes.

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#### 1 Introduction

Methane (CH<sub>4</sub>) is an important greenhouse gas that is emitted by several abiotic sources (e.g. fossil fuel burning and use, biomass burning, geological processes) and biotic sources (e.g. wetlands, agriculture and waste, fresh waters) to the atmosphere (Kirschke et al., 2013; Saunois et al., 2016, 2019). In the past, biotic CH<sub>4</sub> production has been attributed exclusively to strictly anaerobic microorganisms, such as methanogens that are ubiquitous in wetlands, rice paddies, landfills and the intestines of termites and ruminants (Kirschke et al., 2013). The discovery of CH<sub>4</sub> emissions from dead and living plants under oxic conditions (Keppler et al., 2006, 2009) paved the way for the search of new biogenic CH4 sources. Since then, several previously unknown CH<sub>4</sub> sources were discovered, including endothelial cells of rat liver (Boros and Keppler, 2019; Ghyczy et al., 2008), plant cell cultures (Wishkerman et al., 2011), marine algae (Klintzsch et al., 2019; Lenhart et al., 2016), marine and terrestrial cyanobacteria (Bižić et al., 2020), humans (Keppler et al., 2016) and saprotrophic fungi (Lenhart et al., 2012). Fungi play a central role in ecosystems by decomposing organic matter and thereby recycling formerly bound carbon and nutrients (Grinhut et al., 2007). This process is especially important in forests where fungi are essential for wood decay and therefore have a great impact on the carbon and nitrogen cycles in these environments (Ralph and Catcheside, 2002). White rot fungi (e.g. Trametes versicolor or Pleurotus ostreatus) are able to decompose the chemically complex structural component lignin, whereas brown rot fungi (e.g. Serpula lacrymans or Gloeophyllum trabeum) mainly metabolize cellulose and hemicellulose (Ten Have and Teunissen, 2001; Leonowicz et al., 1999; Valášková and Baldrian, 2006). Fungi have already been determined to be involved in the synthesis of CH<sub>4</sub> synthesis during wood decay (Beckmann et al., 2011; Mukhin and Voronin, 2007, 2008) by breakdown of large macromolecules to smaller molecules, thereby providing bacteria and methanogenic archaea with their essential substrate. Elevated levels of CH<sub>4</sub> were found in fungus-ally infected wood stems with oxygen concentrations ranging from 1 to 14 % (Hietala et al., 2015). Here, CH<sub>4</sub> production was associated with anoxic microsites in the xylem, indicating that at least part of the CH<sub>4</sub> was produced by methanogenic archaea. Nevertheless, Lenhart et al., 2012 demonstrated that basidiomycetes are able to produce CH<sub>4</sub> under aerobic conditions without the presence of methanogenic archaea. Therefore, fungi might be an so far underestimated source of  $CH_4$  in the global  $CH_4$  cycle. Applications of Stable carbon isotopes (expressed as  $\delta^{13}$ C values) have often been used to investigate sources and sinks of CH<sub>4</sub> on the global scale (Whiticar, 1993). As different CH<sub>4</sub> sources have distinct characteristic  $\delta^{13}$ C values fingerprints, they δ<sup>13</sup>C-CH<sub>4</sub> values might be used to quantify the individual contributions of various sources regionally and/or globally (Dlugokencky et al., 2011; Hein et al., 1997; Nisbet et al., 2016; Quay et al., 1999; Tyler, 1986; Whiticar, 1999). The short lifetime of CH<sub>4</sub> in the atmosphere (range from 9.7 ± 1.5 to 11.2 ± 1.3 years) (Naik et al., 2013; Prather et al., 2012; Voulgarakis et al., 2013) assures that global isotopic 8<sup>13</sup>C-CH<sub>4</sub>-patterns represent the average of recent inputs by various sources and allows the quantification of respective source strengths (Mikaloff Fletcher et al., 2004b, 2004a). Additionally, stable isotopes provide information about the formation processes of CH<sub>4</sub>. Traditionally, three formation categories of  $\delta^{13}$ C-CH<sub>4</sub> values have been identifiedelassified: biogenic, with typical  $\delta^{13}$ C-CH<sub>4</sub> values ranging from ~-55 mUr to -70 mUr, thermogenic (ranging from ~-25 mUr to -55 mUr) and pyrogenic (ranging from ~-13 mUr to -25 mUr) (Kirschke

et al., 2013). However, isotopic patternsstable isotope values of recently identified CH<sub>4</sub> sources, i.e. human CH<sub>4</sub> emissions (-56 mUr to -95 mUr) (Keppler et al., 2016), plant\_derived CH<sub>4</sub> (-52 mUr to -69 mUr) (Keppler et al., 2006), and abiotic UV<sub>2</sub> induced CH<sub>4</sub> formation by plants (-52 mUr to -67 mUr) (Vigano et al., 2009) also need to be considered.

In this study, we investigated the stable carbon isotope source signatures of CH<sub>4</sub> and CO<sub>2</sub> released by the two basidiomycetes Pleurotus sapidus (white rot fungus) and Laetiporus sulphureus (brown rot fungus). Both fungi were cultivated under sterile conditions on three different substrates (pine wood, grass, and corn) with varying bulk  $\delta^{13}$ C values. We examined the influence of fungal species and growth substrate on  $\delta^{13}$ C-CH<sub>4</sub> and  $\delta^{13}$ C-CO<sub>2</sub> values and compared the  $\delta^{13}$ C-CH<sub>4</sub> values from the two fungal species with those of other known sources reported from the literature.

#### 2 Material and Methods

# 75 2.1 Selected fungi

P. sapidus (Pleurotaceae, DSMZ 8266) and L. sulphureus (Polyporacaeae, DSMZ 1014) were chosen for this experiment because of their capability to emit CH<sub>4</sub> (Lenhart et al., 2012), their ecological and physiological characteristics (P. sapidus is a white rot fungus and L. sulphureus is a brown rot fungus;) and well-established practical handling under laboratory conditions.

