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1 **Abstract**

2 Many studies have demonstrated the importance of symbiotic microbial communities for the host
3 with beneficial effects for nutrition, development, and the immune system. The majority of these
4 studies have focused on bacteria residing in the gastrointestinal tract, while the fungal community
5 has often been neglected. Gut anaerobic fungi of the class Neocallimastigomycetes are a vital part of
6 the intestinal microbiome in many herbivorous animals and their exceptional abilities to degrade
7 indigestible plant material means that they contribute significantly to fermentative processes in the
8 enteric tract. Gorillas rely on a highly fibrous diet and depend on fermentative microorganisms to
9 meet their daily energetic demands. To assess whether Neocallimastigomycetes occur in gorillas we
10 analyzed 12 fecal samples from wild Western lowland gorillas (*Gorilla gorilla gorilla*) from Dzanga-
11 Sangha Protected Areas, Central African Republic, and subjected potential anaerobic fungi sequences
12 to phylogenetic analysis. The clone library contained ITS1 fragments that we related to 45 different
13 fungi clones. Of these, 12 gastrointestinal fungi in gorillas are related to anaerobic fungi and our
14 phylogenetic analyses support their assignment to the class Neocallimastigomycetes. As anaerobic
15 fungi play a pivotal role in plant fiber degradation in the herbivore gut, gorillas might benefit from
16 harboring these particular fungi with regard to their nutritional status. Future studies should
17 investigate whether Neocallimastigomycetes are also found in other non-human primates with high
18 fiber intake, which would also benefit from having such highly efficient fermentative microbes.

19

20 Keywords: gut microbiome, Neocallimastigales, gorillas, diet

21

22 **Introduction**

23 Symbiotic microbial communities residing in the intestinal tract, referred to as the gut microbiome,
24 are assemblages of bacteria, fungi, protozoa, and archaea that provide crucial functions for host
25 nutrition (e.g. Sekirov et al., 2010; Robert & Bernalier-Donandille, 2003), development (e.g. McFall-
26 Ngai, 2002), and immune systems (e.g. Hooper et al., 2012; Round & Mazmanian, 2009). Since many

27 microbes collected from environmental samples are uncultivable (Torsvik & Ovreas, 2002), advances
28 in culture-independent methods, particularly metagenomic approaches based on high-throughput
29 sequencing, allow the detection of a far more detailed microbial diversity than traditional culture
30 based approaches (e.g. Caporaso et al., 2012). These methods have led to an increased
31 understanding of the factors shaping the composition of microbial communities. There is common
32 agreement that the two main factors influencing the microbial community structure are host
33 phylogeny and diet (e.g. Sanders et al., 2014; Muegge et al., 2011). For example, a study investigating
34 the gut microbiome of 60 different mammal species shows that conspecifics harbor bacterial
35 communities more similar to each other than to those of a different host species and that these
36 communities cluster according to host taxonomy. Principal coordinates analyses also provide
37 evidence for the significant impact of diet on gut microbiome structure, because bacterial
38 communities cluster in accordance with diet and gut type (Ley et al., 2008).

39

40 Neocallimastigomycetes are obligate anaerobic fungi that were first isolated in ruminants (Orpin,
41 1975). Their occurrence has also been confirmed in various non-ruminant herbivores like African
42 elephants (*Loxodonta africana*), horses (*Equus ferus caballus*), black rhinoceroses (*Diceros bicornis*),
43 red kangaroos (*Macropus rufus*) and in the herbivorous green iguana (*Iguana iguana*) (Nicholson et
44 al., 2010; Ligginstoffer et al., 2010; Mackie et al., 2004). Intestinal anaerobic fungi are remarkable in
45 their capacities to degrade plant material that is indigestible by the host. They harbor highly efficient
46 hydrolases (cellulases, xylanases, mannosases, esterases, glucosidases, and glucanases) aggregated in
47 extracellular enzyme-complexes, termed cellulosomes. These fungal enzymes are assumed to exceed
48 the fermentative capacities of bacterial enzymes (Lee et al., 2000). Additionally, anaerobic fungi are
49 among the first to colonize plant fragments (Edwards et al., 2008) and are able to mechanically
50 penetrate plant cell walls (Doi & Kosugi, 2004; Fontes & Gilbert, 2010). Due to this initial colonization
51 of plant particles and the mechanical breakdown of large plant particles as well as plant cell walls

52 anaerobic fungi facilitate the accessibility to fermentable substrates for residential bacteria that take
53 part in the hydrolysis of plant fiber in the gastrointestinal tract (Bauchop, 1981).

