Phylogenomic analysis of the Neocallimastigomycota: Proposal of Caecomycetaceae fam. nov., Piromycetaceae fam. nov., and emended description of the families Neocallimastigaceae and Anaeromycetaceae

Radwa A. Hanafy^{1,2†}, Yan Wang^{3,4†}, Jason E. Stajich⁵, Carrie J. Pratt¹, Noha H. Youssef¹, and Mostafa H. Elshahed^{1*}

1Department of Microbiology and Molecular Genetics, Oklahoma State University, Stillwater, OK.
2Department of Chemical & Biomolecular Engineering, University of Delaware, Newark DE, USA
3Department of Ecology & Evolutionary Biology, University of Toronto, Toronto, ON M5S 3B2, Canada
4Department of Biological Sciences, University of Toronto Scarborough, Toronto, ON M1C 1A4, Canada
5Department of Microbiology and Plant Pathology, University of California, Riverside, Riverside, CA.

1
 2
 3

[†]These authors contributed equally to this work.

21

22

23

24

25

26

27

28

29

30

31

32

33

34

35

36

37

38

39

40

41

42

Abstract. The anaerobic gut fungi (AGF) represent a coherent phylogenetic clade within the Mycota. Twenty genera have been described so far. Currently, the phylogenetic and evolutionary relationships between AGF genera remain poorly understood. Here, we utilized 53 transcriptomic datasets from 14 genera to resolve AGF inter-genus relationships using phylogenomics, and to provide a quantitative estimate (amino acid identity) for intermediate rank assignments. We identify four distinct supra-genus clades, encompassing genera producing polyflagellated zoospores, bulbous rhizoids, the broadly circumscribed genus *Piromyces*, and the Anaeromyces and affiliated genera. We also identify the genus Khoyollomyces as the earliest evolving AGF genus. Concordance between phylogenomic outputs and RPB1 and D/D2 LSU, but not RPB2, MCM7, or ITS1, phylogenies was observed. We combine phylogenomic analysis, and AAI outputs with informative phenotypic traits to propose accommodating 13/20 AGF genera into four families: Caecomycetaceae fam. nov. (encompassing genera Caecomyces and Cyllamyces), Piromycetaceae fam. nov. (encompassing the genus Piromyces), emend the description of fam. Neocallimastigaceae to only encompass genera Neocallimastix, Orpinomyces, Pecramyces, Feramyces, Ghazallomyces, and Aestipascuomyces, as well as the family Anaeromycetaceae to include the genera Oontomyces, Liebetanzomyces, and Capellomyces in addition to Anaeromyces. We refrain from proposing families for the deeply branching genus Khovollomyces, and for genera with uncertain position (Buwchfawromyces, Joblinomyces, Tahromyces, Agriosomyces, Aklioshbomyces, and Paucimyces) pending availability of additional isolates and sequence data. Our results establish an evolutionarygrounded Linnaean taxonomic framework for the AGF, provide quantitative estimates for rank assignments, and demonstrate the utility of RPB1 as additional informative marker in Neocallimastigomycota taxonomy.

44

45

46

47

48

49

50

51

52

53

54

55

56

57

58

59

60

61

62

63

64

65

Introduction Members of the anaerobic gut fungi (AGF) represent a phylogenetically, metabolically, and ecologically coherent clade in the kingdom Mycota [1]. Twenty genera and thirty-six different species have been described so far [2]. A recent review has provided detailed description of current genera and resolved historical inaccuracies and synonymies within the Neocallimastigomycota [2]. Further, criteria for the identification and characterization, as well as guidelines for genus- and species-level rank assignment for novel AGF isolates have recently been formulated [3]. In spite of such progress, the phylogenetic and evolutionary relationships between various genera within the Neocallimastigomycota are currently unclear. Similarities in specific microscopic traits (zoospore flagellation, thallus development, and rhizoidal growth patterns) across genera have been identified; and the significance of using such traits for proposing higher order relationship has been debated [4-6]. As well, phylogenetic analysis using two ribosomal loci: the internal transcribed spacer region 1 (ITS1) and D1/D2 region of the large ribosomal subunit (D1/D2 LSU) has yielded multiple statistically-supported supra-genus groupings, although such topologies were often dependent on the locus examined, region amplified, taxa included in the analysis, and tree-building algorithm employed [7-9]. Therefore, while phenotypic and phylogenetic analyses suggest the existence of supragenus relationships within the *Neocallimastigomycota*, the exact nature of such groupings is yet unclear. Approaches that utilize whole genomic and/or transcriptomic (henceforth referred to as -omics) datasets represent a promising tool towards resolving such relationships [10-14]. Comparative genomics approaches (e.g. calculation of shared Kmer (Kmer overlap) [15, 16],

average nucleotide identity (ANI) [17], identification of genomic syntenic blocks [18]) have

been increasingly utilized in taxonomic studies, aided by the development of lower cost high

67

68

69

70

71

72

73

74

75

76

77

78

79

80

81

82

83

84

85

86

87

88

throughput sequencing technologies and the wider availability of bioinformatic analysis tools. More importantly, the development and implementation of phylogenomic approaches have been crucial in resolving high-rank [13], and intra-clade (e.g. [19]) phylogenies within fungi. Phylogenomic analysis involves the identification of groups of single-copy orthologous genes in the group of interest followed by individually multiple alignments of each orthologous gene. aligning such genes. Analysis to determine a species tree can then be performed on either the concatenated alignment of all genes to obtain a single phylogeny of the group in question, or on the individual alignments via coalescence of individual gene trees. In addition, the inferred gene trees canoutput from such approaches could also be compared to single gene phylogenies to assess their value and potential utility for taxonomic assessment and ecological surveys. Within a Linnaean taxonomic framework, taxonomic associations between genera are accommodated in the intermediate ranks of families, orders, and classes. Currently, AGF genera are recognized in a single family (Neocallimastigaceae), order (Neocallimastigales), and class (Neocallimastigomycetes) in the phylum Neocallimastigomycota [20, 21]. It is interesting to note that a nomenclature novelty entry in Index Fungorum database (IF550425) proposes an additional family (Anaeromycetacea) with the genus Anaeromyces as its sole member, although no detailed justification for such a proposal was provided. Indeed, all current genera in the AGF, including Anaeromyces, are assigned to the family Neocallimastigaceae in recent publications [2, 3], reviews [4, 5.31-34 36], and databases (Mycobank, and Index Fungorum). Regardless, it is clear that the current intermediate rank taxonomic outline of AGF genera has not been proposed based on a detailed comparative phenotypic and phylogenetic analysis of relationships between genera. Rather, it reflects the cumulative and progressive recognition of the phylogenetic and phenotypic distinction of the Neocallimastigomycota when compared to all other fungal clades.

90

91

92

93

94

95

96

97

98

99

100

101

102

103

104

105

106

107

108

109

110

111

The earliest studies on AGF taxonomy [22] proposed accommodating them into a family (Neocallimastigaceae) within the chytrid order Spizellomycetales, a reflection of zoospore ultrastructure similarity; and emended the description of Spizellomycetales order to include zoopsores with multiple flagella. Ten years later, Li et al. [23] used cladistic analysis of 42 morphological and ultrastructural characters to demonstrate the distinction of the AGF when compared to members of the *Chytridiomycetes*, hence elevating the anaerobic gut fungi from a family to an order (Order *Neocallimastigales*). Molecular analysis using concatenated proteincoding genes as well as rRNA genes [21, 24, 25], and several morphological and ultrastructural differences from other *Chytridiomycetes* [26] necessitated their recognition as a phylum (Neocallimastigomycota) with one class (Neocallimastigomycetes), a view that has recently been corroborated via phylogenomic analysis [13]. Indeed, currently published taxonomic outlines, e.g. [20], and databases (e.g. GenBank [27], and Mycocosm [28]) recognize the AGF at the rank of phylum within the Mycota. The last decade has witnessed a rapid expansion in the number of described genera within the *Neocallimastigomycota* [2, 4, 5, 29-34]. Due to such expansion, as well as the continuous recognition of the value of genome-based taxonomy in resolving relationships and circumscribing ranks in fungal taxonomy [10, 13, 14]; we posit that a lineage-wide phylogenomic assessment is warranted to resolve inter-genus relationships and explore the need for intermediate ranks to establish a proper Linnaean-based outline for the phylum. Here, we conducted transcriptomic sequencing on multiple additional AGF genera isolated and characterized in our laboratory, and combined these datasets with previously available AGF transcriptomes and genomes to resolve the inter-genus relationships within the *Neocallimastigomycota*. Based on our results, we propose accommodating AGF described

genera into four distinct families to reflect the observed inter-genus relationships. In addition, we provide quantitative amino acid identity (AAI) for circumscribing such families, and test the utility of multiple single genes/loci as additional markers for resolving AGF phylogeny.