# 80 2.2 Cultivation of fungi and incubation experiments

Pine wood (*Pinus sylvestris*), grass (mixture of *Lolium perenne*, *Poa pratensis*, *Festuca rubra*) and corn (*Zea mays*) were selected as growth substrates. Pine wood was chosen to investigate if white rotbrown and brownwhite rot fungi differ in  $\delta^{13}$ C-CH<sub>4</sub> and  $\delta^{13}$ C-CO<sub>2</sub> values released during wood decay. Therefore, dead pine wood branches were collected from the forest floor and shredded to small wood chips with a length of about 5 cm (Natura 1800L; Glora, Witten, Germany). The wood chips were dried at 60°C for 48h and stored in a flask (Weck, Hanau, Germany). Grass (C<sub>3</sub> plant) and corn (C<sub>4</sub> plant) were selected because of their different stable isotope values. As the metabolic pathway for carbon fixation is biochemically different in C<sub>3</sub> and C<sub>4</sub> plants, plant biomass differs in  $\delta^{13}$ C values, which in turn might lead to different  $\delta^{13}$ C values of CH<sub>4</sub> and CO<sub>2</sub> released by fungi. Therefore, typical garden lawn was manually cut, dried at 70 °C, and stored in a flask. The corn substrate consisted of conventional corn flour.

The substrates were autoclaved and filled into 2.7 l flasks (Weck, Hanau, Germany) and inoculated with pure fungal submerged cultures under sterile conditions according to Lenhart et al., 2012. After addition of the fungi, the flasks were closed with lids and a rubber band sealing. To allow gas exchange during the growth time of the fungi (about two weeks), a hole in the centre of every lid was fitted with a cotton stopper. Before the start of the incubation experiments, the flasks were aerated under sterile conditions in order to start the incubation at atmospheric CH<sub>4</sub> mixing ratios. Additionally, to seal the flasks airtight the cotton stoppers were replaced by sterile silicone stoppers (Saint-Gobain Performance Plastics, Charny, France).

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For the incubation experiments, P. sapidus und L. sulphureus were incubated on the three substrates, while substrates were incubated as control treatments. Before the incubation experiments, the substrates were sterilized by autoclaving at 121 °C and 2 bar pressure for 20 minutes. The incubation experiments were conducted as three replicates per treatment. The duration of the incubation accounted for up to 40 h. All incubations were conducted at room temperature (23  $\pm$  1.5 °C). At every sampling point, 40 ml air was taken from the flasks for gas concentration measurements and an additional 40 ml were taken for  $\delta^{13}$ C-CH<sub>4</sub> stable isotope ratio mass spectrometry (IRMS) analysis. The gas samples were taken with airtight 60 ml PE syringes (Plastipak, BD, Franklin Lakes, USA) and transferred into 12 ml evacuated Exetainers (Labco, High Wycombe, UK). Subsequently a volume of atmospheric air equivalent to the volume of the removed sample was added into each flask directly after sampling. Mixing ratios and stable isotope values of CH<sub>4</sub> were corrected according to the dilution.

When calculating the fungal CH<sub>4</sub> and CO<sub>2</sub> production rates, we subtracted substrate derived CH<sub>4</sub> and CO<sub>2</sub> production rates (determined in the control treatments) from the respective fungi containing samples. Additionally, for the calculation of the fungal production rates only sample points showing a linear increase in CH<sub>4</sub> and CO<sub>2</sub> were taken into account.

To account for differences in the metabolic activity of the fungi, we additionally measured respiration rates, assuming that metabolic activity correlates with respiration and therefore CO<sub>2</sub> emissions of the fungi. Hence, we related fungal derived CH<sub>4</sub> emissions to respiration by calculating the CH<sub>4</sub>: CO<sub>2</sub> emission ratio.

# 2.3 Analysis of CH<sub>4</sub> and CO<sub>2</sub> via gas-chromatography

Samples were analysed using a gas chromatograph (GC, Bruker Greenhouse Gas Analyser 450-GC) equipped with a flame ionization detector (FID) and an electron capture detector (ECD) for the detection of CH<sub>4</sub> and CO<sub>2</sub>, respectively. The detector temperatures were set at 300 °C (FID) and 350 °C (ECD). Five reference gases (Deuste Steininger GmbH) were used for calibrating the GC-system. The reference gases were in the range of 1 parts per million by volume (ppmv) to 21 ppmv and 304 ppmv to 40,000 ppmv for CH<sub>4</sub> and CO<sub>2</sub>, respectively. Gas peaks were integrated using Galaxie software (Varian Inc., Palo Alto, CA, USA).

#### 2.4 Definition of $\delta$ values and isotope apparent fractionation

In this paper, all stable carbon isotope ratios are expressed in the conventional 'delta' δ notation, meaning the relative difference
of the isotope ratio of a substance compared to the standard substance Vienna Peedee Belemnite (V-PDB) (Eq. (1)).

$$\delta^{13}C = \frac{\frac{^{13}C}{^{12}C}}{\frac{^{13}C}{^{12}C}} - 1 \tag{1}$$

The apparent fractionation ( $\epsilon_{app}$ ) between fungal  $\delta^{13}C\text{-CH}_4$  or  $\delta^{13}C\text{-CO}_2$  values and the  $\delta^{13}C$  values of the substrates was calculated according to Eq. (2).

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$$\epsilon_{\text{app CH4 or CO2}} = \frac{\left(\delta^{13}C + 1\right)_{\text{fungal CH}_4 \text{ or CO}_2}}{\left(\delta^{13}C + 1\right)_{\text{substrate}}} - 1$$
 (2)

We follow the proposal of Brand and Coplen, 2012 and use the term 'urey' (Ur) as the isotope delta unit, in order to conform with the guidelines for the International System of Units (SI). Hence, isotope delta values that were formerly given as -70 ‰, are expressed as -70 mUr.