54

55 Currently, Neocallimastigomycetes include one order, Neocallimastigales, with one family
56 (Neocallimastigaceae) that encompasses six long known genera (*Neocallimastix*, *Caecomyces*,
57 *Orpinomyces*, *Piromyces*, *Anaeromyces*, and *Cyllamyces*) and three newly described genera
58 (*Buwchfawromyces*: Callaghan et al., 2015; *Oontomyces*: Dagar et al., 2015 and *Pecoromyces*: Hanafy
59 et al., 2017). However, studies of various herbivorous animals propose a revised taxonomy with
60 several new groups (Tuckwell et al., 2005; Fliegerová et al., 2010; Liggenstoffer et al., 2010; Nicholson
61 et al., 2010; Herrera et al., 2011; Kittelmann et al., 2012). Studies suggest that the abundance and
62 composition of different anaerobic fungi genera are dependent on host taxonomy, type of gut
63 fermentation, and fiber content in the diet (Liggenstoffer et al., 2010; Kumar et al., 2013; Denman et
64 al., 2008).

65

66 Despite the growing number of studies investigating the gut microbiome in primates, the fungal
67 community has received disproportionately little attention. Many early studies focused on specific
68 mycotic infections (reviewed in Migaki et al., 1982), and a more recent study targeted a broader
69 diversity of enteric fungi in Western lowland gorillas (*Gorilla gorilla gorilla*). This molecular survey of
70 pathogenic eukaryotes detected 52 fungal species, all belonging to the taxa Ascomycota and
71 Basidiomycota (Hamad et al., 2014). However, no study has yet investigated Neocallimastigomycetes
72 in primates, even though there is good reason to hypothesize that some primates harbour these
73 fungi. Most primates rely on a mainly plant based diet (Chapman & Chapman, 1990), yet, like all
74 mammals, they lack the enzymes to degrade plant structural polysaccharides themselves and thus
75 rely on endosymbiotic microorganisms for an adequate nutritional intake (Mackie, 2002).

76

77 Studies of gorilla feeding ecology reveal that they consume high fiber staple and filler fallback foods
78 such as terrestrial herbaceous vegetation, figs, bark, and pith year-round (Western lowland gorillas:
79 Remis, 2003; Doran-Sheehy et al., 2009). Although chimpanzees (*Pan troglodytes*) also consume high
80 fiber plant material such as pith in times of fruit scarcity (Wrangham et al., 1991), there is strong
81 support for the hypothesis that chimpanzees can maintain a higher quality diet with overall less fiber
82 intake when compared to Western lowland gorillas (Tutin et al., 1991; Wrangham et al., 1998). In line
83 with these observations, gorillas show morphological and physiological adaptations that suggest
84 heavy reliance on high fiber foods. For example, their molar morphology indicates a high capacity for
85 processing tough food (Ungar et al., 2007). Further, gorillas have an enlarged colon surface area and a
86 longer mean gut retention time when compared to less folivorous chimpanzees (Chivers & Hladik,
87 1980; Milton & Demment, 1988; captive Western lowland gorillas: Remis & Dierenfield, 2004) even
88 when accounting for body mass (Harrison & Marshall, 2011). Moreover, daily energy consumed that
89 potentially originates from microbial fermentation in the hindgut is an estimated 57.3 % for western
90 lowland gorillas and 24.7 % for chimpanzees (Popovich et al., 1997; Conklin-Brittain et al., 2006).
91 Gorillas further fulfill two major prerequisites for the potential of harboring anaerobic fungi: a
92 dedicated enlarged digestive chamber for microbial fermentation (hindgut) and a relatively long
93 retention time for plant material.