117

118

119

120

121

122

123

124

125

126

127

128

129

130

131

132

133

134

135

136

137

138

Materials and Methods Cultures. Transcriptomes and genomes from fifty-two strains representing fourteen AGF genera were analyzed (Table 1). Of these, transcriptomes of twenty strains, representing six genera for which no prior sequence data were available were sequenced as part of this study. Many of the analyzed strains have previously been described as novel genera or species by the authors [5, 30-32, 34] (Table 1). Others possessed identical features to previously described type strains and were designated as conferre (cf.) strains (Table 1). Few were identified to the genus level and given an alphanumeric strain name designation (Table 1). RNA extraction, Sequencing, quality control, and transcripts assembly. Isolates were grown in rumen fluid medium with cellobiose as the sole carbon source [35] to late log/early stationaryphase (approximately 48 to 60 \(\text{h}\) post inoculation). Cultures were vacuum filtered to obtain fungal biomass then grounded with a pestle under liquid nitrogen. Total RNA was extracted using Epicentre MasterPure yeast RNA purification kit (Epicentre, Madison, WI) according to manufacturer's instructions and stored in RNase-free Tris-EDTA buffer. Transcriptomic sequencing using Illumina HiSeq2500 platform and $2 \square \times \square 150$ bp paired-end library was conducted using the services of a commercial provider (Novogene Corporation, Beijing, China), or at the Oklahoma State University Genomics and Proteomics center. The RNA-seq data were quality trimmed and de novo assembled with Trinity (v2.6.6) using default parameters. For each data set, redundant transcripts were clustered using CD-HIT [36] with identity parameter of 95% (-c 0.95). The obtained nonredundant transcripts were subsequently used for peptide and coding sequence prediction using the TransDecoder (v5.0.2) (https://github.com/TransDecoder/TransDecoder) with a minimum peptide length of 100 amino acids. Assessment of transcriptome completeness per strain was conducted using BUSCO [37]

140

141

142

143

144

145

146

147

148

149

150

151

152

153

154

155

156

157

158

159

160

161

with the fungi odb10 dataset modified to remove 155 mitochondrial protein families as previously suggested [38]. **Phylogenomic analysis.** The phylogenomic analysis includes 20 newly sequenced and 32 existing AGF genomic and transcriptome sequences (Table 1) [38-43]. Five Chytridiomycota genomes were also included as the outgroup (Chytriomyces sp. strain MP 71, Entophlyctis helioformis JEL805, Gaertneriomyces semiglobifer Barr 43, Gonapodya prolifera JEL478, and Rhizoclosmatium globosum JEL800 [44, 45]). The "fungi_odb10" dataset including 758 phylogenomic markers for Kingdom Fungi was retrieved from BUSCO v4.0 package, and used in our analysis. Profile hidden-Markov-models of these markers were created and used to identify homologues in all included fifty-eight fungal proteomes using hmmer3 (v3.1b2) employed in the PHYling pipeline (https://doi.org/10.5281/zenodo.1257002). A total of 670 out of the 758 "fungi_odb10" markers were identified with conserved homologs in the 57 AGF and Chytrids genomes, which were then aligned and concatenated for the subsequent phylogenomic analyses. The final input data include 491,301 sites with 421,690 distinct patterns. The IQ-TREE v1.7 package was used to find the best-fit substitution model and reconstruct the phylogenetic tree with the maximum-likelihood approach. **Average amino acid identity.** We calculated Average Amino acid Identity (AAI) values for all possible pairs in the dataset using the predicted peptides output from TransDecoder.LongOrfs. AAI values were generated using the aai.rb script available as part of the enveomics collection [46]. Through reciprocal all versus all protein Blast, AAI values represent indices of pairwise genomic relatedness [47]. Since its introduction in 2005 [47] as a means for standardizing taxonomy at ranks higher than species, AAI has been extensively used in bacterial and archaeal genome-based taxonomic studies [48-50]. However, AAI has been utilized only sparsely in the

163

164

165

166

167

168

169

170

171

172

173

174

175

176

177

178

179

180

181

182

183

184

fungal world (e.g. [51, 52], with genome-based quantitative comparisons (e.g. Jaccard index of genomic distance (the fraction of shared k-mers), identification of syntenic blocks, and Average Nucleotide Identity (ANI) [15, 18]) being more heavily utilized and often for delineating lower taxonomic level (e.g. species) boundaries. AAI, however, has the advantage of being readily conducted on the predicted peptides from transcriptomic datasets, as it uses amino acid sequences. The ease of obtaining transcriptomic rather than genomic sequences for AGF (mostly due to the extremely high AT content in intergenic regions and the extensive proliferation of microsatellite repeats, often necessitating employing multiple sequencing technologies for successful genomic assembly) makes the use of AAI for delineation of taxonomic boundaries more appealing. Single gene phylogenetic analysis. Two ribosomal loci (D1/D2 LSU, and ITS1) and four protein-coding gene trees (RNA polymerase II large subunit (RPB1), RNA polymerase II second largest subunit (RPB2), Minichromosome maintenance complex component 7 (MCM7), and Elongation factor 1-alpha (EF1 α)) were evaluated. Sequences for ITS1 and D1/D2 LSU were either obtained from prior studies [5, 9, 30-32, 34, 53] or were bioinformatically extracted from genomic assemblies [54]. Amino acids sequences of RPB1, RPB2, MCM7 and EF1α were obtained from the *Anaeromyces robustus* genome (GenBank assembly accession number: GCA 002104895.1), and used as bait for Blastp searches against all predicted proteomes in all transcriptomic datasets. Sequences for each protein, as well as for the rRNA loci were aligned using MAFFT with default parameters. The alignments were used as inputs to IQ-TREEtree [55, 56] first to predict the best substitution model (using the lowest BIC criteria) and to generate maximum likelihood trees under the predicted best model. The "-alrt 1000" option for performing the Shimodaira-Hasegawa approximate-likelihood ratio test (SH-aLRT), "-abayes"

option for performing approximate Bayes test, and the "–bb 1000" option for ultrafast bootstrap (UFB) were added to the IQ-TREE command line, which resulted in the generation of phylogenetic trees with three support values (SH-aLRT, aBayes, and UFB) on each branch.

Nucleotide sequencing accession number. Raw Illumina RNA-seq read sequences are deposited in GenBank under the BioProject accession number PRJNA847081 and BioSample accessions numbers SAMN28920465- SAMN28920484. Individual SRA accessions are provided in Table 1.

194

195

196

197

198

199

200

201

202

203

204

205

206

207

208

209

210

211

212

213

214

215

Results **Sequencing.** Transcriptomic sequencing yielded 15.6 to 23.8 (average 19.82) million reads that were assembled into 22,649 to 106,687 total transcripts, 20,599 to 103,405 distinct transcripts (clustering at 95%; average 40,099), and 13,858 to 28,405 predicted peptides (average 19,667) (Table S2). Assessment of transcriptome completion using BUSCO yielded high values (73.63 to 99.5%) for all assemblies (Table S1). Resolving inter-genus relationships in the Neocallimastigomycota. Multiple supra-genus relationships were well supported in all phylogenomic outputs. Four distinct clades were observed (Figure 1 and Table 2). Clade one constituted members of the genera *Pecoramyces*, Orpinomyces, Neocallimastix, Feramyces and Aestipascuomyces. Within this large clade, a strong support for *Pecoramyces* and *Orpinomyces* association, as well as for *Neocallimastix*, Aestipascuomyces, and Feramyces association was observed (Figure 1). Phenotypically, this clade encompasses all the AGF genera producing polyflagellated zoospores; and all members of the clade, with the exception the genus *Pecoramyces* produce polyflagellated zoospores. Clade two constituted members of the genera Cyllamyces and Caecomyces. Phenotypically, this clade encompasses the two genera exhibiting a bulbous rhizoidal growth pattern in the *Neocallimastigomycota*. Clade three constituted members of the genus *Piromyces*. Compared to all other AGF genera, the genus *Piromyces* currently exhibits high intra-genus sequence divergence based on ITS1 and LSU analysis [3]. The genus was first defined to encompass all phenotypes with monocentric thalli, filamentous rhizoidal system, and monoflagellated zoospores [57]. However, subsequent isolation efforts clearly demonstrated that such phenotype is prevalent in a wide range of phylogenetically disparate genera across the *Neocallimastigomycota* [4, 5, 29]. Currently, *Piromyces* encompasses all taxa phylogenetically

217

218

219

220

221

222

223

224

225

226

227

228

229

230

231

232

233

234

235

236

237

238

affiliated with the first described monocentric, monoflagellated, and filamentous isolate (*Piromyces communis* [57]). Clade four constituted members of the genera *Anaeromyces*, Liebetanzomyces, and Capellomyces. The clade encompasses genera with filamentous rhizoidal system, and monoflagellated zoospores. The genus Anaeromyces produces polycentric thalli, while the genera *Liebetanzomyces*, and *Capellomyces* produce monocentric thalli. Few genera clustered outside these four clades described above. The genera *Paucimyces* and Aklioshbomyces formed distinct branches at the base of clades 1 and 2, respectively (Figure 1). Finally, the position of the genus *Khoyollomyces* was unique and solitary, being consistently located at the base of the tree, suggesting its deep-branching and relatively ancient origin. Estimating AAI identities. AAI values were estimated using the entire dataset of predicted peptides (Figure 2). Intra-genus AAI values ranged between 72.58-99.6% (Average 92.16 ± 8.55). However, the low intra-genus divergence estimates were only confined to the broadly circumscribed genus *Piromyces*. Indeed, excluding *Piromyces* from this analysis, intra-genus AAI values ranged between 87.78-99.6%, (Average 95.67 ± 3.41). Pairwise AAI values for members of different genera within the same clade (intra-clade inter-genus AAI values) ranged between 75.44-85.48% (Average 79.58 ± 2.47). Maximum intra-clade inter-genus divergence was observed between members of the genera Neocallimastix and Pecoramyces (Average 77.5 ± (0.91) and the genera *Neocallimastix* and *Orpinomyces* (Average 77.4 \pm 0.59) in clade 1, while minimal intra-clade inter-genus divergence were observed between Caecomyces and Cyllamyces in clade 2 (83.7% \pm 0.4); as well as the genera Anaeromyces and Capellomyces (Average 84.5 \pm 0.57), the genera Anaeromyces and Liebetanzomyces (Average 83.9 \pm 0.3), and the genera Capellomyces and Liebetanzomyces (Average 85.1 ± 0.18) in clade 4. Inter-clade AAI values averaged 73.15 ± 1.57 , and ranged between 65.27% (between members of the genera *Piromyces*

240

241

242

243

244

245

246

247

248

249

250

251

252

253

254

and Neocallimastix) and 76.64 % (between members of the genera Capellomyces and Pecoramyces). Single gene phylogenetic analysis for resolving AGF inter-genus relationships. We tested whether supra-genus clades topology as well as within clades inter-genus relationships observed in phylogenomic analysis were retained in single gene phylogenies (Figure 3-8). One ribosomal locus (D1/D2 LSU) and one protein-coding gene (RPB1) retained the monophylly of all four clades described above (Figure 3, 5, Table S2). As well, both D1/D2 LSU and RPB1 phylogenies resolved all inter-genus relationships within all clades in the *Neocallimastigomycota* (Figure 3, 5). On the other hand, ITS1, RPB2, MCM7, and EF1α phylogenies each recovered three out of the four supra-genus clades delineated above. The monophylly of clade 1 was not retained in ITS1 and RPB2 phylogenies (Figure 4, 6, Table S2), the monophylly of clade 4 was not retained in MCM7 phylogeny (Figure 7), and the monophylly of clade 3 was not retained in EF1α phylogeny (Figure 8). Further, within the clades that were supported, few inter-genus relationships were compromised in ITS1 (genus Anaeromyces), and EF1 α (genera *Neocallimastix, Orpinomyces,* and *Pecoramyces*) phylogenies.