#### 2.5 Measurements of δ13CH4 and δ13CO2 values

Stable carbon isotope values of CH<sub>4</sub> and CO<sub>2</sub> were measured using a continuous flow isotope mass spectrometry system (CF-IRMS). A HP 6890N GC (Agilent, Santa Clara, USA) was linked to a preconcentration unit for CH<sub>4</sub> measurements and an autosampler A200S (CTC Analytics, Zwingen, Switzerland) for CO<sub>2</sub> analysis. The GC was equipped with a CP-PoraPLOT Q capillary column (Varian, Palo Alto, USA) (27,5 m x 0.25 mm i.d., film thickness 8 μm). The GC was operated with an injector temperature of 200°C, isothermal oven temperature of 30°C, split injection (10:1) and a constant carrier gas flow of 1.8 ml
 min<sup>-1</sup> (methane-free helium). The GC was coupled to a Delta<sup>PLUS</sup>XL isotope ratio mass spectrometer (ThermoQuest Finnigan, Bremen, Germany) via an oxidation reactor and a GC Combustion III Interface (ThermoQuest Finnigan, Bremen, Germany). The oxidation reactor was employed with the following properties: ceramic tube (Al<sub>2</sub>O<sub>3</sub>), length 320 mm, 1.0 mm i.d., with Ni/Pt wires inside activated by oxygen, reactor temperature 960 °C.

For CH<sub>4</sub> measurements with the preconcentration unit, headspace gas samples were transferred to an evacuated 40 ml sample loop. Methane was trapped on Hayesep D, separated from other compounds by the GC and then introduced into the IRMS system via an open split. The working referencemonitor gas was carbon dioxide of high purity (carbon dioxide 4.5, Messer Griesheim, Frankfurt, Germany) with a known  $\delta^{13}$ C value of -23.6 mUr (calibrated at MPI for Biogeochemistry in Jena, Germany). All  $\delta^{13}$ C values were corrected using two working-CH<sub>4</sub> reference gases of high purity carbon dioxidestandards (Isometric instruments, Victoria, Canada) with  $\delta^{13}$ C values of -23.9 ± 0.2 mUr and -54.5 ± 0.2 mUr that were calibrated against IAEA and NIST reference substances. The normalization of the sample values was done according to Paul et al., 2007.

# ${\bf 2.6~Bulk~isotope~analysis~of~fungal~substrates}$

Stable carbon isotope values of the bulk substrate were measured using an Elemental Analyzer Flash EA 11112 (Thermo Fischer Scientific, Germany) coupled to a Delta V IRMS (Thermo Fischer Scientific, Germany). Therefore, 0.06 mg of the substrate were was put into a tin cup and combusted in the Elemental Analyzer. The resulting gases were separated in a GC by a CP-PoraPLOT Q capillary column (Varian, Palo Alto, USA) (27,5 m x 0.25 mm i.d., film thickness 8 µm) and then reached the Delta V IRMS via a Conflo IV Universal Continuous Flow Interface (Thermo Fischer Scientific, Germany). Isotope values were corrected using USGS 40 and USGS 41 standards.

## 2.7 Determination of isotopic source signature of CH<sub>4</sub> and CO<sub>2</sub> applying keeling Keeling plots

For the determination of  $\delta^{13}$ C source values of CH<sub>4</sub> and CO<sub>2</sub> the keeling Neeling plot method was used (Keeling, 1958; Pataki et al., 2003) (Eq. (3)):

$$\delta^{13}C_{a} = c_{b}(\delta^{13}C_{b} - \delta^{13}C_{s}) \left(\frac{1}{c_{s}}\right) + \delta^{13}C_{s}$$
(3)

where  $c_a$  is the mixing ratio of CH<sub>4</sub>/CO<sub>2</sub> in the headspace,  $\delta^{13}C_a$  is the  $\delta^{13}C$  value of CH<sub>4</sub>/CO<sub>2</sub> in the headspace,  $c_b$  is the mixing ratio of background CH<sub>4</sub>/CO<sub>2</sub>,  $\delta^{13}C_b$  is the  $\delta^{13}C$  value of background CH<sub>4</sub>/CO<sub>2</sub> and  $\delta^{13}C_s$  the  $\delta^{13}C$  source value of the CH<sub>4</sub>/CO<sub>2</sub>. For a more detailed description of the application of keeling Feeling plots for determination of CH<sub>4</sub> source signature we refer to the study by Keppler et al., 2016.

 $\delta^{13}$ C-CH<sub>4</sub> source signatures were calculated after the <u>keeling Keeling</u> plot method for each flask. Results of the <u>keeling Keeling</u> plots are then given as the arithmetic mean of the three individual flasks per treatment with standard deviations (n=3).

 $\delta^{13}$ C-CH<sub>4</sub> source signatures of each flask of *P. sapidus*-grown on pine and *L. sulphureus* grown on pine were corrected for CH<sub>4</sub> emissions and  $\delta^{13}$ C-CH<sub>4</sub> values of the "pine" control samples using the following mass balance approach (Eq. (4)).

$$\delta^{13}C_{fungi\ corrected} = \frac{\left(P(CH_4)_{fungi} * \delta^{13}C_{fungi}\right) \cdot \left(P(CH_4)_{pine} * \delta^{13}C_{pine}\right)}{\left(P(CH_4)_{fungi} \cdot P(CH_4)_{pine}\right)} \tag{4}$$

, where P(CH<sub>4</sub>)  $_{\text{fungi/pine wood}}$  is the CH<sub>4</sub> emitted by the fungi or pine wood and  $\delta^{13}$ C  $_{\text{fungi/pinewood}}$  is the  $\delta^{13}$ C-CH<sub>4</sub> source signature of the fungi or pine wood derived from  $_{\text{keeling-Keeling}}$  plots. Corrected  $\delta^{13}$ C-CH<sub>4</sub> source values for *P. sapidus* and *L. sulphureus* are given as the arithmetic mean of the three individual flasks per treatment with standard deviations (n=3).

The determination coefficient ( $R^2$ ) of the keeling Keeling plots showed values higher than 0.93, except for *P. sapidus* grown on grass ( $R^2$ =0.51). The lPlease note that the, in comparison to the other incubation experiments, lower  $R^2$  value for *P. sapidus* grown on grass is probably a result of the marginal results from a lack of changes of  $\delta^{13}$ C-CH<sub>4</sub> values due to the only a small increase emission of the CH<sub>4</sub> mixing ratio compared to the background CH<sub>4</sub> mixing ratio. Therefore, the low  $R^2$  does not necessarily indicate a weaker poorrelationship relation between CH<sub>4</sub> mixing ratio and  $\delta^{13}$ C-CH<sub>4</sub>.