94

95 We explore fungal communities in feces of wild Western lowland gorillas using culture-independent
96 molecular methods. Specifically, we aim to amplify ITS1 rDNA fragments of Neocallimastigales from
97 DNA isolated from fecal samples. Given their year-round exploitation of high-fibrous foods, we
98 hypothesize that gorillas benefit from harboring highly efficient fermentative microorganisms such as
99 anaerobic fungi in their intestinal tract. Based on their digestive morphology, we predict that it is very
100 likely that Neocallimastigales are part of the gorilla gut microbiome.

101

102 **Methods**

103 ***Study site, subjects and sample collection***

104 We collected fecal samples from two habituated groups of wild Western lowland gorillas at two field
105 sites: Bai Hokou and Mongambe in Dzanga-Ndoki National Park, Dzanga-Sangha Protected Areas,
106 Central African Republic, from September 2014 to January 2015. Both field sites comprise semi-
107 deciduous forests and are characterized by seasonal variations in rainfall with a dry season lasting
108 from December to February (for detailed description see Masi, 2007). We collected samples from
109 known individuals as soon as possible after defecation, i.e. as soon as it was safe to collect the sample
110 without disturbing the animal, which was usually within minutes.

111

112 We fixed fecal material in 96% ethanol in 8 ml tubes (approximate ratio 2/3 ethanol to 1/3 sample
113 material) and stored the samples at ambient temperature at the field sites until we transported them
114 to the University for Veterinary Medicine and Pharmaceutical Sciences, Brno, Czech Republic, where
115 we kept them in ethanol at -20°C until analysis. We preserved fecal material in ethanol due to the
116 lack of other storage possibilities at the field sites. DNA has been successfully isolated and amplified
117 from such fixed samples (Frantzen et al., 1998; Hale et al., 2015) and preserving samples in highly
118 concentrated ethanol at ambient temperatures appears to have little influence on the microbial
119 community (Song et al., 2016).

120

121 Our study is a preliminary investigation for which we processed 12 gorilla samples, representing 11
122 individuals. We picked gorilla samples randomly from the samples we collected during the study.

123

124 ***Sample Processing***

125 **DNA Isolation.** After evaporating ethanol at 40°C (heat block) overnight, we isolated DNA from the
126 fecal material with the FastDNA™ Spin Kit for Soil (MP Biomedicals, USA) according to the
127 manufacturer's protocol with the following changes: to break fungi chitin walls, we homogenized the
128 sample by bead-beating it three times for 30 sec at 6 m/s with 30 sec on ice between homogenization

129 steps (Cheng et al., 2009). We eluted DNA with 70 µl instead of 100 µl of the elution solution
130 provided with the kit and stored eluates at -20°C.

131

132 **PCR Amplification of Fungal Barcodes.** We used the fungal universal forward primer ITS1F
133 (CTTGGTCATTTAGAGGAAGTAA) in combination with a primer specific for anaerobic fungi NeoQ PCR R
134 (GTGCAATATGCGTTCTGAAGATT) to amplify ITS1 fragments (Fliegerová et al., 2010). We prepared PCR
135 reactions with a final volume of 25 µl using the QIAGEN Multiplex PCR Kit (Qiagen, Germany)
136 containing 12.5 µl Master Mix, 8.0 µl dH₂O, 2.5 µl dye 0.01 µM of each primer and 1 µl DNA. We set
137 cycling conditions of the touchdown PCR protocol as 95°C for 5 min; 20 cycles consisting of 95°C for
138 30 sec, 60.5°C for 30 sec with -0.2°C per cycle, 72°C for 30 sec; followed by another 20 cycles
139 consisting of 95°C for 30 sec, 57°C for 30 sec, 72°C for 30 sec and a final extension of 5 min at 68°C.
140 We visualized PCR products on 1% agarose gels and subjected fragments of expected size to cloning
141 procedure after purification with ExoSap (Affymetrix Inc., USA).

