

1 *Running Title: Blastocystis update*

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5 *Title: Current status of Blastocystis: a personal view*

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18 Abstract

19

20 Despite *Blastocystis* being one of the most widespread and prevalent
21 intestinal eukaryotes, its role in health and disease remains elusive. DNA-
22 based detection methods have led to a recognition that the organism is much
23 more common than previously thought, at least in some geographic regions
24 and some groups of individuals. Molecular methods have also enabled us to
25 start categorizing the vast genetic heterogeneity that exists among
26 *Blastocystis* isolates, wherein the key to potential differences in the clinical
27 outcome of *Blastocystis* carriage may lie.

28

29 In this review we summarize some of the recent developments and advances
30 in *Blastocystis* research, including updates on diagnostic methods, molecular
31 epidemiology, genetic diversity, host specificity, clinical significance,
32 taxonomy, and genomics. As we are now in the microbiome era, we also
33 review some of the steps taken towards understanding the place of
34 *Blastocystis* in the intestinal microbiota.

35

36 Keywords: parasite; gut; Stramenopiles; public health; clinical microbiology

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43 1. INTRODUCTION

44

45 It is now over 100 years since Alexeieff [1] first described the intestinal
46 eukaryote *Blastocystis* but, despite the efforts of numerous researchers
47 (especially in recent years), there are still many unknowns surrounding this
48 organism. Most important of these is whether *Blastocystis* causes disease in
49 humans. For every report linking *Blastocystis* with gastrointestinal or other
50 symptoms there is another that finds no such link. There are a number of
51 factors that have contributed to this apparent lack of progress and these will
52 form the basis of this review. We would like to warn the reader at this early
53 stage that we ourselves are convinced only that there are no definitive data
54 yet available to resolve this issue.

55

56 2. TAXONOMY AND EVOLUTION

57

58 In culture, *Blastocystis* is generally spherical with no obvious surface features.
59 When stained, the most common morphological form seen has a large central
60 vacuole of unknown function and the cytoplasm with all the organelles is
61 visible as a thin peripheral layer between the vacuole and the cell membrane
62 (Figure 1). While many morphological forms have been described, the
63 significance of most is unclear, the boundaries between them are not discrete,
64 and some may well represent degenerating forms [2]. We refer the reader to
65 earlier reviews for more details [3-5]. The life-cycle is typical of most gut
66 protists, with a resistant cyst form for transmission and a trophic form that
67 divides by binary fission. More complex and alternative life-cycles have been

68 described (discussed in [5]) but in our opinion there is no conclusive evidence
69 for anything other than this simple two-stage cycle.

70

71 *Blastocystis* has a complicated taxonomic history. It has been viewed as a
72 fungus, a sporozoan and even the cyst of another organism at various points
73 in its history, until 20 years ago [6] when it was finally placed among the
74 Stramenopiles. This is one of the major groups of eukaryotes [7], but one that,
75 to date, contains only a single other human-infective eukaryote, *Pythium*.
76 *Blastocystis* has none of the typical features of a stramenopile, which is in part
77 why identifying its correct relationships took so long.

78

79 Since its classification as a Stramenopile further data have emerged
80 regarding the closest relatives of *Blastocystis*. These turn out to be poorly
81 known flagellated or ciliate-like organisms that live in vertebrate intestines.
82 While most Stramenopiles are free-living and aerobes, *Blastocystis* and its
83 relatives are gut-living and anaerobes, although they do have mitochondrion-
84 like organelles (see later). *Blastocystis* is related specifically to the
85 Proteromonadidae and Slopalinida [8], but these cannot be considered close
86 relatives. However, it seems likely that the common ancestor of these groups
87 of organisms was already living in a gut and an anaerobe.

88

89 The simple spherical morphology of *Blastocystis* mentioned above applies to
90 all members of this genus. This means that morphology is of no use in
91 defining species. Traditionally, *Blastocystis* species have been defined by the
92 identity of their host, with all human *Blastocystis* being assigned to

93 *Blastocystis hominis*. However, even before DNA sequences identified
94 *Blastocystis* as a Stramenopile it had become clear that significant
95 heterogeneity existed among human *Blastocystis*. Using serology,
96 isoenzymes and karyotyping, human *Blastocystis* were being divided into
97 subgroups [4], and this picture of variation was reinforced by direct and
98 indirect DNA sequence analyses [9]. Subsequent data have only added to the
99 diversity and have refined our understanding of this genus.