## 2.8 Statistics

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Mixing ratios and production rates of CH<sub>4</sub>, CO<sub>2</sub>, δ<sup>13</sup>C-CH<sub>4</sub> and δ<sup>13</sup>C-CO<sub>2</sub> values and δ<sup>13</sup>C source values are presented as arithmetic mean of three independent replicates with standard deviations (SD; n = 3). The SDs are given with a confidence interval of 1-σ. Linear regression analysis, arithmetic means and SDs were calculated using Excel (Microsoft Excel for Office 365 MSO). Two-way analysis of variance (ANOVA) and a post hoc test (Fisher least significant difference) (SigmaPlot 12.2.0.45, USA) were carried out to test for "species" and "substrate" related effects on δ<sup>13</sup>C-CH<sub>4</sub> and δ<sup>13</sup>C-CO<sub>2</sub> source values for each treatment. Differences at the p < 0.05 level were referred to as significant.</li>

#### 3 Results and Discussion

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In this section, we firstly present the results of  $CH_4$  and  $CO_2$  production from the two fungal species grown on the three different substrates. This includes emission rates of  $CH_4$  and  $CO_2$  from the control treatments of pine wood, grass and corn as well as the molar ratio of  $CH_4$  and  $CO_2$ . Secondly, we then present the respective stable isotope values measured for  $CH_4$  and  $CO_2$  during the incubation experiments and calculate the stable isotope source values of  $CH_4$  and  $CO_2$  released by the fungi applying keeling. Feeling plots. We then compare these values with stable carbon isotope values of the bulk organic matter by calculating the apparent fractionation. Finally, we compare  $\delta^{13}C$  source values of fungal derived  $CH_4$  with sources values known for other  $CH_4$  sources from the literature.

# 3.1 Release of CH<sub>4</sub> and CO<sub>2</sub> from P. sapidus and L. sulphureus

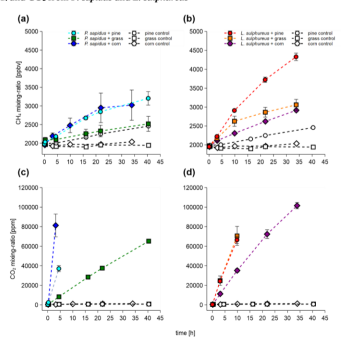


Figure 1: Mixing ratios of CH<sub>4</sub> and CO<sub>2</sub> of *P. sapidus* (a, c) and *L. sulphureus* (b, d) grown on pine wood, grass, and com. Mixing ratios are presented as mean values with standard deviation SD (n=3).

All incubation experiments where in which fungi were grown on the different substrates showed a significant increase in CH<sub>4</sub> compared to the respective substrate control (Fig. 1 a, c). Calculated emission rates for CH<sub>4</sub> and CO<sub>2</sub> are presented in Table 1. L. sulphureus grown on grass  $(7.5 \pm 1.3 \text{ nmol h}^{-1})$  showed the highest emission rate of CH<sub>4</sub>, followed by L. sulphureus grown on pine  $(6.2 \pm 0.3 \text{ nmol h}^{-1})$ , P. sapidus grown on corn  $(4.4 \pm 1.9 \text{ nmol h}^{-1})$ , L. sulphureus grown on corn  $(2.6 \pm 0.1 \text{ nmol h}^{-1})$ , P. sapidus grown on pine  $(2.5 \pm 0.2 \text{ nmol h}^{-1})$  and P. sapidus grown on grass  $(1.4 \pm 0.5 \text{ nmol h}^{-1})$ . Please note that CH<sub>4</sub> and CO<sub>2</sub> emission rates are not related to fungal biomass. Therefore, differences in the emission rates might be due to varying fungal biomass of the subsamples. Instead, CH<sub>4</sub> production was related to CO<sub>2</sub> production by determining the molar emission ratio between CH<sub>4</sub> and CO<sub>2</sub> (nµmol CH<sub>4</sub>: mmol CO<sub>2</sub>). CO<sub>2</sub> production thereby reflects the amount of fungal biomass and is also an indicator for the metabolic activity of the fungi.

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Most of the The control flasks did not show significant changes in their CH<sub>4</sub> and CO<sub>2</sub> mixing ratios over time, except for CH<sub>4</sub> in pine wood controls (1.3 ± 0.1 nmol h<sup>-1</sup>). However, in the control flasks of pine wood and corn small CH<sub>4</sub> emission rates of 1.3 ± 0.1 nmol h<sup>-1</sup> and 0.25 ± 0.01 nmol h<sup>-1</sup>, respectively were observed, and in the control 'grass' the CH<sub>4</sub> mixing ratio slightly decreased over time (-0.05 ± 0.04 nmol h<sup>-1</sup>). Whilst the pine wood and corn control flasks showed a small increase in the CH<sub>4</sub> mixing ratio, they did not show an increase in CO<sub>2</sub> mixing ratios. These data rule out a contamination by microbial heterotrophs, as this would cause a measurable CO<sub>2</sub> increase within the flasks. The CH<sub>4</sub> increase in the substrate controls might be attributed to CH<sub>4</sub> release by dead plant material as it was already shown by Keppler et al., 2006 and Vigano et al., 2009. Within the scope of these experiments, no analytic test for microbial contamination was conducted. Nevertheless, Lenhart et al., 2012 clearly showed that with the performed method of cultivation of fungi and incubation experiments no methanogenic archaea were present, using three different methods (Fluorescence in situ hybridization (FISH), confocal laser scanning microscopy (CLSM) and quantitative real time PCR). Furthermore, CH<sub>4</sub> and CO<sub>2</sub> release and the CH<sub>4</sub>: CO<sub>2</sub> emission ratios in our incubations are similar to the experiments of Lenhart et al., 2012 and do not indicate microbial contamination. Therefore, we assume that in our investigations no contamination with bacteria or methanogenic archaea wasere present.