255 **Discussion** 256 Identifying and circumscribing supra-genus relationships within the 257 **Neocallimastigomycota.** Our phylogenomic analysis identified four distinct statistically 258 supported supra-genus clades in the *Neocallimastigomycota* (Table 2, Figure 1). Clades' 259 boundaries were based on phylogenomic tree topologies, while taking taxonomically informative 260 morphological characteristics into account. For example, phylogenomic analyses placed the 261 genus *Paucimyces* at the base of clade 1, and the genus *Aklioshbomyces* at the base of clade 2. 262 Exclusion of *Paucimyces* from clade 1 was based on its production of monoflagellated zoospores 263 [32], as opposed to the polyflagellated zoospores produced by all members of clade 1 (with the 264 exception of *Pecoramyces*). Similarly, exclusion of *Aklioshbomyces* from clade 2 was based on 265 its filamentous rhizoidal growth pattern; which contrasts the bulbous growth pattern exclusive to 266 both genera (*Caecomyces* and *Cyllamyces*) constituting clade 2. 267 AAI values were further examined to quantitatively circumscribe these clades. A clear 268 delineation of the clade boundary was evident using AAI values (Figure 2). Within genus, AAI 269 values ranged between 87.78-99.6% (or 72.58-99.6% if including values for the broadly 270 circumscribed genus Piromyces). Inter-genus/ Intra-clade AAI estimates ranged between 75.44-271 85.48%, while inter-clade values ranged between 65.27-76.64% (Figure 2). These values are 272 similar to AAI values estimated for delineating the Ascomycetes family Hypoxylaceae [51], but 273 are higher than the arbitrary cutoffs used for delineating taxa in the prokaryotic world (~45-65% 274 for family, ~65-95% for genus [48]). Therefore, we suggest using 85.0%, and 75.0% AAI cutoff 275 values as a guide for circumscribing genera, and families, respectively, in the 276 Neocallimstigomycota. Currently, the genus Piromyces represents the sole genus in clade 3. AAI 277 estimates using the currently available *Piromyces* species –omics datasets suggest broader inter-

279

280

281

282

283

284

285

286

287

288

289

290

291

292

293

294

295

296

297

298

299

300

genus AAI range when compared to other genera (Figure 2). This is a reflection of the fact that the genus was originally circumscribed based on phenotypic, rather than a combination of phenotypic and molecular data. Future availability of additional -omics data coupled to a detailed comparative morphotypic analysis of its described species could possibly lead to splitting this genus (the sole member of clade 3 here) into several clades. Up to this point, only ITS1 and D1/D2 LSU loci have been evaluated for assessment of phylogenetic positions of genera within the Neocallimastigomycota, as well as for ecological culture-independent surveys [7, 9]. To test the utility of other phylomarkers commonly utilized in fungal taxonomy, we assessed additional four protein-coding genes, and examined concordance between each of the six loci (ribosomal ITS1 and D1/D2 LSU, and RPB1, RPB2, MCM7, and EF-1 α) and phylogenomic trees topologies. Our results demonstrate that D1/D2 LSU, currently regarded as the phylomarker of choice for genus-level delineation [9, 58] and utilized as a marker in culture-independent diversity surveys [9], is equally useful in resolving supra-genus clades delineated by phylogenomics (Figures 3, S1). As well, our results add the protein-coding gene RPB1 to the list of phylomarkers that could be used for inter-genus, and supra-clade delineation (Figures 5, S2). As such, values of 8.5%, and 2.1% for LSU, and RPB1, respectively (these values correspond to the 75-percentile value for intra-clade inter-genus divergence based on the distance matrix from the alignments used to generate the maximum likelihood trees in Figures 3, 5) seem to circumscribe these clades. The high sequence similarity in the protein-coding gene RPB1 is quite surprising since, typically, higher levels of divergence are usually observed in protein coding genes when compared to the non-protein-coding ribosomal genes [59]. Other phylomarkers tested here were only successful in resolving three of the four clades, and some also compromised intra- and inter-genus relationships (Figures 4, 6-7).

302

303

304

305

306

307

308

309

310

311

312

313

314

315

316

317

318

319

320

321

322

323

Such failure to resolve genus-level relationships appears to be a function of high sequence similarities in these genes. For example, the inter-genus divergence values between *Orpinomyces* and *Pecoramyces* RPB2 sequences ranged between 0-1.8%, which are comparable to the values within the genus *Orpinomyces*. This has resulted in failure of RPB2 to resolve the *Orpinomyces*-Pecoramyces relationship. The unreliability of the ITS1 locus for clade delineation has been described before, and is mainly due to length variability between genera and high within-strain sequence divergence [7, 9]. Phylogenetic position of taxa currently lacking genome or transcriptome sequences. The fifty-three transcriptomic datasets examined cover fourteen out of the twenty currently described AGF genera. The remaining six genera (Oontomyces, Buwchfawromyces, Agriosomyces, Ghazallomyces, Tahromyces, and Joblinomyces) are all currently represented by a single species. Further, most of these genera appear to exhibit extremely limited geographic and animal host distribution patterns [4, 5, 9, 29]. The phylogenetic position of these six genera could hence be only evaluated using available D1/D2 LSU (and to some extent ITS1) sequence data from taxa description publications. D1/D2 LSU and ITS1 phylogenies strongly support placement of the genus Ghazallomyces as a member of clade 1 (Figure 3, 4) [5]. Further, the genus produces polyflagellated zoospores (an exclusive trait for clade 1), filamentous rhizoid (similar to all taxa in clade 1), and monocentric thalli (similar to all genera in clade 1, except *Orpinomyces*), further supporting its recognition as member of clade 1[5]. Similarly, phylogenetic analysis using D1/D2 LSU and ITS1 supports the placement of genus *Oontomyces* as a member of clade 4 (Figure 3, 4). Members of the genus *Oontomyces* exhibit similar phenotypes (monocentric thalli, monoflagellated zoospores, and filamentous rhizoidal growth patterns) to the genera *Liebetanzomyces* and *Capellomyces* in the clade [29].

325

326

327

328

329

330

331

332

333

334

335

336

337

338

339

340

341

342

343

344

345

346

Interestingly, phylogenetic analysis using the D1/D2 region of LSU rRNA genes places three of the genera for which no –omics data is available (Buwchfawromyces, Tahromyces, and Joblinomyces) in a single distinct monophyletic clade (Figure 4). Future availability of –omics data is needed to confirm such topology. Finally, while the genus Agriosomyces has a distinct position in both ITS1 and LSU phylogenies (Figure 4, ITS), no clear association to any of the clades was apparent. As such, -omics data is hence needed to resolve the position of this genus. Rank assignment for supra-genus clades in the Neocallimastigomycota. Our analysis identifies and circumscribes four distinct clades in the *Neocallimastigomycota*. What taxonomic rank should be assigned to accommodate these clades? The Linnaean classification system places groups of genera into families. A recently proposed definition identifies fungal families as "a compilation of genera with at least one inherent morphological feature that they commonly share or which makes them distinct" [60]. The clades described in this study agree with such a definition, being a compilation of genera forming a distinct and monophyletic lineage with strong statistical support, and most of which share a common distinctive morphological feature (Table 2). We propose retaining all currently described AGF genera in a single order (Neocallimastigales), and a single class (Neocallimastigomycetes) in the phylum Neocallimastigomycota. Such proposition is based on the lack of fundamental differences in their cellular structures, metabolic capabilities, ecological distribution, and life cycle phases in all currently described genera, coupled to the observed AAI values, when compared to the few studies utilizing this approach in fungi [51]. Beyond the four clades described above, we refrain from proposing an additional family for the D1/D2 LSU-defined and well-supported clade encompassing the genera