100

101 Analyses of human *Blastocystis* by different researchers always resulted in
102 the detection of variation, but each group came up with its own nomenclature
103 for the groupings it identified. To resolve this confusion a consensus
104 terminology was agreed [9] and this classification of human *Blastocystis* into
105 numbered subtypes has simplified communication among workers in this field.
106 At the time of the consensus two things were clear: 1. that humans were host
107 to a number of distinct small subunit rRNA gene (SSU-rDNA)-based subtypes
108 of *Blastocystis*, and 2. that most of these subtypes were also found in other
109 mammalian or avian hosts. This meant the host-linked binomial species
110 names were untenable, as the same organism was being called by multiple
111 names. For example, one grouping of *Blastocystis hominis* proved to be
112 genetically indistinguishable from *Blastocystis ratti*; both are now known as
113 *Blastocystis* subtype 4 (ST4).

114

115 The current taxonomy of *Blastocystis* follows a distinct structure for mammal
116 and bird organisms compared to all others [10]. The mammalian/avian
117 *Blastocystis* are subdivided into seventeen subtypes (STs), nine of which

118 (ST1–ST9) have been found in humans. There is host range overlap
119 observed for many of these organisms (Figure 2). *Blastocystis* from reptiles,
120 amphibia and invertebrates retain Linnean binomial names for the most part.
121 This is largely because little investigation of diversity and host range of these
122 *Blastocystis* has been undertaken to date and so the same impetus to change
123 the nomenclature has not existed. Whether a similar situation involving broad
124 host-range and large genetic diversity will be uncovered in those organisms
125 remains to be seen; it seems likely, and therefore the nomenclature of
126 *Blastocystis* in those hosts may require a similar solution.

127

128 3. GENETIC DIVERSITY AND HOST SPECIFICITY

129

130 Subtypes of *Blastocystis* are discrete and no intermediate variants have been
131 uncovered to date despite extensive sampling from around the world.
132 However, many host species remain to be sampled, so this picture may
133 change. Guidance on how and when to define a new subtype has been
134 published [11]. The recommendation is that a minimum of 5% sequence
135 divergence from the SSU-rDNA of known subtypes is required before defining
136 a new subtype is appropriate. One of the reasons for establishing this
137 boundary is that *Blastocystis* subtypes are often assigned based on the
138 sequence with the closest similarity in sequence database searches, without
139 taking into account the degree of similarity. So a sequence that actually
140 represents a new subtype may be assigned to an existing subtype. This
141 misattribution has been a problem in some existing cases, for example ST13,
142 as discussed in reference [10]. Unfortunately, information attached to entries

143 in GenBank databases are rarely corrected and this can result in
144 misidentifications being propagated forward in the literature.

145

146 The 5% level of divergence to define a new subtype was chosen in part
147 because variation within subtypes can also be substantial, up to at least 3%
148 [11]. Therefore a single 'outlier' sequence that appears to be distinct and
149 potentially a new subtype could eventually merge into an adjacent subtype as
150 more sequences become available. Only as more subtyping data accumulate
151 will the validity of this arbitrary threshold be tested. Note that 5% divergence is
152 the recommendation for establishing new subtypes, where sampling is likely
153 to be limited. The divergence between some existing subtypes (for example,
154 ST6 and ST9) is actually less than 5%. However, sampling is sufficient to give
155 us confidence that these are indeed distinct lineages rather than variants of
156 the same subtype. In other words, 5% divergence has been chosen as quite a
157 stringent criterion and more data may lead to the revision of new subtype
158 definitions in the future..