For *P. sapidus* grown on corn and *L. sulphureus* grown on grass, no further linear increase in CH<sub>4</sub> was observed after 22 h and 10 h, respectively. This might be due to a reduced decay of organic matter and slower fungal metabolism because of higher CO<sub>2</sub> and lower O<sub>2</sub> mixing ratios.

A drastic increase in CO<sub>2</sub> mixing ratios relative to the controls was observed in all flasks containing fungi (Fig. 1 b, d). The CO<sub>2</sub> emission rates are shown in Table 1. CO<sub>2</sub> production rates ranged from 176 ± 4 μmol h<sup>-1</sup> to 2910 ± 410 μmol h<sup>-1</sup> for *P. sapidus* grown on grass and *P. sapidus* grown on corn, respectively. These highly variable CO<sub>2</sub> production rates might reflect different fungal biomass and metabolic activity (mineralisation of organic matter). In the control treatments, tiny increases in the CO<sub>2</sub> mixing ratio were detected ranging from 0.64 ± 0.12 μmol h<sup>-1</sup> to 0.91 ± 0.14 μmol h<sup>-1</sup>. Only one flask (corn control) showed a somewhat higher increase in CO<sub>2</sub> (7.76 μmol h<sup>-1</sup>), which is most likely caused by microbial contamination of the flask. However, no increase in the CH<sub>4</sub> mixing ratio was detected (see supplementary material). Therefore, this control flask was excluded from further calculations.

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Mean CH<sub>4</sub> and CO<sub>2</sub> emission rates and CH<sub>4</sub>: CO<sub>2</sub> emission ratios of all treatments are presented in Table 1. Higher ratios indicate a higher CH<sub>4</sub> production during decay of the substrates. Thereby, both fungal species and substrate both affect the CH<sub>4</sub>: CO<sub>2</sub> emission ratio (p><0.001). For P. sapidus, CH<sub>4</sub>: CO<sub>2</sub> emission ratios are more variable (1.4 to 8.0 nmol µmol CH<sub>4</sub>/mmol CO<sub>2</sub>) compared to L. sulphureus (6.7 – 9.6 nµmol CH<sub>4</sub>/mmol CO<sub>2</sub>). This variation might be due to differences in the fungi's enzyme sets required for organic matter decay, as P. sapidus is a white rot fungus and L. sulphureus is a brown rot fungus. At present the biochemical pathways that lead to CH<sub>4</sub> are still unknown, although compounds such as the sulphurbound methyl-group of methionine and glucose have been identified to act as carbon precursors of fungal-derived CH<sub>4</sub> (Lenhart et al., 2012).

Lenhart et al., 2012 found CH<sub>4</sub>: CO<sub>2</sub> ratios of fungi that ranged between 8 µmmol CH<sub>4</sub>/mmol CO<sub>2</sub> and 17 nµmol CH<sub>4</sub>/mmol CO<sub>2</sub>, which is in a good accordance within the same order of magnitude as the CH<sub>4</sub>: CO<sub>2</sub> ratios determined in this study. Please It should be noted; that CH<sub>4</sub>: CO<sub>2</sub> ratios of Lenhart et al., 2012 were given in ppbv CH<sub>4</sub>: % CO<sub>2</sub> and for better comparability CH<sub>4</sub>: CO<sub>2</sub> ratios were converted to fit the units used in this study (µmmol CH<sub>4</sub>: mmol CO<sub>2</sub>).

**Table 1:**  $CH_4$  and  $CO_2$  production rates and molar  $CH_4$ :  $CO_2$  emission ratios of the fungi incubated on different substrates. Values are presented as mean values of three independent replicates with SD (n = 3), except for the control "corn" (n=2).

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Fungi	Substrate	CH <sub>4</sub> production rate	CO <sub>2</sub> production rate	CH4: CO2 ratio	
		[nmol h <sup>·1</sup> ]	[µmol h <sup>-1</sup> ]	[ <u>µ</u> mol/mmol]	
P. sapidus	pine	$2.5 \pm 0.2$	901 ± 79	$2.8 \pm 0.4$	
	grass	$1.4\pm0.5$	$176 \pm 4$	$8.0\pm2.8$	
	corn	$4.4 \pm 1.9$	$2910 \pm 419$	$1.4\pm0.5$	
L. sulphureus	pine	$6.2 \pm 0.3$	$724 \pm 42$	$8.6 \pm 1.0$	
	grass	$7.5 \pm 1.3$	$771 \pm 103$	$9.6\pm0.5$	
	corn	$2.6 \pm 0.1$	$385 \pm 20$	$6.7 \pm 0.4$	
control	pine	$1.3 \pm 0.1$	$0.64 \pm 0.12$	-	
	grass	$-0.05 \pm 0.04$	$0.91 \pm 0.14$	-	
	corn	0.25	0.66	-	

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# 3.2 Stable carbon isotope values of CH<sub>4</sub> and CO<sub>2</sub>

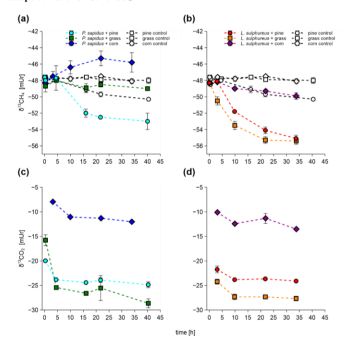


Figure 2: Stable carbon isotope values of CH<sub>4</sub> and CO<sub>2</sub> of *P. sapidus* (a, c) and *L. sulphureus* (b, d) grown on pine, grass, and corn. Values are presented as mean values with SD (n=3), except for  $\delta^{13}$ CO<sub>2</sub> values of *L. sulphureus* grown on corn (n=2).