348

349

350

351

352

353

354

355

356

357

358

359

360

361

362

363

364

365

366

367

368

369

Buwchfawromyces, Tahromyces, and Joblinomyces pending the availability of confirmatory phylogenomic data. As well, we refrain from proposing new families for the genera Khyollomyces, Aklioshbomyces, Paucimyces, and Agriosomyces, due to their current solitary positions in phylogenomic trees (Figure 1), although such proposition would be justified by the isolation of characterization of additional novel taxa and the availability of -omics data from such taxa. Such genera should be regarded as orphan taxa for the present time. The proposed novel families would be named after the first described genus within the clade: Clade 1 = Neocallimastigaceae comprising the genera Neocallimastix (Braune 1913 [61], Vavra and Joyon 1966 [62], Heath et al. 1983, [22]), Ghazallomyces (Hanafy et al. 2021) [5], Orpinomyces (Breton et al. 1989 [63], Barr et al. 1989 [64]), *Pecoramyces* (Hanafy et al. 2017) [30], Feramyces (Hanfay et al. 2018 [31]), and Aestipascuomyces (Stabel et al. 2020, [34]); Clade 2 = Caecomycetaceae fam. nov., comprising the genera Caecomyces (Gold et al. 1988) [57] and Cyllmayces (Ozkose et al. 2001) [33], clade 3 = Piromycetaceae fam. nov., comprising the genus *Piromyces* (Gold et al. 1988) [57]; and clade 4 = *Anaeromycetaceae*, comprising the genera Anaeromyces (Breton et al. 1990) [65], Capellomyces (Hanafy et al. 2021) [5], Liebetanzomyces (Joshi et al. 2018) [66], and *Oontomyces* (Dagar et al. 2015) [29]. Such arrangement would necessitate amending the description of the family *Neocallimastigaceae*, currently encompassing all twenty genera, to include only the six genera stated above, rather than all twenty currently described AGF genera, as well as assigning the genera Anaeromyces (Breton et al. 1990), Capellomyces [5], Liebetanzomyces (Joshi et al. 2018) [66], and Oontomyces (Dagar et al. 2015) to the previously proposed (IF550425) nomenclature novelty family *Anaeromycetaceae*. **Typification** Emended description of fam. Neocallimastigaceae.

371

372

373

374

375

376

377

378

379

380

381

382

383

384

385

386

387

388

389

390

391

392

Obligate anaerobic fungi with monocentric or polycentric thalli and filamentous rhizoidal system. Zoospores are polyflagellated in all described genera, with the exception of the monoflagellated genus *Pecoramyces*. The clade is defined by phylogenomic, phylogenetic and morphological characteristics. Currently accommodates the genera *Neocallimastix* (Braune 1913 [61], Vavra and Joyon 1966 [62], Heath et al. 1983, [22]), Ghazallomyces (Hanafy et al. 2021) [5], Orpinomyces (Breton et al. 1989 [63], Barr et al. 1989 [64]), Pecoramyces (Hanafy et al. 2017) [30], Feramyces (Hanfay et al. 2018 [31]), and Aestipascuomyces (Stabel et al. 2020, [34]). The emended description of the family *Neocallimastigaceae* is generally similar to that provided for the family Neocallimastigaceae [22], and order Neocallimsatigales [23], with amendments to exclude bulbous rhizoidal growth, and to circumscribe its boundaries to encompass a monophyletic clade of six genera. The clade is circumscribed by phylogenomic analysis, AAI values, and confirmed by LSU and RPB1 phylogenetic analyses, as well as morphological characteristics. The emended family encompasses the genera *Neocallimastix* (Braune 1913 [61], Vavra and Joyon 1966 [62], Heath et al. 1983) [22], Orpinomyces (Breton et al. 1989, Barr et al. 1989) [70, 71], *Pecramyces* (Hanafy et al 2017) [32], *Feramyces* (Hanafy et al 2018) [33], Ghazallomyces (Hanafy et al 2020) [5], and Aestipascuomyces (Stabel et al 2020) [8]. Type genus: Neocallimastix Braune 1913 [61], Vavra and Joyon 1966 [62], Heath et al. 1983, [22]. Mycobank ID: MB25486. **Description of** *Caecomycetaceae* **fam. nov.** Obligate anaerobic fungi that produce monoflagellated zoospores, monocentric or polycentric thalli that are either uni- or multisporangiate, and a bulbous rhizoidal system with spherical holdfasts. The clade is

394

395

396

397

398

399

400

401

402

403

404

405

406

407

408

409

410

411

412

circumscribed by phylogenomic analysis, AAI values, and confirmed by LSU and RPB1 phylogenetic analyses, as well as morphological characteristics. Currently accommodates the genera Caecomyces (Gold et al. 1988) [57] and Cyllmayces (Ozkose et al. 2001) [33]. Type genus: Caecomyces (Gold et al 1988) [57]. Mycobank ID: MB844401 **Description of** *Piromycetaceae* **fam. nov.** Obligate anaerobic fungi that produce monoflagellated zoospores, monocentric thalli, and filamentous rhizoidal system. The clade is circumscribed by phylogenomic analysis, AAI values, and confirmed by LSU and RPB1 phylogenetic analyses, as well as morphological characteristics. Currently accommodates the genus *Piromyces* (Gold et al. 1988) [57]. Type genus: Piromyces (Gold et al. 1988) [57]. Mycobank ID: MB844402 Emended description of Anaeromycetaceae fam. nov. Obligate anaerobic fungi that produce monoflagellated zoospores, monocentric or polycentric thalli, and filamentous rhizoidal system. The clade is circumscribed by phylogenomic analysis, AAI values, and confirmed by LSU and RPB1 phylogenetic analyses, as well as morphological characteristics. Currently accommodates the genera Anaeromyces (Breton et al. 1990) [65], Capellomyces (Hanafy et al. 2021) [5], Liebetanzomyces (Joshi et al. 2018) [66], and Oontomyces (Dagar et al. 2015) [29]. Type genus: Anaeromyces, Breton et al. 1990 [65]. Mycobank ID: MB550425.

413 Tables.

414 Table 1. List of strains used in this study.

Genus	species	Strain	Genome BioProject accession number	Transcriptome BioProject accession number	SRA accession number	Assembled transcriptome TSA accession number	Reference
Aestapascuomyces	dupliciliberans	R1		PRJNA847081	SRR19612713		This study
Aklioshbomyces	papillarum	WTS1		PRJNA847081	SRR19612712		This study
Anaeromyces	contortus	ABS23		PRJNA847081	SRR19612701		This study
Anaeromyces	contortus	C3G		PRJNA489922		GGWR00000000.1	[67, 68]
Anaeromyces	contortus	C3J		PRJNA489922		GGWO00000000.1	[67, 68]
Anaeromyces	contortus	G3G		PRJNA489922		GGWP00000000.1	[67, 68]
Anaeromyces	contortus	Na		PRJNA489922		GGWN00000000.1	[67, 68]
Anaeromyces	contortus	O2		PRJNA489922		GGWQ00000000.1	[67, 68]
Anaeromyces	mucronatus	YE505		PRJNA437872			[38]
Anaeromyces	robustus	S4	PRJNA330692	PRJNA250973			[69]
Caecomyces	communis	churrovis	PRJNA347164	PRJNA393353			[39, 41]
Caecomyces	communis	FD27		PRJNA847081	SRR19612700		This study
Caecomyces	communis	TB33		PRJNA847081	SRR19612699		This study
Caecomyces	communis	Iso3		PRJNA489922		GGXE00000000.1	[67, 68]
Caecomyces	communis	Brit4		PRJNA489922		GGWS00000000.1	[67, 68]
Capellomyces	forminis	Cap2a		PRJNA847081	SRR19612698		This study
Cyllamyces	aberensis	TSB2		PRJNA847081	SRR19612697		This study
Feramyces	austinii	WSF2		PRJNA489922		GGWT00000000.1	[67, 68]
Feramyces	austinii	WSF3		PRJNA489922		GGWU00000000.1	[67, 68]
Khyollomyces	ramosus	ZO44		PRJNA847081	SRR19612696		This study
Liebetanzomyces	polymoprphus	Orc37		PRJNA847081	SRR19612695		This study
Neocallimastix	frontalis	EC30		PRJNA847081	SRR19612694		This study

Neocallimastix	frontalis	Hef5		PRJNA489922		GGXJ00000000.1	[67, 68]
Neocallimastix	frontalis	27		PRJNA437872			[38]
Neocallimastix	cameroonii	G1	PRJNA262392	PRJNA251043			[69]
Neocallimastix	cameroonii	lanati	PRJNA658393	PRJNA677809			[43]
Neocallimastix	cameroonii	G3		PRJNA489922		GGXC00000000.1	[67, 68]
Orpinomyces	joyonii	AB6		PRJNA847081	SRR19612711		This study
Orpinomyces	joyonii	AB3		PRJNA847081	SRR19612710		This study
Orpinomyces	joyonii	ABC-24		PRJNA847081	SRR19612709		This study
Orpinomyces	joyonii	D3A		PRJNA489922		GGWV00000000.1	[67, 68]
Orpinomyces	joyonii	D3B		PRJNA489922		GGWW00000000.1	[67, 68]
Orpinomyces	joyonii	D4C		PRJNA489922		GGWX00000000.1	[67, 68]
Orpinomyces	joyonii	SG4		PRJNA437872			[38]
Paucimyces	polynucleatus	BB3		PRJNA847081	SRR19612708		This study
Pecoramyces	ruminantium	C1A	PRJNA200719	PRJNA284193			[67, 70]
Pecoramyces	ruminantium	S4B		PRJNA489922		GGWY00000000.1	[67, 68]
Pecoramyces	ruminantium	FS3C		PRJNA489922		GGXF00000000.1	[67, 68]
Pecoramyces	ruminantium	FX4B		PRJNA489922		GGWZ00000000.1	[67, 68]
Pecoramyces	ruminantium	YC3		PRJNA489922		GGXA00000000.1	[67, 68]
Pecoramyces	ruminantium	Orc32		PRJNA847081	SRR19612707		This study
Pecoramyces	ruminantium	AS31		PRJNA847081	SRR19612706		This study
Pecoramyces	ruminantium	AS32		PRJNA847081	SRR19612705		This study
Pecoramyces	ruminantium	F1	PRJNA517297	PRJNA517315			[71]
Piromyces	finnis	finn	PRJNA330696	PRJNA268530			[69]
Piromyces	finnis	DonB11		PRJNA847081	SRR19612704		This study
Piromyces	cryptodigmaticus	Axs23		PRJNA847081	SRR19612703		This study
Piromyces	cryptodigmaticus	A1		PRJNA489922		GGXB00000000.1	[67, 68]
Piromyces	potentiae	B4		PRJNA489922		GGXH00000000.1	[67, 68]
Piromyces	sp. NZB19	Ors32		PRJNA847081	SRR19612702		This study