159

160 As mentioned earlier, nine distinct subtypes have been found in humans
161 (Figure 2). However 95% of human infections sampled belong to one of just
162 four of these subtypes (STs 1-4; [12]) and only one of the human subtypes
163 has not yet been found in another host: ST9 can claim (at present) to be
164 restricted to humans. The four most common STs in humans have also been
165 detected in other hosts. Most frequently these hosts are other primates, but
166 they have also been found in various hoofed mammals, rodents and even
167 birds [10]. Conversely, the rarer subtypes in humans (STs 5-8) are more

168 commonly found in other hosts: ST5 in hoofed animals, STs 6 and 7 in birds,
169 and ST8 in non-human primates. It has been suggested that these rarer
170 subtypes in humans are of zoonotic origin and there is some evidence to
171 support this: ST8 has frequently been found in zookeepers that work with non-
172 human primates [13], and ST5 is prevalent in piggery workers in Australia
173 [14], for example. However, there is no reason to suspect that human
174 infections involving the common STs (STs 1-4) originate from non-human
175 sources except in rare cases.

176

177 Exposure to *Blastocystis*-infected animals alone is not sufficient to result in an
178 infection. For example, ST10 is very common in livestock [10] but is yet to be
179 reported in humans. This suggests that variables other than just body
180 temperature are determining the ability of *Blastocystis* to colonize the human
181 gut; the gut flora may have an impact, for example.

182

183 The degree of genetic diversity within subtypes is quite variable. ST3 is
184 probably the most diverse of the well-studied subtypes – varying by ca. 3% in
185 the SSU-rDNA sequences - while ST4 shows the least variation, especially in
186 humans [15]. Diversity in these subtypes has been further explored using a
187 multi-locus sequence typing approach based on variation in several regions of
188 the mitochondrion-like organelle's genome [15]. MLST data are not yet
189 published for other subtypes. How genetic variability within a subtype is
190 reflected in phenotypic and functional variability is as yet unclear. However,
191 differences in adhesion and drug resistance between strains of *Blastocystis*
192 ST7 have been reported [16].

193

194 Intra-subtype variation has provided further insight into host specificity. For
195 example, ST3 is common in both humans and non-human primates [13].
196 However, MLST analysis divided ST3 into four clades and almost all human
197 samples fell into only one of these clades [15]. Where this was not the case,
198 the individuals concerned had work exposure to non-human primates, again
199 suggesting zoonotic transmission had occurred [15]. It would be interesting to
200 know whether such host specificity exists between variants within other
201 subtypes that are found in a wide range of mammals and exhibit genetic
202 diversity, like ST10 for example [17].

203

204 MLST has the potential to provide insight into geographic aspects of genetic
205 variation as well. However, this could be confounded by the increasing
206 population mobility in today's world: geographic differences will be starting to
207 break down. To date, it is only subtyping that has provided evidence of
208 geographic differences in *Blastocystis* distribution. Specifically, it has become
209 clear that ST4 has a restricted distribution, being rare or absent in South
210 America, North Africa, and the Middle East, while being the second most
211 common subtype in Europe (summarized in [12]). The reasons for this are
212 obscure, but when combined with the relatively low genetic diversity of ST4 in
213 humans the evidence suggests that ST4 may only have entered the human
214 population relatively recently (perhaps in Europe) and is yet to spread around
215 the world [12]. ST4 is also found in other hosts [10], but there is no link
216 between these hosts and Europe.

217

218

219 4. DIAGNOSIS AND MOLECULAR CHARACTERIZATION

220

221 For most parasites, both direct and indirect diagnostic methods have been
222 developed. Direct methods include those based on morphology (microscopy)
223 and detection of DNA (typically PCR) or antigens (IFA, antigen ELISA, etc.),
224 while indirect methods are based mainly on detection of antibodies [18]. While
225 the potential utility of serology in the indirect detection of *Blastocystis*
226 infections remains unclear, some studies have used serology to look for
227 quantitative differences in antibody responses between symptomatic and
228 asymptomatic individuals ([19-20]; see also below).

229

230 With regard to direct detection methods, the use of diverse diagnostic
231 modalities of varying sensitivity may very well have impaired attempts to
232 define the role of *Blastocystis* in health and disease [21-23]. Molecular
233 methods developed to detect *Blastocystis* in genomic DNA extracted directly
234 from fresh stool have highlighted the sensitivity shortcomings of diagnostic
235 methods such as the traditional 'ova and parasites' (O&P) work-up (used to
236 detect cysts of protozoa and larvae and eggs of helminths), culture methods,
237 and permanent staining of fixed fecal smears [24-26].