142

143 **Cloning Library Construction.** We constructed a clone library with the TOPO TA Cloning Kit for
144 Sequencing (Life Technologies, USA) following the manufacturer's protocol for vector preparation and
145 the transformation of competent *E. coli* cells. We picked 289 clone colonies and transferred them into
146 20 µl PCR H₂O to screen them for the presence of the insert by PCR. We prepared PCR reaction
147 mixtures of 25 µl containing 12.5 µl Master Mix (PCR BIO Taq Mix Red, PCR Biosystems, UK), 9.5 µl
148 dH₂O, 1 µl of clone colony solution and 0.01 µM of ITS1F and NeoQ PCR R primers. We set cycling
149 conditions for ITS1 insert amplification as 95°C for 5 min, followed by 30 cycles of 95°C for 30 sec,
150 55°C for 30 sec, 72°C for 30 sec and a final elongation for 5 min at 72°C. We checked PCR products
151 using gel electrophoresis, purified products of the right length with ExoSap and subjected them to
152 Sanger sequencing (Macrogen Europe, The Netherlands).

153

154 **Sequence Analysis**

155 We first edited sequences with BioEdit software (version 7.2.3) and subsequently used GenBank's
156 Basic Local Alignment Search Tool (BLAST; default setting highly similar sequences (megablast)) to
157 identify their nearest relatives. We only subjected sequences that we could relate to anaerobic fungi
158 to further analysis. Given that sequence similarity among different anaerobic fungi strains can be very
159 high (Goudarzi et al., 2015) we first aligned a selection of 12 clone sequences, as representatives for
160 all related anaerobic fungi strains, to assess their resemblance (ClustalX, Bioedit; Hall, 1999; Table 1,
161 Appendix S1). We subsequently chose a subset of the nine most divergent sequences for
162 phylogenetic analysis to determine the taxonomic relationships of potential ape anaerobic fungi
163 strains with known Neocallimastigales. By applying the MAFFT algorithm with default settings (online
164 version 7, ©Kato, 2013) we computed alignments that included ITS1 fragments generated in this
165 study and reliable ITS1 sequences representing the improved taxonomic framework for
166 Neocallimastigales fungi (Kittelmann et al., 2012; Dagar et al., 2015; Appendix S2). In addition to
167 these reference sequences classified as Neocallimastigales we included the uncultured *fungus clone*
168 *AFI-1* sequence isolated from Bactrian camel (*Camelus bactrianus*) rumen (Acc. No: JX944983). High
169 degrees of sequence dissimilarities and length polymorphisms between Neocallimastigales genera
170 resulted in multiple large gaps in the original 452 bp alignment. Given that the applied Maximum
171 Likelihood algorithm treats gaps like missing data we aimed to reduce ambiguity by manually deleting
172 those gaps to different degrees, resulting in two further alignments, one of 241 bp and another of
173 only 197 bp.

174

175 We constructed phylogenetic trees in PhyML (Guindon et al., 2010) based on the original MAFFT
176 alignment and two further alignments. Based on the results of Modeltest 3.7 (Posada & Crandall,
177 1998), we used the GTR+G substitution model for tree calculation using maximum likelihood for the
178 unedited alignment and computed phylogenies based on the two manually edited alignments under
179 the HKY+G model. We also constructed a ML tree under the T92+G (Tamura, 1992) model in MEGA 6

180 (Tamura et al., 2013) to account for uneven GC content in our sequences. We based bootstrap
181 analyses for each tree on 1000 pseudo-replicates.

182

183 **Ethical Note**

184 We collected all gorilla samples non-invasively and with no harm to the study subjects. Permission to
185 conduct research in the Dzanga-Sangha Protected Areas was granted by the Ministère de
186 l'Enseignement Supérieur et de la Recherche Scientifique and the Ministère des Eaux, Forêts,
187 Chasses, Pêches, chargé de l'Environnement.