Stable carbon isotope values of CH<sub>4</sub> and CO<sub>2</sub> measured from the incubation experiments are presented in Fig. 2. All incubations show a trend towards more negative δ<sup>13</sup>C-CH<sub>4</sub> values (less <sup>13</sup>C) with time except for *P. sapidus* grown on corn, where a tendency towards more positive δ<sup>13</sup>C-CH<sub>4</sub> values was observed (Fig. 2 a, b). During the incubation, δ<sup>13</sup>C-CH<sub>4</sub> values changed from -47.7 ± 0.1 mUr (for incubation of *P. sapidus* grown on pine/grass) and -48.2 ± 0.1 mUr (for incubation of *P. sapidus* grown on corn and *L. sulphureus* grown on pine/grass/corn) to -53.0 ± 0.7 mUr (*P. sapidus* grown on pine), -48.7 ± 0.3 mUr (*P. sapidus* grown on grass), -45.8 ± 1.2 mUr (*P. sapidus* grown on corn), -55.1 ± 0.4 mUr (*L. sulphureus* grown on pine), -55.4 ± 0.4 mUr (*L. sulphureus* grown on grass) and -49.9 ± 0.4 mUr (*L. sulphureus* grown on corn). The controls showed no significant shift in δ<sup>13</sup>C-CH<sub>4</sub> values except for the control "pine", where an increase in the CH<sub>4</sub> mixing ratio along with more

negative values of  $\delta^{13}$ C-CH<sub>4</sub> values occurred over time. This was accounted for when calculating the  $\delta^{13}$ C-CH<sub>4</sub> source signatures for *P. sapidus* grown on pine and *L. sulphureus* grown on pine (see materials and methods 2.7).

The δ<sup>13</sup>C-CO<sub>2</sub> values showed a trend towards more negative values within the first three to four hours of incubation (Fig. 2 c, d). After this time only minor changes of the δ<sup>13</sup>C-CO<sub>2</sub> values occurred. Final δ<sup>13</sup>C-CO<sub>2</sub> values of the incubation were -24.9 ±
260 0.6 mUr (*P. sapidus* grown on pine), -28.6 ± 0.9 mUr (*P. sapidus* grown on grass), -12.0 ± 0.3 mUr (*P. sapidus* grown on corn), -24.1 ± 0.1 mUr (*L. sulphureus* grown on pine), -27.7 ± 0.5 mUr (*L. sulphureus* grown on grass) and -13.0 ± 0.5 mUr (*L. sulphureus* grown on corn).

Table 2: Calculated  $\delta^{13}$ C-CH<sub>4</sub> and  $\delta^{13}$ C-CO<sub>2</sub> source signatures,  $\delta^{13}$ C values of the substrates, and  $\epsilon_{app CH4}$  and  $\epsilon_{app CO2}$ . Values 265 are presented as mean values with the SD (n=3).

Fungi	Substrate	δ <sup>13</sup> C-CH <sub>4</sub> source [mUr]	δ <sup>13</sup> C-CO <sub>2</sub> source [mUr]	δ <sup>13</sup> C substrate [mUr]	Eapp CH4	Eapp CO2
		[IIIO1]	source [mor]	[IIIO1]	[mUr]	[IIIOF]
P. sapidus	pine	-65.3 ± 1.1	$-24.1 \pm 0.1$		-38.4 ± 1.2	$4.0 \pm 0.1$
	grass	$-52.9 \pm 1.6$	$-27.4 \pm 1.3$		$-21.8 \pm 1.7$	$4.6\pm1.3$
	corn	$-39.8 \pm 2.0$	$\text{-}12.0 \pm 0.3$		$\text{-}28.5 \pm 2.0$	$-0.3 \pm 0.3$
L. sulphureus	pine	$-61.4 \pm 0.5$	$-25.0 \pm 0.5$		$\text{-}34.4 \pm 0.6$	$3.0 \pm 0.4$
	grass	$-69.2 \pm 1.9$	$\text{-}29.0 \pm 0.5$		$\text{-}38.6 \pm 2.0$	$2.9 \pm 0.5$
	corn	$-53.4 \pm 1.1$	$\text{-}12.8 \pm 0.3$		$-42.2 \pm 1.1$	$\text{-}1.1 \pm 0.3$
control	pine			$-28.0 \pm 0.5$		
	grass			$-31.5 \pm 0.6$		
	corn			$-11.7 \pm 0.1$		

#### 3.3 \delta^{13}C-CH4 and \delta^{13}C-CO2 source signatures of fungi

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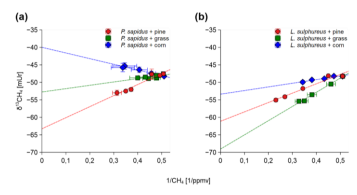


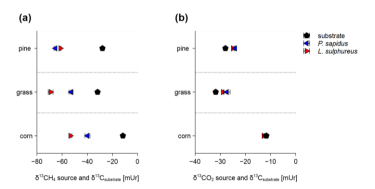
Figure 3: Keeling plots shown for *P. sapidus* (a) and *L. sulphureus* (b) grown on three substrates. Sample points in the graphs are given as the arithmetic mean of  $\delta^{13}$ C-CH<sub>4</sub> or  $\delta^{13}$ C-CO<sub>2</sub> values with SD (n=3) on the y-axis and the arithmetic mean of the inverted mixing ratio of CH<sub>4</sub> or CO<sub>2</sub> with SD (n=3) on the x-axis.

The  $\delta^{13}$ C-CH<sub>4</sub> source signatures determined via a keeling Keeling plot analysis (Fig. 3) that are presented in Table 2 and range from -69.2  $\pm$  1.9 mUr (*L. sulphureus* grown on grass) to -39.8  $\pm$  2.0 mUr (*P. sapidus* grown on corn) are presented- in Table 2. Average  $\delta^{13}$ C-CH<sub>4</sub> source signatures for each fungal species, considering all three substrates, are -52.6 mUr for *P. sapidus* and -61.3 mUr for *L. sulphureus*. These results suggest that the fungal species significantly influence the isotopic values of the emitted CH<sub>4</sub> (p<0.001). A possible explanation for this observation could be the different enzyme sets of both fungi decomposing different components of the growth substrates, as *P. sapidus* belongs to white rot fungi and *L. sulphureus* is a brown rot fungus. However, detailed investigations of the metabolic pathways leading to CH<sub>4</sub> formation were beyond the scope of this study.