238

239 Simple stains like Lugol's iodine can be used as a quick aid to the
240 identification of *Blastocystis* in fecal smears or concentrates; the organism is
241 otherwise difficult to differentiate from other structures seen in unstained
242 preparations due to the lack of diagnostic morphological features. Trichrome

243 staining is one of several permanent stains used for detection of trophic forms
244 of protozoa in feces. *Blastocystis* stains characteristically with Trichrome, and
245 this method had a specificity and sensitivity of 100% and 82%, respectively, in
246 a study by Stensvold et al. [24].

247

248 Despite being the primary diagnostic tool worldwide, the use of microscopy to
249 detect *Blastocystis* has limited utility in clinical microbiology laboratories and
250 in generating data for clinical and epidemiological purposes: 1) Microscopy of
251 fecal concentrates - the commonly applied O&P method - has very low
252 sensitivity in detecting *Blastocystis* [24, 27]; 2) there is no consensus on the
253 importance of the cell numbers (see below) or the various morphological
254 forms reported; and 3) microscopy cannot distinguish between genetically
255 highly dissimilar organisms (STs), which may differ in their clinical
256 significance, a situation potentially similar to *Entamoeba histolytica* and
257 *Entamoeba dispar*. Nevertheless, there are situations in which microscopy
258 may serve a purpose, such as those aiming to verify the presence of
259 *Blastocystis* in various types of non-human samples, including those of
260 environmental and animal origin, to inform hypotheses on transmission. For
261 instance, a recent study used microscopy to identify *Blastocystis* in various
262 environmental samples, including food, water, and fomites [28].

263

264 Xenic *in vitro* culture (XIVC) is defined as culture in the presence of an
265 undefined bacterial flora. *Blastocystis* can be grown and propagated xenically
266 in a variety of media [29, 30]. Perhaps due to its simplicity and low cost,
267 Jones' medium has been popular for both detecting and maintaining

268 *Blastocystis*; another medium often used for isolation is Robinson's [29], while
269 we have also used LYSGM (a variant of TYSGM-9; [31]) for propagation when
270 large numbers of cells are needed. XIVC as a diagnostic tool using Jones'
271 medium has a sensitivity ranging from 52%—79% compared with real-time
272 PCR assays [26, 32].

273

274 The diagnostic utility of Ag-ELISA and immunofluorescent antibody staining
275 methods for the detection of *Blastocystis*, including commercial kits such as
276 ParaFlor B (Boulder Diagnostics, Boulder, CO, USAa), coproELISA™
277 Blastocystis (Savyon Diagnostics, Ashdod, Israel), and Blasto-Fluor
278 (Antibodies Inc., Davis, CA, USA), is as yet unclear, since these assays have
279 been used in only a limited number of studies and applied to only a very
280 limited number of samples [33-37]. The utility of such assays remains
281 unknown as the range of subtypes they detect is unclear.

282

283 The first diagnostic PCR for *Blastocystis* was introduced in 2006 [25] but it
284 was later suspected to exhibit preferential amplification of some subtypes over
285 others. Since then, three diagnostic real-time PCR assays have been
286 reported. A real-time PCR based on an unknown *Blastocystis* target using
287 FRET probes was validated against ST1, ST3, and ST4 [38]. A SYBR green
288 real-time PCR used the SSU rRNA gene for detection of *Blastocystis*-specific
289 DNA (ST1–ST9), and subsequent subtyping was performed by melting curve
290 analysis [26]. The relatively large PCR product used (320 to 342 bp,
291 depending on the subtype) may impair the sensitivity of this test—especially
292 when DNA quality is not optimal—and the specificity of the assay was 95%.