Piromyces	sp. PR1	E2	PRJNA82799		[69]
Piromyces	rhizinflatus	YM600		PRJNA437872	[38]

420 Table 2. Clades circumscribed in this study.

Clades					
	Genera	Average intra-genus (range) Average inter-genus (range) Average intra-clade (range) (range)		Average inter-clade (range)	Phenotype
Clade 1	Pecoramyces, Orpinomyces, Neocallimastix, Aestipascuomyces, Feramyces	96.89 (87.78-99.49)	82.95 (75.44-78.99)	73.21 (65.27-76.64)	Polyflagellated zoospores except for <i>Pecoramyces</i>
Clade 2	Cyllamyces, Caecomyces	94.01 (88.02-98.37)	84.05 (83.08-83.67)	72.8 (67.39-74.91)	Bulbous rhizoidal growth pattern
Clade 3	Piromyces	79.35 (72.58-99.06)	79.35 (72.58-99.06)	72.61 (65.27-75.61)	Monocentric thalli, monoflagellated zoospores, filamentous rhizoidal growth pattern
Clade 4	Anaeromyces, Liebetanzomyces, Capellomyces	96.55 (93.07-99.6)	84.41 (83.58-85.48)	73.75 (67.25-76.64)	Filamentous rhizoidal growth pattern, monoflagellated zoospore, all monocentric thallus except <i>Anaeromyces</i>
	Aklioshbomyces	NA	NA	73.54 (69.1-75.42)	
	Paucimyces	NA	NA	74.26 (68.14-76.98)	
	Khyollomyces	NA	NA	71.88 (66.47-73.41)	

422

423

424

425

426

427

428

429

430

431

432

433

434

435

436

437

438

439

440

441

442

443

Figure legends. **Figure 1.** Phylogenomic tree of *Neocallimastigomycota* based on 670 genome-wide markers highlighting the family-level relationships within the phylum. The tree was reconstructed using the maximum likelihood approach implemented in the IQ-TREE package. Number on each branch represents the ultrafast bootstrap value suggesting the robustness of the taxa joining. The scale bar at the bottom indicates the number of substitutions per site in the analysis. Isolate names at tree tips are color coded by clade (clade 1, purple; clade 2, layender; clade 3, orange; clade 4, light blue). Figure 2. Upper triangle matrix (A) and box and whisker plots (B) for the AAI values obtained for all possible pairwise comparisons of the datasets analyzed in this study. (A) Isolate names in rows and columns are color coded by clade (clade 1, purple; clade 2, lavender; clade 3, orange; clade 4, light blue). The AAI values for each clade are shown within a thick border. Intra-genus values are shown in red text with pink highlight, intra-clade/ inter-genus values are shown in blue text with light blue highlight, while inter-clade values are shown in green text with light green highlight. Values for the three genera unaffiliated with the 4 clades are highlighted in grey. (B) Box and whisker plots constructed using the values in (A). Intra-genus values (red) are shown both including and excluding the genus *Piromyces*. Intra-clade/ inter-genus values are shown in blue. Inter-clade values are shown in green. Each box plot spans the region between the 25-percentile to 75-percentile, while the whiskers limit the minimum and maximum scores excluding the outliers. The thick line inside the box marks the median, while the 'x' corresponds to the average value. Figure 3. Maximum likelihood phylogenetic tree constructed using the D1/D2 region of the LSU rRNA genes of all cultured and described *Neocallimastigomycota* genera. Sequences were either

445

446

447

448

449

450

451

452

453

454

455

456

457

458

459

460

461

462

463

464

465

obtained from prior studies [5, 9, 30-32, 34, 53] or were bioinformatically extracted from genomic assemblies [54], and GenBank accession numbers are shown for each branch label. Sequences were aligned using MAFFT with default parameters. IQ-tree [55, 56] was used to choose the best substitution model (TN+F+G4 was chosen using the lowest BIC criteria) and to generate the maximum likelihood tree. Support values at each node correspond to SH-aLRT, aBayes, and ultrafast bootstrap. Clades are coded using the same color code in Figure 2 (clade 1, purple; clade 2, lavender; clade 3, orange; clade 4, light blue), and boxes with the same colors are used to delimit each clade. The support values at the nodes corresponding to each clade are shown in bold red text, and the node itself is shown as a red dot. The tree was rooted (root not shown) using the D1/D2 region of the LSU rRNA gene from *Chytriomyces* sp. WB235A (GenBank accession number DQ536493.1). **Figure 4.** Maximum likelihood phylogenetic tree constructed using the ITS1 region of all cultured and described Neocallimastigomycota genera. Sequences were either obtained from prior studies [5, 9, 30-32, 34, 53] or were bioinformatically extracted from genomic assemblies [54], and GenBank accession numbers are shown for each branch label. Sequences were aligned using MAFFT with default parameters. IQ-tree [55, 56] was used to choose the best substitution model (TN+F+G4 was chosen using the lowest BIC criteria) and to generate the maximum likelihood tree. Support values at each node correspond to SH-aLRT, aBayes, and ultrafast bootstrap. Branch labels are color coded using the same color code in Figure 2 (clade 1, purple; clade 2, lavender; clade 3, orange; clade 4, light blue), and boxes with the same colors are used to delimit each clade. The support values at the nodes corresponding to each clade are shown in bold red text, and the node itself is shown as a red dot. The tree was rooted (root not shown)

467

468

469

470

471

472

473

474

475

476

477

478

479

480

481

482

483

484

485

486

487

488

using the ITS1 region from *Chytriomyces* sp. JEL176 (GenBank accession number AY349118.1). Figure 5. Maximum likelihood phylogenetic tree constructed using the protein sequences of the largest subunit of DNA-dependent RNA polymerase II (RPB1). Amino acids sequence of RPB1 was obtained from the *Anaeromyces robustus* genome (GenBank assembly accession number: GCA_002104895.1), and used as bait for Blastp searches against all predicted proteomes in all transcriptomic datasets. Sequences were aligned using MAFFT with default parameters. IQ-tree [55, 56] was used to choose the best substitution model (LG+R2 was chosen using the lowest BIC criteria) and to generate the maximum likelihood tree. Support values at each node correspond to SH-aLRT, aBayes, and ultrafast bootstrap. Branch labels are color coded using the same color code in Figure 2 (clade 1, purple; clade 2, lavender; clade 3, orange; clade 4, light blue), and boxes with the same colors are used to delimit each clade. The support values at the nodes corresponding to each clade are shown in bold red text, and the node itself is shown as a red dot. The tree was rooted (root not shown) using the RPB1 sequence from *Batrachochytrium* dendrobatidis JAM81 (GenBank accession number EGF82086.1). **Figure 6.** Maximum likelihood phylogenetic tree constructed using the protein sequences of the second largest subunit of DNA-dependent RNA polymerase II (RPB2). Amino acids sequence of RPB2 was obtained from the *Anaeromyces robustus* genome (GenBank assembly accession number: GCA 002104895.1), and used as bait for Blastp searches against all predicted proteomes in all transcriptomic datasets. Sequences were aligned using MAFFT with default parameters. IQ-tree [55, 56] was used to choose the best substitution model (LG+R3 was chosen using the lowest BIC criteria) and to generate the maximum likelihood tree. Support values at each node correspond to SH-aLRT, aBayes, and ultrafast bootstrap. Branch labels are color

490

491

492

493

494

495

496

497

498

499

500

501

502

503

504

505

506

507

508

509

510

511

coded using the same color code in Figure 2 (clade 1, purple; clade 2, lavender; clade 3, orange; clade 4, light blue), and boxes with the same colors are used to delimit each clade if the clade is supported. The support values at the nodes corresponding to each clade are shown in bold red text, and the node itself is shown as a red dot. The tree was rooted (root not shown) using the RPB2 sequence from Batrachochytrium dendrobatidis JELA23 (GenBank accession number OAJ42635.1). Figure 7. Maximum likelihood phylogenetic tree constructed using the protein sequences of the DNA replication licensing factor MCM7. Amino acids sequence of MCM7 was obtained from the Anaeromyces robustus genome (GenBank assembly accession number: GCA 002104895.1), and used as bait for Blastp searches against all predicted proteomes in all transcriptomic datasets. Sequences were aligned using MAFFT with default parameters. IQ-tree [55, 56] was used to choose the best substitution model (LG+R3 was chosen using the lowest BIC criteria) and to generate the maximum likelihood tree. Support values at each node correspond to SH-aLRT, aBayes, and ultrafast bootstrap. Branch labels are color coded using the same color code in Figure 2 (clade 1, purple; clade 2, lavender; clade 3, orange; clade 4, light blue), and boxes with the same colors are used to delimit each clade if the clade is supported. The support values at the nodes corresponding to each clade are shown in bold red text, and the node itself is shown as a red dot. The tree was rooted (root not shown) using the MCM7 sequence from *Batrachochytrium* dendrobatidis JAM81 (GenBank accession number XP 006677581.1). **Figure 8.** Maximum likelihood phylogenetic tree constructed using the protein sequences of the elongation factor 1-alpha (EF-1A). Amino acids sequence of EF-1A was obtained from the Anaeromyces robustus genome (GenBank assembly accession number: GCA_002104895.1), and used as bait for Blastp searches against all predicted proteomes in all transcriptomic datasets.