Furthermore, a significant effect of the growth substrate on δ<sup>13</sup>C-CH<sub>4</sub> source signatures was observed (*p*<0.001). δ<sup>13</sup>C-CH<sub>4</sub> source signatures by *P. sapidus* were more positive compared to those of *L. sulphureus* when grown on grass (Δ=16.3 mUr) and corn (Δ=13.6 mUr) (Fig. 4). When grown on pine wood, δ<sup>13</sup>C-CH<sub>4</sub> source signatures were similar with *P. sapidus* showing slightly more negative values (Δ=-3.9 mUr). Methane emitted by both fungi grown on corn was generally more enriched in <sup>13</sup>C (less negative δ<sup>13</sup>C-CH<sub>4</sub> source values) compared to the fungi grown on pine wood and grass. This might be easily explained by the δ<sup>13</sup>C values of the growth substrates corn (-11.7 mUr, typical for C<sub>4</sub>-plants) being roughly 20 mUr less negative in their δ<sup>13</sup>C values compared to the C<sub>3</sub>-plants pine wood (-28.0 mUr) and grass (-31.5 mUr).

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**Figure 4:** Calculated source signatures of  $\delta^{13}$ C-CH<sub>4</sub> values (a) and  $\delta^{3}$ C-CO<sub>2</sub> values (b) by *P. sapidus*, *L. sulphureus* and the  $\delta^{13}$ C values of the substrate. Values The data points are presented as represent the mean values of the individual keeling Reeling plots with SD (n=3).

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Comparison of calculated  $\delta^{13}C$ -CH<sub>4</sub> source signatures with measured bulk  $\delta^{13}C$  values of the substrates shows that CH<sub>4</sub> emitted by both fungi is generally depleted in  $^{13}C$  compared to the respective substrates (Fig. 4a). Based on this data we further calculated the apparent fractionation ( $\epsilon_{app}$  cH<sub>4</sub>) between the  $\delta^{13}C$ -CH<sub>4</sub> source signatures and the bulk  $\delta^{13}C$  values of the growth substrates. The apparent fractionation was calculated as up to the presentso far no metabolic pathway for the formation of CH<sub>4</sub> in fungi is known and therefore currently only the initial  $\delta^{13}C$  signatures of the substrates and the calculated  $\delta^{13}C$ -CH<sub>4</sub> source signatures of the fungi can be compared. The values of  $\epsilon_{app}$  cH<sub>4</sub> are presented in Table 2 and that range from -21.8 mUr (*P. sapidus* grown on grass) to -42.2 mUr (*L. sulphureus* grown on corn) are presented in Table 2. When grown on pine wood  $\epsilon_{app}$  cH<sub>4</sub> values are similar for *P. sapidus* (-38.4 ± 1.2 mUr) and *L. sulphureus* (-34.4 ± 0.6 mUr). The differences in  $\epsilon_{app}$  cH<sub>4</sub> values between both fungal species are more distinct when grown on grass (*P. sapidus*: -21.8 ± 1.7 mUr, *L. sulphureus*: -38,6 ± 2.0 mUr) and corn (*P. sapidus*: -28,5 ± 2.0 mUr, *L. sulphureus*: -42.2 ± 1.1 mUr).

The calculated  $\delta^{13}\text{C-CO}_2$  source signatures of both fungi (Table 2) range from -29.0 ± 0.5 mUr (*L. sulphureus* grown on grass) to -12.0 ± 0.3 mUr (*P. sapidus* grown on corn).  $\delta^{13}\text{C-CO}_2$  source signatures are in a similar range for both fungi for all three substrates. However, Although CO<sub>2</sub> emitted by *L. sulphureus* is slightly more depleted in <sup>13</sup>C for all three substrates compared to *P. sapidus*. Hence, an effect of fungal species on the stable carbon isotope values of CO<sub>2</sub> is significant (p=0.008). Also, the used substrates were found to influence  $\delta^{13}\text{C-CO}_2$  values significantly (p<0.001).

The  $\delta^{13}$ C-CO<sub>2</sub> source signatures of the fungi show only small deviations from the bulk  $\delta^{13}$ C values of the respective substrates (Fig. 4b). However, for both fungi grown on pine wood and grass,  $\delta^{13}$ C-CO<sub>2</sub> values are slightly less negative (a few mUr) compared to the bulk substrate. This observation is rather unexpected, as <u>usually</u>  $\delta^{13}$ C-CO<sub>2</sub> values are <u>usually</u> more negative

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with respect to  $\delta^{13}$ C values of growth substrates due to fractionation during the metabolism (Bowling et al., 2008). However, when grown on corn  $\delta^{13}$ C-CO<sub>2</sub> source signatures by both fungi are-slightly more negative compared to the substrate and calculated  $\varepsilon_{app\ CO2}$  values (Table 2) are -1.1 ± 0.3 mUr and +4.6 ± 1.3 mUr for *L. sulphureus* grown on corn and *P. sapidus* grown on grass, respectively.

The results of the incubation experiments show that there are distinct differences in the patterns of δ<sup>13</sup>C-CH<sub>4</sub> and δ<sup>13</sup>C-CO<sub>2</sub> solutions released by both fungi. While the δ<sup>13</sup>C-CO<sub>2</sub> source signatures are similar to the δ<sup>13</sup>C values of the substrate (with ε<sub>app</sub> CD<sub>2</sub> values up to 4.6 mUr), the δ<sup>13</sup>C-CH<sub>4</sub> source signatures deviate strongly from the respective substrate, with ε<sub>app CH4</sub> values of up to -42.2 mUr. This either indicates that metabolic pathways leading to the formation of CH<sub>4</sub> and CO<sub>2</sub> have different fractionation and/or that fungal CH<sub>4</sub> and CO<sub>2</sub> are derived from different precursor compounds of the respective substrate. The used growth substrates used for this study (pine wood, grass<sub>2</sub>-and-corn) consist of various components including mainlycontain distinct amounts of cellulose, hemicellulose<sub>3</sub>-and lignin and other compounds at different proportions (in contrast toif only using pure glucose or cellulose as growth substrate). Hence, the δ<sup>13</sup>C-CH<sub>4</sub> and δ<sup>13</sup>C-CO<sub>2</sub> source signatures might be dependent on the specific metabolic pathways of the substrate by the fungal species but also on as well as the chemical composition of the growth substrate. Therefore, we suggest that Tehe selected fungi and used growth substrates provide a first solid basis for the potential range of δ<sup>13</sup>C-CH<sub>4</sub> values that might occur in nature.