188

189 **Results**

190 *Fungal diversity*

191 We analyzed 238 clones with inserts of appropriate length from the clone libraries of amplified ITS1
192 fragments. The sequences we generated were associated with 45 different fungal rDNA sequences
193 deposited in GenBank. Of the 238 clones we obtained, 78 were moderately similar to 12 different
194 uncultured Neocallimastigales clones. These potential anaerobic fungi ITS1 fragments originated from
195 8 of 12 processed samples, with sequences similar to the Uncultured Neocallimastigales clone Iguana
196 01BMIEK (Acc. No. GQ843155) being the most abundant and the only one that occurs in all 8
197 samples. Other prospective anaerobic fungi ITS1 fragments that we amplified fit with uncultured
198 Neocallimastigales clones detected in hindgut-fermenting Equidae, ruminant Bovidae, and the
199 pseudo-ruminant hippopotamus (Table 1; Appendix S1).

200

201 The remaining fungal ITS1 fragments from gorillas that we cannot associate with anaerobic fungi
202 clones are linked to sequences of the fungal classes Ascomycota and Basidiomycota (Appendix S1).
203 These clones comprise 33 sequences that are related to 15 different strains of Ascomycota with
204 moderate to high similarities (91 – 100%), covering five known orders and three strains of unclassified
205 Ascomycota. Another four sequences that we obtained show high similarities (96 – 100%) with three

206 different Basidiomycota strains, belonging to three orders. According to BLAST analysis the majority
 207 of our ITS1 fragments are identified as unclassified fungal clones. In total, our sequences are related
 208 to 13 different such unclassified fungal clones that have been isolated from plant tissues, soil, reactor
 209 bio-filter, and woodpecker excavation with similarities ranging 96 – 100% (Appendix S1). An
 210 additional unclassified fungal clone (*Uncultured fungus clone AFI-1*; Acc. No. JX944983, unpublished
 211 sequence) to which 11 of our sequences are highly similar has been isolated from Bactrian camel
 212 (*Camelus bactrianus*) rumen. Finally, one sequence does not match with any of the rDNA sequences
 213 deposited in online data bases.

214

215 *Phylogenetic analysis of anaerobic fungi*

216 Our initial alignments revealed high degrees of resemblance among the potential anaerobic fungi
 217 sequences we obtained from gorilla feces, although they were associated with different uncultured
 218 Neocallimastigales clones (Table 1).

219

220 **Table 1:** Nearest relatives of ITS1 sequences retrieved from Western lowland gorilla feces collected at
 221 the sites Bai Hokou and Mongambe from September 2014 to January 2015.

222 UNC – Uncultured Neocallimastigales clone

223 * Not classified as Neocallimastigales fungus in NCBI (National Center for Biotechnology Information)
 224 sequence database.

Sequence ID (date sample collection)	Field site	Size [bp]	GenBank Accession Number	Nearest relative [Accession Number]	Sequence similarity
Mak_2 (23.10.2014)	Bai Hokou	213	KY697108	UNC NileLechwe03FKYBS [GQ592255]	90
Mal_1 (29.11.2014)	Bai Hokou	283	KY697116	UNC HorseTopper01A6QWL [GQ688452]	89
Mob_11 (12.09.2014)	Bai Hokou	264	KY697114	UNC HorseBug01B20BM [GQ829356]	88
Mob_22 (12.09.2014)	Bai Hokou	279	KY697115	UNC Iguana01BLGEC [GQ843065]	88

Won_5 (01.12.2014)	Mongambe	260	KY697113	UNC Iguana01BMIEK [GQ843155]	90
May_19 (20.09.2014)	Mongambe	253	KY697112	UNC GrantsGazelle02CZ47B [GQ784902]	88
Mob2_2 (27.09.2014)	Bai Hokou	242	KY697109	UNC PigmyHippopotamus03GM37B [GQ607513]	89
Mop_14 (17.10.2014)	Mongambe	243	KY697110	UNC Iguana01A3GEE [GQ842869]	89
Map_14 (24.11.2014)	Mongambe	244	KY697111	Uncultured fungus clone AFI-1 [JX944983]*	100