Sequences were aligned using MAFFT with default parameters. IQ-tree [55, 56] was used to choose the best substitution model (LG+R2 was chosen using the lowest BIC criteria) and to generate the maximum likelihood tree. Support values at each node correspond to SH-aLRT, aBayes, and ultrafast bootstrap. Branch labels are color coded using the same color code in Figure 2 (clade 1, purple; clade 2, lavender; clade 3, orange; clade 4, light blue), and boxes with the same colors are used to delimit each clade if the clade is supported. The support values at the nodes corresponding to each clade are shown in bold red text, and the node itself is shown as a red dot. The tree was rooted (root not shown) using the EF-1A sequence from *Batrachochytrium dendrobatidis* JEL423 (GenBank accession number OAJ38128.1).

References

522

523

- 1. Gruninger RJ, Puniya AK, Callaghan TM, Edwards JE, Youssef N et al. Anaerobic
- 524 fungi (phylum Neocallimastigomycota): advances in understanding their taxonomy, life cycle,
- ecology, role and biotechnological potential. FEMS Microbiol Ecol 2014;90:1-17.
- 526 2. Hanafy RA, Dagar SS, Griffith GW, Youssef NH, Elshahed. MS. Taxonomy of the
- 527 anaerobic gut fungi (Neocallimastigomycota): a review of classification criteria and description
- of current taxa. *Int J Syst Evol Microbiol* 2022; 72:005322.
- 3. Elshahed M, Hanafy R, Cheng Y, Dagar SS, Edwards JE et al. On the characterization
- and rank assignment criteria for the anaerobic gut fungi (Neocallimastigomycota). Int J Syst Evol
- 531 *Microbiol* 2022; *In Press.* DOI 10.1099/ijsem.0.005449
- 532 4. Callaghan TM, Podmirseg SM, Hohlweck D, Edwards JE, Puniya AK et al.
- 533 Buwchfawromyces eastonii gen. nov., sp. nov.: a new anaerobic fungus (Neocallimastigomycota)
- isolated from buffalo faeces. *MycoKeys* 2015;9:11-28.
- 535 5. Hanafy RA, Lanjekar VB, Dhakephalkar PK, Callaghan TM, Dagar SS et al. Seven
- new Neocallimastigomycota genera from wild, zoo-housed, and domesticated herbivores greatly
- expand the taxonomic diversity of the phylum. . *Mycologia* 2020;112::1212-1239.
- 6. Wang X, Liu X, Groenewald JZ. Phylogeny of anaerobic fungi (phylum
- Neocallimastigomycota), with contributions from yak in China. Antonie Van Leeuwenhoek
- 540 2017;110(1):87-103.
- 7. Edwards JE, Hermes GDA, Kittelmann S, Nijsse B, Smidt H. Assessment of the accuracy
- of high-throughput sequencing of the ITS1 region of Neocallimastigomycota for community
- composition analysis. Front Micorobiol 2019;10:2370.

- 8. Stabel M, Schweitzer T, Haack K, Gorenflo P, Aliyu H et al. Isolation and biochemical
- 545 characterization of six anaerobic fungal strains from zoo animal feces *Microorganisms*
- 546 2021;9:1655.
- 9. Hanafy RA, Johnson B, Youssef NH, Elshahed MS. Assessing anaerobic gut fungal
- 548 diversity in herbivores using D1/D2 large ribosomal subunit sequencing and multi-year isolation.
- 549 Environ Microbiol 2020;22:3883-3908.
- 550 10. **Choi J, Kim S-H**. A genome tree of life for the Fungi kingdom. *Proc Natl Acad Sci*
- 551 2017;114:9391-9396.
- 552 11. Montoliu-Nerin M, Sánchez-García M, Bergin C, Kutschera VE, Johannesson H et al.
- In-depth phylogenomic analysis of arbuscular mycorrhizal fungi based on a comprehensive set of
- de novo genome assemblies. Front Fungal Biol, 2021; 2:716385.
- 555 12. Galindo LJ, López-García P, Torruella G, Karpov S, Moreira D. Phylogenomics of a
- new fungal phylum reveals multiple waves of reductive evolution across Holomycota. *Nature*
- 557 *Communications* 2021;12(1):4973.
- 13. Li Y, Steenwyk JL, Chang Y, Wang Y, James TY et al. A genome-scale phylogeny of the
- 559 kingdom Fungi. *Current Biology* 2021;31:1653-1665.
- 14. James TY, Stajich JE, Hittinger CT, Rokas A. Toward a Fully Resolved Fungal Tree of
- 561 Life. *Annual Rev Microbiol* 2020;74:291-313.
- 562 15. Gostinčar C. Towards Genomic Criteria for Delineating Fungal Species. *J Fungi (Basel)*
- 563 2020;6:246.
- 16. Pornputtapong N, Acheampong DA, Patumcharoenpol P, Jenjaroenpun P,
- Wongsurawat T et al. KITSUNE: A tool for identifying empirically optimal K-mer length for
- alignment-free phylogenomic analysis. Front Bioengineer Biotechnol 2020;8:556413-556413.

- 17. Gramaje D, Berlanas C, Martínez-Diz MDP, Diaz-Losada E, Antonielli L et al.
- 568 Comparative genomic analysis of *Dactylonectria torresensis* strains from grapevine, soil and
- weed highlights potential mechanisms in pathogenicity and endophytic lifestyle. *J Fungi (Basel)*
- 570 2020;6:255.
- 18. Li C, Guo Z, Zhou S, Han Q, Zhang M et al. Evolutionary and genomic comparisons of
- 572 hybrid uninucleate and nonhybrid Rhizoctonia fungi. *Communications Biol* 2021;4:201.
- 573 19. Steenwyk JL, Shen XX, Lind AL, Goldman GH, Rokas A. A robust phylogenomic time
- 574 tree for biotechnologically and medically important Fungi in the genera Aspergillus and
- 575 *Penicillium. mBio* 2019;10(4).
- 576 20. Adl SM, Bass D, Lane CE, Lukeš J, Schoch CL et al. Revisions to the classification,
- 577 nomenclature, and diversity of eukaryotes. *J Eukaryotic Microbiol* 2019;66:4-119.
- 578 21. **Hibbett DS, et al.** A higher-level phylogenetic classification of the Fungi. *Mycol Res* 2007;
- 579 111:509-547.
- 580 22. **Heath BI, Bauchop T, Skipp RA**. Assignment of the rumen anaerobe *Neocallimastix*
- 581 frontalis to the Spizellomycetales (Chytridiomycetes) on the basis of its polyflagellate zoospore
- 582 ultrastructure. *Can J Bot* 1983;61:295-307.
- 583 23. Li J, Heath IB, Packer L. The phylogenetic relationships of the anaerobic
- 584 chytridiomycetous gut fungi (Neocallimasticaceae) and the Chytridiomycota. II. Cladistic
- analysis of structural data and description of Neocallimasticales ord.nov. Canadian J Bot
- 586 1993;71:393-407.
- 587 24. James TY, Kauff F, Schoch CL, Matheny PB, Hofstetter V et al. Reconstructing the early
- evolution of Fungi using a six-gene phylogeny. *Nature* 2006;443:818-822.

- 589 25. **Tedersoo L, Sánchez-Ramírez S, Kõljalg U, Bahram M, Döring M et al.** High-level
- 590 classification of the Fungi and a tool for evolutionary ecological analyses. Fungal Diversity
- 591 2018;90:135-159.
- 592 26. **Powell MJ, Letcher PM**. 6 Chytridiomycota0 Chytridiomycota, Monoblepharidomycota,0
- 593 Monoblepharidomycotaand Neocallimastigomycota0 Neocallimastigomycota. In: McLaughlin
- 594 DJ, Spatafora JW (editors). Systematics and Evolution: Part A. Berlin, Heidelberg: Springer
- 595 Berlin Heidelberg; 2014. pp. 141-175.
- 596 27. Benson DA, Cavanaugh M, Clark K, Karsch-Mizrachi I, Ostell J et al. GenBank.
- 597 Nucleic Acids Res 2018;46(D1):D41-d47.
- 598 28. Grigoriev IV, Nikitin R, Haridas S, Kuo A, Ohm R et al. MycoCosm portal: gearing up
- for 1000 fungal genomes. *Nucleic Acids Res* 2014;42(Database issue):D699-704.
- 29. Dagar SS, Kumar S, Griffith GW, Edwards JE, Callaghan TM et al. A new anaerobic
- fungus (*Oontomyces anksri* gen. nov., sp. nov.) from the digestive tract of the Indian camel
- 602 (Camelus dromedarius). Fungal Biol 2015;19:731-737.
- 30. Hanafy RA, Elshahed MS, Liggenstoffer AS, Griffith GW, Youssef NH. Pecoramyces
- 604 ruminantium, gen. nov., sp. nov., an anaerobic gut fungus from the feces of cattle and sheep.
- 605 *Mycologia* 2017;109:231-243.
- 31. Hanafy RA, Elshahed MS, Youssef NH. Feramyces austinii, gen. nov., sp. nov., an
- anaerobic gut fungus from rumen and fecal samples of wild Barbary sheep and fallow deer
- 608 *Mycologia* 2018; 110:513-525
- 609 32. Hanafy RA, Yousseef NH, Elshahed MS. Paucimyces polynucleatus gen. nov., a
- 610 novel polycentric genus of anaerobic gut fungi from the feces of a wild blackbuck antelope. Int J
- 611 Syst Evol Microbiol 2021;71:004832.