293 The third real-time assay, using a hydrolysis probe based on the SSU rRNA
294 gene, was characterized by 100% specificity and ability to detect all nine
295 subtypes identified in humans so far [32]. The use of real-time PCR in large-
296 scale surveys would assist in identifying whether the development of
297 symptoms is related to infection intensity by simple analysis of threshold cycle
298 (C_t) values for individual samples, as this enables quantitation of the amount
299 of *Blastocystis*-specific DNA present. The same DNA samples may also be
300 used for subtyping and MLST protocols, hence allowing the detection and
301 evaluation of genetic diversity as well as the simple presence of *Blastocystis*
302 [22]. *Blastocystis* has also been included as a diagnostic target in commercial
303 gastrointestinal pathogen diagnostic panels such as Feconomics® (Salubris
304 Inc, Boston, USA), EasyScreen™ Enteric Parasite Detection Kit (Genetic
305 Signatures, Sydney, Australia), and NanoChip® (Savyon Diagnostics, Israel).
306

307 While the potency of DNA-based methods is evident, they do not allow the
308 evaluation of whether differences in morphotypes are important. Several
309 different forms of *Blastocystis* have been described, including the avacuolar,
310 vacuolar, multivacuolar, granular, ameboid, and cyst stages. Although there
311 are a few reports of ameboid stages being detected only in symptomatic
312 *Blastocystis* carriers [eg. [39]], there is no consensus regarding the
313 significance of the different forms. Moreover, as mentioned earlier, it is not
314 clear whether some of these forms represent life-cycle stages, or are artifacts
315 resulting from exposure to oxygen or other stresses [2]. Relatively few studies
316 on the cyst stage are available [40-42], which is remarkable given that this is

317 the stage that allows survival of the parasite in the environment and
318 transmission to a new host.

319

320 The high sensitivity of qualitative PCR for detection of *Blastocystis* DNA in
321 stool was reinforced by a recent study of *Blastocystis* in Senegalese children
322 [43], where the prevalence of *Blastocystis* among 93 children with and without
323 gastrointestinal symptoms was 100%. When prevalence is so high there will
324 be little incentive for including *Blastocystis* PCR as a screening tool in the
325 clinical microbiology laboratory. However, where treatment of a patient with
326 *Blastocystis* has been undertaken, PCR methods are useful in post-treatment
327 follow-up to evaluate treatment efficacy.

328

329 This leads to one of the fundamental questions for clinical microbiology labs:
330 When is testing for *Blastocystis* appropriate? Data currently emerging indicate
331 that *Blastocystis* can be more common in individuals with a healthy GI system
332 than in patients with organic and functional bowel diseases (see below).

333 Therefore, the inclusion of *Blastocystis* as a specific target in screening
334 panels, alongside known pathogens such as *Giardia*, *Cryptosporidium*, and
335 *Entamoeba histolytica*, currently appears to make little sense in the clinical
336 microbiology laboratory. The presence of *Blastocystis* in stool samples most
337 likely implies that the carrier has been exposed to fecal-oral contamination,
338 which should prompt the laboratory to look more closely for the presence of
339 pathogens transmitted in the same way. However, since *Blastocystis* may
340 colonize the human colon for more than 10 years [44], it may be impossible to
341 identify when this contamination happened. This has important implications

342 for the interpretation of clinical microbiology lab results. *Blastocystis* is
343 sometimes detected in stool samples of patients with diarrhea or other
344 gastrointestinal symptoms and in the absence of proven pathogens, so
345 clinicians might conclude that *Blastocystis* could be the cause of the
346 symptoms. If it is known that the infection is recent, the organism could
347 certainly be viewed as a potential cause of the symptoms; however, in most
348 cases it will be impossible to rule out that it has been present in the gut for
349 months - even years - and therefore is an incidental finding.

350

351 Another dilemma is the question of whether or not to report the presence of
352 *Blastocystis* in stool samples given that it is so common. Several studies have
353 sought to address this by setting a threshold number of *Blastocystis*
354 organisms detected microscopically per visual field at a specified
355 magnification before scoring the sample as positive; usually this has been set
356 at 5 organisms per 40x field (see references in [5]). However, the rationale for
357 this is unclear. It is known that shedding of both trophic and cyst forms of the
358 organism is irregular [45]. Moreover, several factors may influence the
359 number of organisms seen per visual field, including whether or not the
360 sample was fresh or preserved prior to analysis, and if preserved whether or
361 not the sample was fresh at the time of fixation. Real-time PCR would be
362 more sensitive and less affected by some of these variables.

363

364 In the event that symptoms are eventually linked to specific subtypes,
365 including those individual subtypes as specific targets in diagnostic panels
366 would be more relevant than including a general target for *Blastocystis*.