225

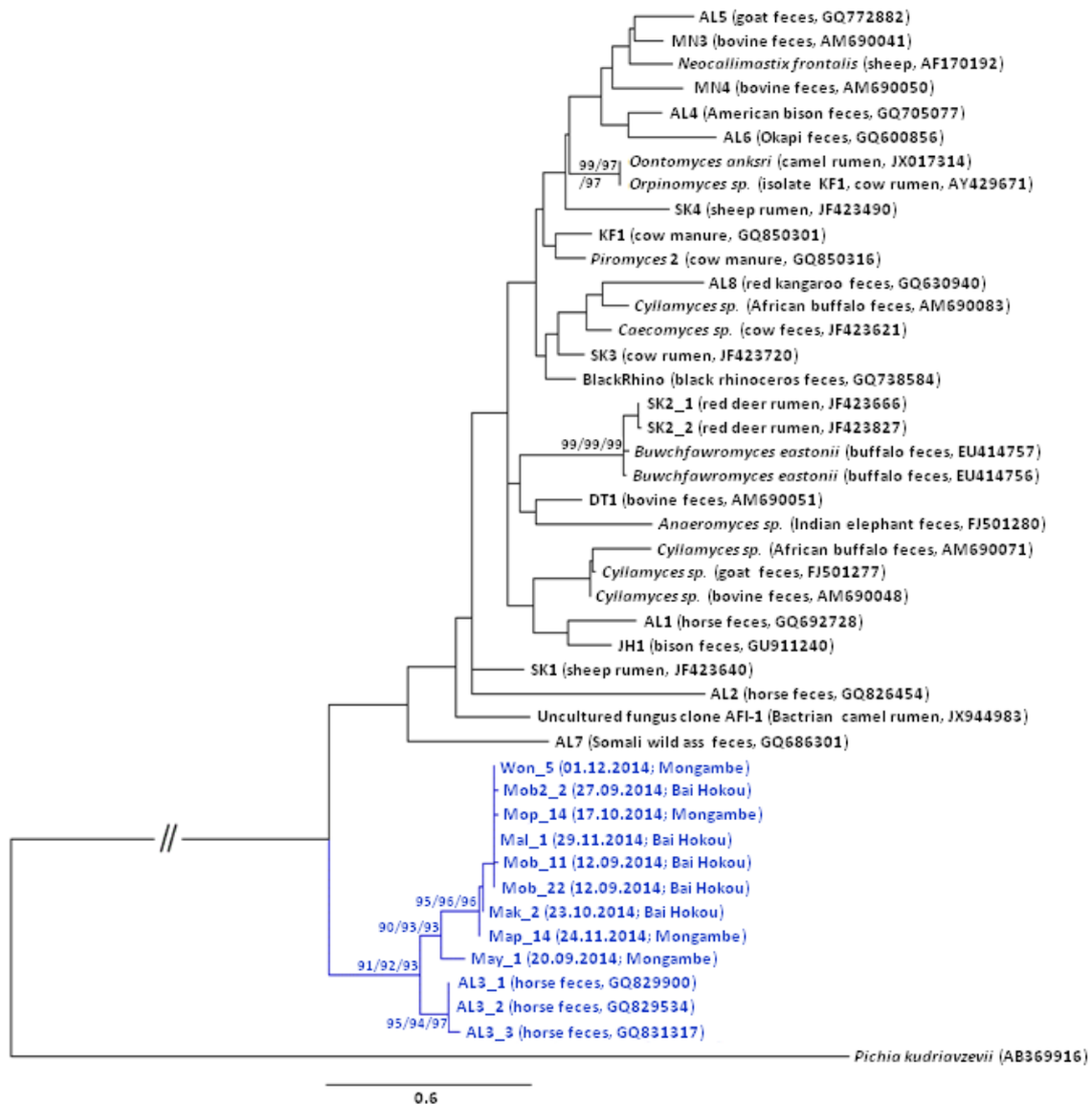
226 In the Maximum Likelihood tree based on the original 452 bp alignment, our ITS1 fragments form a
 227 separate clade which clusters with the clade of the newly described uncultured anaerobic fungi group
 228 AL3 (group NG3 in Liggenstoffer et al., 2010) with significant support (Figure 1). This phylogenetic
 229 relationship is also supported in two other phylogenies that we constructed from 241 bp and 197 bp
 230 alignments. All other reference ITS1 sequences cluster in an unsupported monophyletic clade in
 231 which most of the phylogenetic relationships between the different groups and genera are rather
 232 weakly supported.