#### 325 3.4 Fungal δ<sup>13</sup>C-CH<sub>4</sub> values compared with known CH<sub>4</sub> sources

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Figure 5 summarizes compares the  $\delta^{13}$ C-CH<sub>4</sub> values emitted by fungi in relation to other known CH<sub>4</sub> sources in the environment that have been reported from the literature. The red bars indicate typical biogenic (formerly only considered to be produced by archaea) CH<sub>4</sub> sources with emissions from wetlands, ruminants, landfills and rice paddies where  $\delta^{13}$ C-CH<sub>4</sub> values are usually ranging from -85 mUr to -40 mUr. Abiotic CH<sub>4</sub> sources (including thermogenic or pyrolytic processes) stemming from natural gas, coal mining and biomass burning are characterized by less negative  $\delta^{13}$ C values usually ranging from -55 mUr to -20 mUr. In addition gas hydrates which might be formed by both microbial and abiotic processes cover a wider range of  $\delta^{13}$ C values (-29 mUr to -73 mUr), depending on its forming mechanisms (Kvenvolden, 1995). The  $\delta^{13}$ C source signatures of plant derived CH<sub>4</sub> have been reported to be in the range of -72 mUr to -45 mUr (Keppler et al., 2006; Vigano et al., 2009) depending on the three photosynthetic pathways (C<sub>3</sub>, C<sub>4</sub> or CAM). Furthermore, there is a tendency towards more negative  $\delta^{13}$ C-CH<sub>4</sub> values when the respective plant was treated with UV radiation (Vigano et al., 2009).  $\delta^{13}$ C-CH<sub>4</sub> source signatures of humans which might include both formation by microbes in the gut but also from cellular processes show a rather wide range with values between -95 mUr and -56 mUr (Keppler et al., 2016). The results of our experiments conducted with two fungal species and three different growth substrates provide a range of  $\delta^{13}$ C-CH<sub>4</sub> source values from -69 mUr to -40 mUr. This range overlaps with other eukaryotic sources, most microbial CH<sub>4</sub> sources and even some abiotic CH<sub>4</sub> sources such as natural gas or emissions from coal mining.

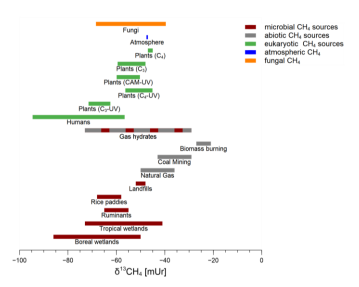


Figure 5: Range of  $\delta^{13}$ C-CH<sub>4</sub> values of microbial CH<sub>4</sub> sources (red), abiotic CH<sub>4</sub> sources (grey), eukaryotic CH<sub>4</sub> sources (green), atmospheric CH<sub>4</sub> (blue) and fungal CH<sub>4</sub> from this study (orange). The red and grey dashed bar indicates a mixture of microbial and abiotic CH<sub>4</sub> formation processes for gas hydrates (Kvenvolden, 1995). Data taken from (Brownlow et al., 2017; Keppler et al., 2006, 2016; Kvenvolden, 1995; Nisbet et al., 2016; Quay et al., 1999; Vigano et al., 2009).

#### 4 Conclusion

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This study provided the first analysis of stable carbon isotope values of  $CH_4$  emitted by two saprotrophic fungi that were grown on three different substrates.  $\delta^{13}C$ - $CH_4$  and  $\delta^{13}C$ - $CO_2$  source values were found to be dependent on the fungal species, as well as the substrates decomposed by the fungi.  $\delta^{13}C$ - $CH_4$  source values of the fungi were found to be in the range of -69 mUr to -40 mUr and therefore overlap with  $\delta^{13}C$ - $CH_4$  values reported for other  $CH_4$  sources such as methanogenic archaea, eukaryotes and from abiotic processes $CH_4$  sources (e.g. natural gas, coal mining). Stable carbon isotope values of  $CH_4$  in combination with flux measurements are often applied for a better understanding of regional and global  $CH_4$  cycling. However, in recent years it has become clear that many biogenic  $CH_4$  sources include complex  $CH_4$  formation processes, resulting in different isotopic fractionation patterns depending on several biochemical and abiotic factors. Thus, studying ecosystems in which more than one major  $CH_4$  source has to be expected (e.g. methanogenic archaea, fungi, cyanobacteria or plants) gets increasingly complicated as distinguishing between each individual source solely by stable carbon isotope values might be highly

challenging. Therefore, additional tools are needed to better identify the sources but also to disentangle sources and sinks. In future research, the stable hydrogen isotopic values of  $CH_4$  ( $\delta^2H$ - $CH_4$  values) or even applications of clumped isotopes might prove suitable tools for better distinction between different  $CH_4$  sources and thus to better constrain the global  $CH_4$  budget.

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<u>Data availability</u>. We provide the data in heiDATA, which is an institutional repository for research data of Heidelberg <u>University (https://doi.org/10.11588/data/DQYPMC)</u>

- 365 Author contributions. MS, KL, and FK conceived the study and designed the experiments; HZ provided fungal cultures, MS performed the experiments under the supervision of FK and KL; CE helped with gas measurements; MG measured stable isotope values of greenhouse gases; MS, FK, and KL analysed the data; MS, FK, HZ, MG, and KL, discussed the results, and MS, KL and FK wrote the paper.
- 370 Competing interests. The authors declare that they have no conflict of interest.

Acknowledgments. We thank Anette Giesemann for analytical measurements of stable carbon isotope values of the bulk substrates. We are grateful to Bianka Daubertshäuser for technical support with the cultivation of the fungi and to Lukas Kohl for encouraging us to perform this study. We acknowledge financial support by the Deutsche Forschungsgemeinschaft.

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Financial support. This research has been supported by the <u>German Research Foundation (DFGeutsche Forschungsgemeinschaft (grant nosGrant Numbers</u>. KE 884/8-2, KE 884/16-2) and (LE3381/1-1).

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