367 Subtype-specific PCRs already exist, and barcoding of *Blastocystis* DNA
368 amplified by generic primers can also be performed [46, 47]. To date,
369 diagnostic PCR methods have been developed and validated only for human
370 clinical samples; no validated PCR method for detecting *Blastocystis* in
371 environmental samples is yet available to the knowledge of the authors.

372

373 Given the extensive cryptic genetic diversity of *Blastocystis* [10, 15, 48], a
374 number of tools have been developed to map its molecular epidemiology.
375 Among these tools, two in particular have been widely used. A PCR assay for
376 detecting subtypes using sequence-tagged-site (STS) primers was developed
377 and refined in the early 1990s [49]. This approach involves the use of seven
378 PCR reactions, one for each of subtypes 1—7, and should be viewed as
379 comprising a diagnostic method for each of these subtypes, circumventing the
380 need for sequencing. The other method involves analysis of SSU rDNA
381 variation. This approach has been developed independently by several
382 groups, each of which used different regions of the SSU rRNA gene as
383 markers [24-25, 50-56]. The barcoding method mentioned above, developed
384 in 2006 by Scicluna et al., is one such example [46]. A comparison of the STS
385 method and barcoding showed that barcoding should be preferred where
386 possible for a variety of reasons [47]. First and foremost, barcoding enables
387 the detection of subtypes beyond STs 1—7 and further scrutiny of genetic
388 diversity. The barcode region has also been validated as a marker of overall
389 genetic diversity of *Blastocystis* [15].

390

391 Barcoding uses the primers RD5 and BhRDr, which amplify ~600 bp at the 5'-
392 end of the SSU rRNA gene. Comparison of phylogenetic trees obtained by
393 analysis of barcoding sequences with those obtained using concatenated
394 sequences obtained by MLST (reflecting loci in the genome of the
395 mitochondrion-like organelle) demonstrated the appropriateness of using the
396 barcode region as a surrogate marker for overall genome diversity in this
397 particular organism [15]. The drawbacks of barcoding compared to the STS
398 method are that sequencing is required and that mixed subtype infections
399 may not always be evident in sequence chromatograms, and, even if they are,
400 they may prove difficult to decipher [47]. On the other hand, barcoding
401 enables more subtle analyses, namely SSU rDNA allele analysis [15]. A
402 public database is available (<http://pubmlst.org/blastocystis/>) that includes a
403 sequence repository for barcode sequences and those obtained by MLST. It
404 also has a BLAST facility, where individual or bulk fasta files can be uploaded
405 and analyzed for rapid identification of subtype and allele number, hence
406 eliminating the need for phylogenetic analysis. To date, 35 SSU rDNA alleles
407 within ST3 have been identified, whereas the number of SSU rDNA alleles for
408 ST4 and some other subtypes remains much more limited. However, some of
409 the allelic variation included is the result of sequencing of cloned DNA;
410 intragenomic SSU rDNA polymorphism has been reported [57, 58], and such
411 polymorphism will likely go unnoticed when sequences obtained directly from
412 PCR products are studied.

413

414 There is no doubt that DNA-based methods now enable us to carry out large
415 and well-designed research studies that are dependent on accurate detection

416 and molecular characterization of *Blastocystis*. Such studies are required to
417 produce data that can shed light on the role of this organism in human health
418 and disease with a view to potentially developing diagnostics, biomarkers, and
419 therapies, including antimicrobial or probiotic agents, as appropriate.

420

421 5. CLINICAL SIGNIFICANCE AND EPIDEMIOLOGY

422

423 Even after more than 100 years, the role of *Blastocystis* in human health and
424 disease remains obscure. While *Blastocystis* has been speculated to be
425 involved in a range of organic and functional bowel diseases, it is clear that
426 asymptomatic carriage is common. This does not mean that *Blastocystis* does
427 not cause disease. The situation may resemble that for *Giardia*, where many
428 infections are asymptomatic (for example [59]), and *Entamoeba histolytica*,
429 where the proportion of symptomatic infections is at most 10% [60]. Case
430 reports and surveys continue to be published with regularity, mostly indicating
431 a link between *Blastocystis* and symptoms, although not always. We do not
432 propose to evaluate all the evidence here. However we do wish to highlight
433 two common issues: 1. Identification of an appropriate control group for
434 survey studies can be problematic; and 2. Excluding all other possible
435 etiologic agents or non-infectious causes of intestinal symptoms is almost
436 impossible.