233

234 The Maximum Likelihood tree constructed under T92 + G substitution model, which accounts for
 235 uneven CG content in sequences, revealed very similar results for the sequence clustering. Again,
 236 fungal clones obtained from gorilla feces grouped with AL3 references with adequate support
 237 (bootstrap value 82; data not shown). However, as in the other three phylogenies, relationships
 238 between the reference sequences of known Neocallimastigales lack significant support.

239

240 **Fig 1:** Phylogenetic relationships of potential gorilla anaerobic fungi sequences in the order of
 241 Neocallimastigales fungi based on Maximum Likelihood. Bootstrap support above 50% is indicated at
 242 nodes for the 452, 241 and 197 bp alignments. Clones obtained in our study and reference sequences
 243 are listed in Table 1 and S2. Dates of sample collection and field site for sequences from gorilla
 244 samples are given in brackets.



245

246 Discussion

247 Our results suggest that anaerobic gut fungi are part of the gorilla gut microbiome. The assignment of
 248 the ITS1 sequences we analyzed as a sister clade to the novel Neocallimastigales lineage AL3 is
 249 significantly supported. Despite the highly significant support for the hypothesis that some of our
 250 gorilla gut fungi belong to the class Neocallimastigomycetes, two factors warrant some caution. First,
 251 fungal ITS1 sequences that we obtained from gorilla feces were only moderately similar to known
 252 Neocallimastigales sequences deposited in the GenBank database. However, new lineages and
 253 species of Neocallimastigales are constantly discovered (Ariyawansa et al., 2015; Hanafy et al., 2017).
 254 Thus, our sequences might represent a new anaerobic fungi lineage. Second, our amplified ITS1

255 fragments were very short. This in combination with the known high variation in the
256 Neocallimastigales ITS1 region (Edwards et al., 2017) limits the reliability of constructed alignments
257 and phylogenies.

258

259 Like in other rapidly evolving non-coding regions insertions – deletions (indels) accumulate over time
260 in the ITS1 sequence. These indels are thought to be more conserved than base substitutions and
261 thus can provide a reliable source of information for phylogenetic reconstructions (Matheny et al.,
262 2006; Abarenkov et al., 2010). Alignment gap deletion decreases tree resolution, particularly when
263 sophisticated alignment algorithms such as MAFFT are applied (Nagy et al., 2012). Our phylogenetic
264 analysis based on alignment without gap removal significantly supports the close relationship of
265 gorilla gut fungi with the anaerobic fungi group AL3. Following the logic that alignment gaps can
266 provide phylogenetic information, this result supports our assignment of gorilla gut fungi to the class
267 of Neocallimastigales. The low bootstrap values in our phylogeny might be the result of difficulties
268 aligning anaerobic fungi sequences given the significant sequence dissimilarities and length
269 polymorphisms between genera (Nicholson et al., 2010). However, our goal was to determine
270 whether anaerobic fungi occur in wild gorillas rather than resolving the Neocallimastigales phylogeny.
271 Our sequences are very closely related to the anaerobic fungi group AL3. This group of
272 Neocallimastigales was first detected in hindgut fermenting equids which have similar digestive
273 physiology to gorillas. Since digestive physiology is a key factor determining anaerobic fungi
274 community structure (Liggenstoffer et al., 2010) it is likely that even distantly related herbivorous
275 animals harbor similar Neocallimastigales strains. This finding, therefore, provides additional support
276 for our hypothesis that Neocallimastigales are part of the gorilla gut microbiome.