613

614

615

616

617

618

619

620

621

622

623

624

625

626

627

628

629

630

631

632

633

634

33. Ozkose E, Thomas BJ, Davies DR, Griffith GW, Theodorou MK. Cyllamyces aberensis gen.nov. sp.nov., a new anaerobic gut fungus with branched sporangiophores isolated from cattle. Can J Bot 2001;79:666-673. 34. Stabel M, Hanafy RA, Schweitzer T, Greif M, Aliyu H et al. Aestipascuomyces dupliciliberans gen. nov, sp. nov., the first cultured representative of the uncultured SK4 clade from Aoudad Sheep and Alpaca. Microorganisms 2020;8:1734. 35. Calkins S, Elledge NC, Hanafy RA, Elshahed MS, Youssef N. A fast and reliable procedure for spore collection from anaerobic fungi: Application for RNA uptake and long-term storage of isolates. J Microbiol Meth 2016;127:206-213. 36. Fu L, Niu B, Zhu Z, Wu S, Li W. CD-HIT: accelerated for clustering the next-generation sequencing data. *Bioinformatics* 2012;28:3150-3152. 37. Manni M, Berkeley MR, Seppey M, Simão FA, Zdobnov EM. BUSCO update: novel and streamlined workflows along with broader and deeper phylogenetic coverage for scoring of eukaryotic, prokaryotic, and viral genomes. *Mol Biol Evol* 2021;38:4647-4654. 38. Gruninger RJ, Nguyen TTM, Reid ID, Yanke JL, Wang P et al. Application of transcriptomics to compare the carbohydrate active enzymes that are expressed by diverse genera of anaerobic fungi to degrade plant cell wall carbohydrates. Front Microbiol 2018;9:1581. 39. Brown JL, Swift CL, Mondo SJ, Seppala S, Salamov A et al. Co cultivation of the anaerobic fungus Caecomyces churrovis with Methanobacterium bryantii enhances transcription of carbohydrate binding modules, dockerins, and pyruvate formate lyases on specific substrates. *Biotechnol Biofuels* 2021;14(1):234. 40. Haitjema CH, Gilmore SP, Henske JK, Solomon KV, de Groot R et al. A parts list for fungal cellulosomes revealed by comparative genomics. Nat Microbiol 2017;2:17087.

- 41. Henske JK, Gilmore SP, Knop D, Cunningham FJ, Sexton JA et al. Transcriptomic
- characterization of *Caecomyces churrovis*: a novel, non-rhizoid-forming lignocellulolytic
- anaerobic fungus. *Biotechnol Biofuels* 2017;10:305.
- 42. Li Y, Li Y, Jin W, Sharpton TJ, Mackie RI et al. Combined genomic, transcriptomic,
- proteomic, and physiological characterization of the growth of *Pecoramyces* sp. F1 in
- monoculture and co-culture with a syntrophic methanogen. Front Microbiol 2019;10:435.
- 43. Wilken SE, Monk JM, Leggieri PA, Lawson CE, Lankiewicz TS et al. Experimentally
- validated reconstruction and analysis of a genome-scale metabolic model of an anaerobic
- Neocallimastigomycota fungus. mSystems 2021;6(1).
- 644 44. Chang Y, Wang S, Sekimoto S, Aerts AL, Choi C et al. Phylogenomic Analyses Indicate
- 645 that Early Fungi Evolved Digesting Cell Walls of Algal Ancestors of Land Plants. *Genome Biol*
- 646 Evol 2015;7:1590-1601.
- 45. Mondo SJ, Dannebaum RO, Kuo RC, Louie KB, Bewick AJ et al. Widespread adenine
- N6-methylation of active genes in fungi. *Nat Genet* 2017;49:964-968.
- 46. **Rodriguez-R L, Konstantinidis K**. The enveomics collection: a toolbox for specialized
- analyses of microbial genomes and metagenomes. PeerJ Preprints; 2016.
- 47. **Konstantinidis KT, Tiedje JM**. Towards a genome-based taxonomy for prokaryotes. J
- 652 Bacteriol 2005;187:6258-6264.
- 48. Konstantinidis KT, Rosselló-Móra R, Amann R. Uncultivated microbes in need of their
- 654 own taxonomy. *The ISME Journal* 2017;11(11):2399-2406.
- 49. Parks DH, Rinke C, Chuvochina M, Chaumeil P-A, Woodcroft BJ et al. Recovery of
- 656 nearly 8,000 metagenome-assembled genomes substantially expands the tree of life. *Nature*
- 657 *Microbiol* 2017;2:1533-1542.

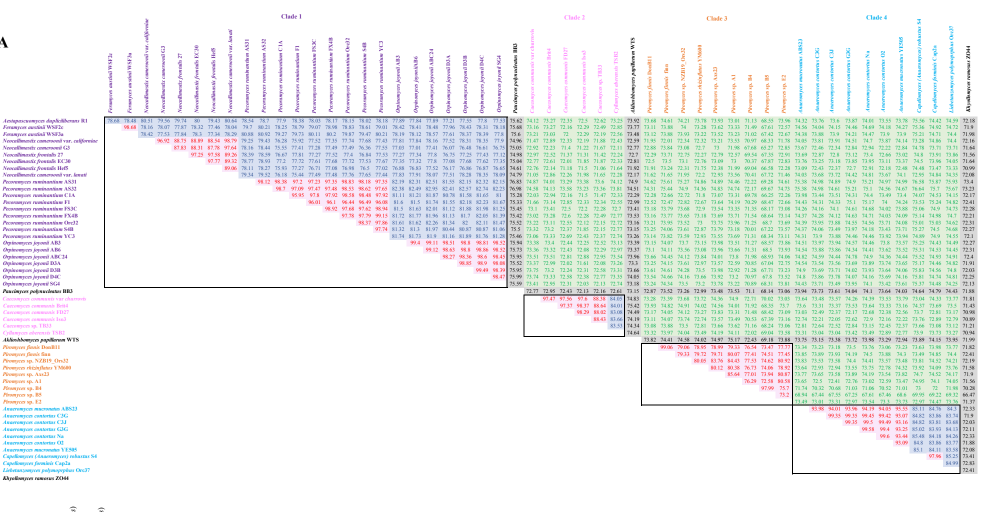
- 658 50. Parks DH, Chuvochina M, Waite DW, Rinke C, Skarshewski A et al. A standardized
- bacterial taxonomy based on genome phylogeny substantially revises the tree of life. *Nat*
- 660 Biotechnol 2018;36:996-1004.
- 51. Wibberg D, Stadler M, Lambert C, Bunk B, Spröer C et al. High quality genome
- sequences of thirteen Hypoxylaceae (Ascomycota) strengthen the phylogenetic family backbone
- and enable the discovery of new taxa. Fungal Diversity 2021;106:7-28.
- 52. Wang K, Sipilä T, Overmyer K. A novel *Arabidopsis* phyllosphere resident *Protomyces*
- species and a re-examination of genus Protomyces based on genome sequence data. IMA Fungus
- 666 2021;12:8.
- 53. Hanafy RA, Johnson B, Elshahed MS, Youssef NH. Anaeromyces contortus, sp. nov., a
- new anaerobic gut fungal species (Neocallimastigomycota) isolated from the feces of cow and
- 669 goat. Mycologia 2018;110:502-512.
- 54. Solomon KV, Haitjema CH, Henske JK, Gilmore SP, Borges-Rivera D et al. Early-
- branching gut fungi possess a large, comprehensive array of biomass-degrading enzymes.
- 672 Science 2016;351:1192-1195.
- 55. **Hoang DT, Chernomor O, von Haeseler A, Minh BQ, Vinh LS**. UFBoot2: Improving the
- Ultrafast Bootstrap Approximation. *Mol Biol Evol* 2018;35(2):518-522.
- 56. Nguyen LT, Schmidt HA, von Haeseler A, Minh BQ. IQ-TREE: a fast and effective
- stochastic algorithm for estimating maximum-likelihood phylogenies. *Mol Biol Evol*
- 677 2015;32:268-274.
- 57. **Gold JJ, Heath IB, Bauchop T**. Ultrastructural description of a new chytrid genus of
- caecum anaerobe, *Caecomyces equi* gen. nov., sp. nov., assigned to the Neocallimasticaceae
- 680 *BioSystems* 1988;21:403–415.

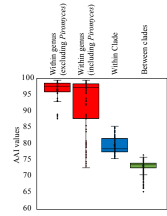
- 58. Dagar SS, Kumar S, Mudgil P, Singh R, Puniya AK. D1/D2 Domain of large-subunit
- ribosomal DNA for differentiation of Orpinomyces spp. Appl Environ Microbiol 2011;77:6722-
- 683 6725.
- 59. Chethana KWT, Jayawardena RS, Hyde KD. Hurdles in fungal taxonomy: Effectiveness
- of recent methods in discriminating taxa. *Megataxa* 2020;1(2):114-122.
- 686 60. **Jeewon R**. Establishing species boundaries and new taxa among fungi: recommendations to
- resolve taxonomic ambiguities. *Mycosphere* 2016;7:1669-1677.
- 688 61. **Braune R**. Untersuchungen uber die im Wiederkauermagen vorkommenden Protozoen
- 689 (Investigations into the protozoa occurring in the ruminant stomach). Archiv für Protistenkunde
- 690 1913;32:I I 1-170.
- 691 62. **Vavra J, Joyon L**. Etude sur la morphologie, le cycle ivolutif et la position systematique de
- 692 Callimastix cyclopsis Weissenberg 1912. (Study on the morphology, the evolutionary cycle and
- the systematic position of Callimastix cyclopsis Weissenberg 1912). *Protistologica* 1966;2:5-15.
- 694 63. Breton A, Bernalier A, Bonnemoy F, Fonty G, Gaillard B et al. Morphological and
- 695 metabolic characterization of a new species of strictly anaerobic rumen fungus: *Neocallimastix*
- 696 joyonii. FEMS Microbiol Lett 1989;58:309-314.
- 697 64. **Barr DJ, KudO H, Jakober KD, Cheng KJ**. Morphology and development of rumen fungi:
- 698 Neocallimastix sp., Piromyces communis, and Orpinomyces bovis gen.nov., sp.nov. Can J Bot
- 699 1989; 67 2815–2824
- 700 65. Breton A, Bernalier A, Dusser M, Fonty G, Gaillard-Martinie B et al. Anaeromyces
- 701 mucronatus nov. gen., nov. sp. A new strictly anaerobic rumen fungus with polycentric thallus.
- 702 FEMS Microbiol Lett 1990;70:177-182.