437

438 While distinctive intestinal pathology has been clearly linked to the intestinal
439 protists *Giardia*, *Cryptosporidium*, and *Entamoeba*, there is little – if any –
440 evidence for direct pathology caused by *Blastocystis*. Phagocytosis of red

441 blood cells is a well-known feature of *Entamoeba histolytica* that correlates
442 with virulence; there is only one study reporting phagocytosis in *Blastocystis*
443 [61]. No *Blastocystis* proteins such as glycoproteins or lectins that could
444 facilitate attachment to the gut epithelial layer have been identified, although
445 Denoeud et al. [57] have speculated that *Blastocystis* hydrolases might be
446 able to alter the colonic mucus layer (see below). It is generally accepted that
447 *Blastocystis* is non-invasive as well as lacking the ability to phagocytize the
448 microbiota or host-derived material.

449

450 When examining tissue sections from pig intestines, Fayer et al. [62] found
451 *Blastocystis* primarily in the lumen, usually associated with digested food
452 debris, and although sometimes in close proximity to or appearing to adhere
453 to the epithelium, there were no cells penetrating to the epithelium or the
454 lamina propria. These observations were confirmed by Wang et al. [63], who
455 did not observe any obvious pathology in histological sections of porcine gut
456 mucosal biopsies. In the latter study, *Blastocystis* cells were observed as
457 vacuolar/granular forms found within luminal material or in close proximity to
458 epithelial cells, with no evidence of attachment or invasion. When *Blastocystis*
459 is observed adhering to the epithelium in histological preparations it should be
460 kept in mind that histological procedures are likely to dissolve and eliminate
461 the mucus layer that is potentially separating *Blastocystis* from the mucosa in
462 vivo.

463

464 Despite the absence of invasion, discrete non-specific colonic inflammation
465 has been reported in a patient with both urticaria and what was characterized

466 as 'heavy *Blastocystis* colonization'; *Blastocystis* eradication resulted in
467 symptom resolution [64]. There are also some reports of *Blastocystis* having
468 been found extra-intestinally, but in those cases it has not been possible to
469 rule out that the presence of *Blastocystis* at these sites was merely a result of
470 incidental or secondary colonization resulting from damage generated by
471 other microorganisms or anatomical anomalies [65-68].

472

473 *Blastocystis* is one of several organisms to have been linked to Irritable Bowel
474 Syndrome (IBS), including post-infectious IBS [69-71]. Genome analysis by
475 Poirier et al. [72] identified various genes encoding hydrolases and serine and
476 cysteine proteases, and the authors speculated that these potential virulence
477 factors could be triggers of IBS by alteration of the mucus layer and
478 interaction with tight junctions.

479

480 Cross-sectional studies testing the hypothesis that *Blastocystis* is linked to
481 IBS mostly assume that, if the organism is associated with the disease, it
482 should be more common in patients with IBS symptoms. The outcomes of
483 such studies have been mixed, with some finding a higher prevalence of
484 *Blastocystis* in IBS patients and some finding no difference or even lower
485 prevalence (summarized in [12]). A few have looked at the subtype
486 distribution, but although they have generally found differences between IBS
487 and non-IBS patients, there is no consistency regarding the subtypes
488 associated with IBS (summarized in [12]). IBS itself presents a diverse
489 picture, with patients having diarrhea, constipation or a mixture of symptoms

490 [69]. Even fewer investigations have been performed to look at potential links
491 between *Blastocystis* and subgroups within IBS.

492

493 IBS patients are likely to have multiple tests performed before a diagnosis is
494 made and, because of this, a common finding may well be *Blastocystis* in the
495 stool, which might then be suspected of being the agent responsible for the
496 symptoms if no other candidates have been uncovered. So *Blastocystis* may
497 be more commonly detected in IBS patients simply because the investigations
498 are more thorough. Post-infectious IBS - a term describing the development of
499 IBS following treatment of an infection with antimicrobials [71] – adds another
500 complication, as the actual trigger for IBS may have been eliminated by
501 antimicrobial treatment, leaving *Blastocystis* behind to take the blame. It is
502 also impossible to exclude that *Blastocystis* was the initial trigger of IBS even
503 if it is no longer present. The potential links, if any, between *Blastocystis* and
504 IBS may be impossible to prove or disprove without large longitudinal cohort
505 studies.