277

278 While our analysis suggests that Neocallimastigales reside in the gastrointestinal tract of gorillas, we
279 have no indication so far that other African great apes harbor anaerobic fungi (unpublished data:
280 chimpanzee fecal samples, analyzed by D. Schulz). We predicted that anaerobic fungi are a part of the

281 gorilla gut microbiome based on gorilla diet and digestive physiology. Western lowland gorillas,
282 although more frugivorous than mountain gorillas (*Gorilla gorilla beringei*), consume high fiber foods
283 throughout the year (Rothman et al., 2008; Remis et al., 2001). The occurrence of anaerobic fungi in
284 gorillas could therefore be interpreted as an adaptation to a high fiber diet. Along with other adaptive
285 morphological and physiological digestive features (Harrison and Marshall, 2011) this might enable
286 gorillas to survive on a low quality diet (Tutin et al., 1991). Other non-human primates that similarly
287 rely on a highly or even strictly leafy diet could likewise benefit from harboring anaerobic fungi in
288 their intestines. This remains to be investigated.

289

290 Gorillas fall back on more low-quality foods in periods of low preferred fruit abundance and in
291 general consume much more fiber than chimpanzees (Wrangham et al, 1998, Tutin et al., 1991).
292 Further, chimpanzees have smaller fiber digestions coefficients and their fecal microbial communities
293 have diminished fiber degradation capacities compared to gorillas (Popovich et al., 1997; Conklin-
294 Brittain et al., 2006; Kišidayová et al., 2009). Neocallimastigales play a pivotal role in digesting
295 structural polysaccharides, particularly with regard to their ability to enhance access to fermentable
296 substrate for hydrolyzing bacteria. Thus, the higher fiber degradation capacities of the gorilla gut
297 microbiome might be the consequence of higher rates of bacterial fermentation facilitated by
298 anaerobic fungi. However, given the limitations of sampling and methodology in our study, we draw
299 this conclusion only cautiously.

300

301 Similar to the findings of a previous study (Hamad et al., 2014), we detected several Ascomycota and
302 Basidiomycota strains in our gorilla samples. There is no concordance on the species level between
303 Ascomycota strains we obtained and clones isolated by Hamad et al. (2014). However, four
304 (Eurotiales, Hypocreales, Saccharomycetales, and Capnodiales) of six genera found by Hamad and
305 colleagues are also present in our samples. Our results for Basidiomycota differ greatly from
306 previously isolated strains in gorillas. While we isolated only four strains, it seems that the diversity of

307 Basidiomycota in the colonic fungal community of gorillas is actually far greater (Hamad et al., 2014).
308 While some of the Basidiomycota strains detected in gorillas are human pathogens, a few of the
309 identified Ascomycota, namely members of the order *Saccharomycetales* that are usually associated
310 with plants, possess fermentative capacities (Hamad et al., 2014). It is, however, unclear whether
311 these aerobic fungi constitute transients passing through the enteric tract with food particles or if
312 they are residents and part of the gut microbiome with benefits for the host. We find the latter
313 explanation unlikely due to the low redox potential of the anaerobic conditions in the intestinal tract
314 (Espey, 2013).

315

316 In conclusion, our analyses provide evidence that Neocallimastigales is part of the gorilla gut
317 microbiome. Our results emphasize the need to include enteric fungi when investigating the
318 composition of the primate gut microbiome and we suggest that more research is needed to improve
319 our understanding of the role of enteric fungi in the digestive tract. More extensive studies of fungal
320 communities of several wild primate populations employing next generation sequencing techniques
321 is warranted to enhance our knowledge of how differences in the fungal gut microbiome reflect
322 differences in host diet and distribution. The results of such studies will contribute significantly to our
323 understanding of the complexity of primate microbiomes and their adaptive values.

324

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339

340 **Supplementary Material**

341 Intestinal fungi strains identified and similarity with amplified ITS1 fragments (Appendix S1) and an-
342 aerobic fungi reference sequences (Appendix S2) and all sequences included in phylogenetic analysis
343 (Appendix S3) are available online. If reasonable we will grant all further data requests from interest-
344 ed researchers.

345

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