703 66. Joshi A, Lanjekar VB, Dhakephalkar PK, Callaghan TM, Griffith GW et al. 704 Liebetanzomyces polymorphus gen. et sp. nov., a new anaerobic fungus (Neocallimastigomycota) 705 isolated from the rumen of a goat. MycoKeys 2018;40:89-110. 706 67. Murphy CL, Youssef NH, Hanafy RA, Couger MB, Stajich J et al. Horizontal gene 707 transfer forged the evolution of anaerobic gut fungi into a phylogenetically distinct gut-dwelling 708 fungal lineage. Appl Environ Microbiol 2019;85::e00988-00919. 709 68. Wang Y, Youssef N, Couger M, Hanafy R, Elshahed M et al. Comparative genomics and 710 divergence time estimation of the anaerobic fungi in herbivorous mammals. msystems 711 2019;00247-19. 712 69. Haitjema CH, Gilmore SP, Henske JK, Solomon KV, Groot Rd et al. A parts list for 713 fungal cellulosomes revealed by comparative genomics. Nature Microbiol 2017;2:17087. 714 70. Youssef NH, Couger MB, Struchtemeyer CG, Liggenstoffer AS, Prade RA et al. The 715 genome of the anaerobic fungus *Orpinomyces* sp. strain C1A reveals the unique evolutionary 716 history of a remarkable plant biomass degrader. Appl Environ Microbiol 2013;79:4620-4634... 717 71. Li Y, Li Y, Jin W, Sharpton TJ, Mackie RI et al. Combined genomic, transcriptomic, 718 proteomic, and physiological characterization of the growth of *Pecoramyces* sp. F1 in 719 monoculture and co-culture with a syntrophic methanogen Front Micorobiol 2019;10:435.

720

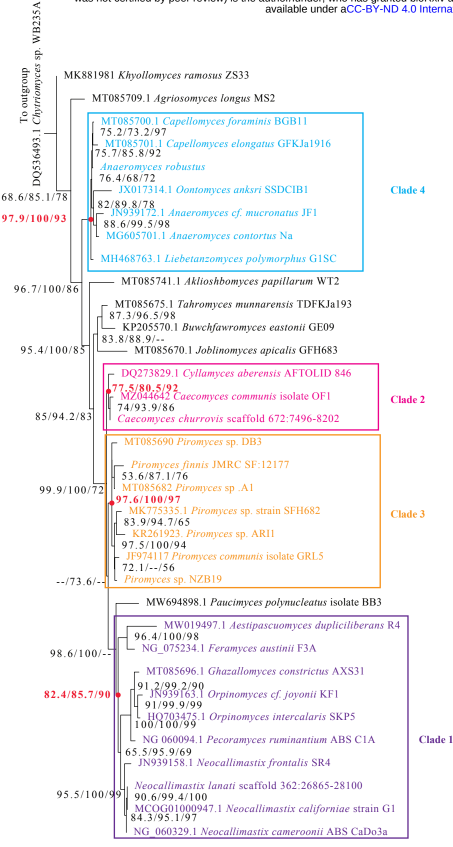




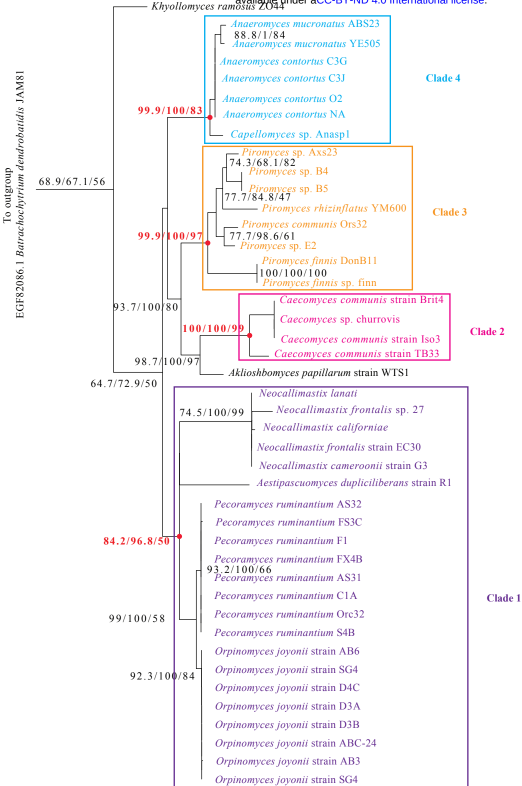


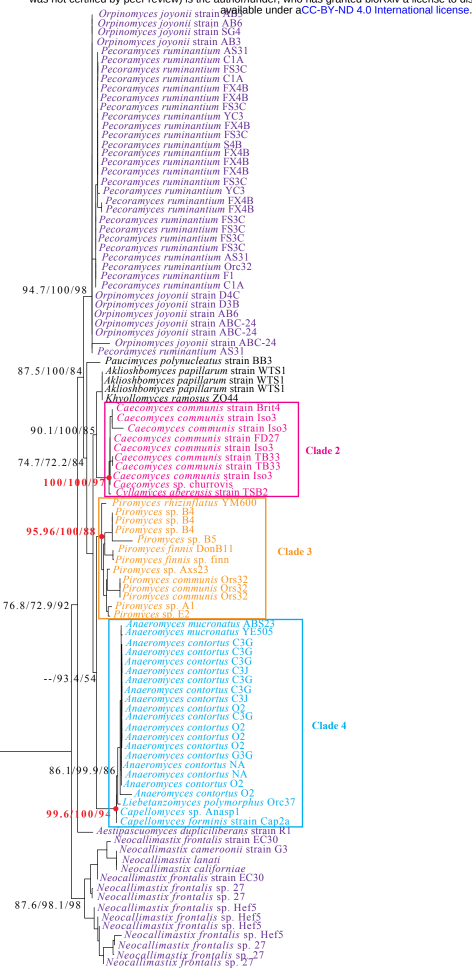
Α

B



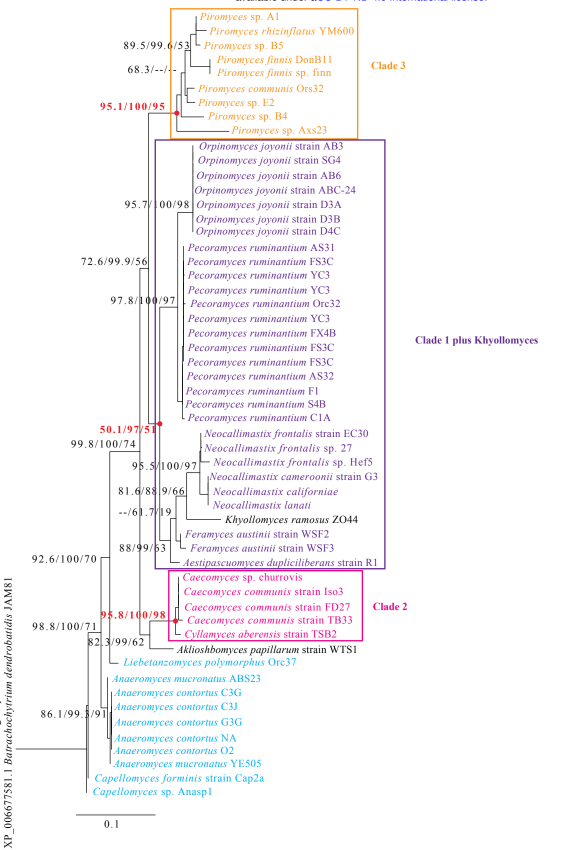
1.0



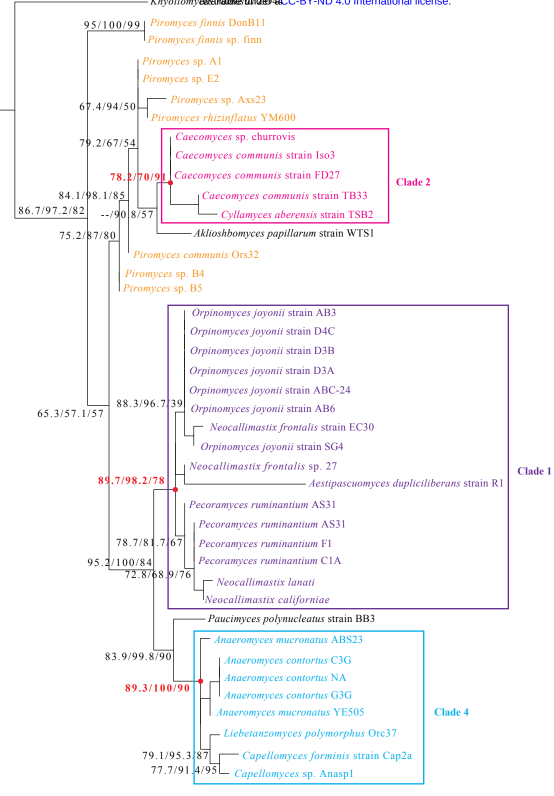


JAJ42635.1 Batrachochytrium dendrobatidis JEL423

Outgroup to



Outgroup to



OAJ38128.1 Batrachochytrium dendrobatidis JEL423

Outgroup to