506

507 One of the most interesting recent findings is that *Blastocystis* could be a
508 marker of gastrointestinal health rather than a cause of disease. This may in
509 fact not be surprising, given that we have been unable to reach a consensus
510 on a role for the organism in disease despite the large number and wide
511 range of investigations undertaken. A recent study identified *Blastocystis* as a
512 common member of the healthy human gut microbiota, with greater than 50%
513 of the healthy background population colonized [44]. Moreover, long-term
514 colonization trends were also noted; the same strains were present in the

515 same hosts for up to 10 years [44]. A lower prevalence of *Blastocystis* in IBS
516 patients (n = 189) compared with healthy controls (n = 297), 14.5% versus
517 22% respectively (p = 0.09), was also highlighted in a recent study [73]; the
518 prevalence of *Dientamoeba fragilis* also differed significantly between the two
519 groups, with *D. fragilis* being similarly more common in individuals without
520 gastrointestinal symptoms. Another study, this time involving 96 healthy
521 controls and 100 patients with Inflammatory Bowel Disease (IBD) - a disease
522 affecting about 12,000 individuals in Denmark alone, 0.2% of the population -
523 detected a significantly lower prevalence of *Blastocystis* in IBD patients
524 compared with healthy controls (p < 0.05), with only 5/100 IBD patients being
525 colonized by *Blastocystis* compared with 18/96 controls [74-75]. Interestingly,
526 four of the five positive IBD patients were in an inactive stage of the disease;
527 only 1/42 patients with active IBD was a carrier.

528

529 Whether it is linked to gastrointestinal health or disease, it is clear that
530 *Blastocystis* is much more common than previously reported, reaching a
531 prevalence of 100% in some cohorts [43]. Individuals in communities with high
532 prevalence may become and remain infected from a very young age, while in
533 other communities, particularly where the overall prevalence is low, many
534 individuals may acquire *Blastocystis* later in life. For now, it is uncertain
535 whether the age at colonization - including whether *Blastocystis* becomes a
536 stable member of the intestinal microbiota from early on - is of any clinical
537 importance. It could be that in some regions of the world, *Blastocystis* might
538 be an 'emerging pathogen'.

539

540 While recent observations suggest that *Blastocystis* colonization may be
541 inversely correlated with intestinal disease [44], we now know that the
542 bacterial component of the gut microbiota in IBS, IBD, and other intestinal
543 diseases is significantly different to that of the healthy human gut [69, 76].
544 Importantly, this may in fact indicate that *Blastocystis* is dependent on other
545 components of the microbiota to colonize and maintain a stable colonization in
546 the human gut. To test this prediction, we recently obtained access to data
547 from the MetaHIT Consortium (<http://www.metahit.eu/>), originally generated to
548 identify associations between intestinal bacterial communities and disease
549 patterns, including obesity, diabetes, and IBD [77]. From the data, we were
550 able to extract *Blastocystis*-specific DNA signatures, which enabled us to (1)
551 identify the relative prevalence of *Blastocystis* in each of the study groups,
552 and (2) to perform a preliminary investigation of the association between
553 *Blastocystis* and bacterial communities, in this case the so-called
554 ‘enterotypes’ [77]. Our analysis [78] showed that: 1) *Blastocystis* was indeed
555 negatively associated with disease and absent in all 13 patients with Crohn’s
556 disease (although not all studies have found this; [79]); and 2) very
557 intriguingly, *Blastocystis* was negatively associated with the *Bacteroides*
558 enterotype ($p < 0.0001$, unpublished data). This finding may be linked to the
559 fact that the *Bacteroides* enterotype—compared with the *Prevotella* and the
560 *Ruminococcus* enterotypes—is characterized by low microbial diversity, and
561 this could therefore indicate that *Blastocystis* requires high overall microbial
562 diversity to become established in the human colon. However, it could also be
563 that some other unknown feature(s) of the enterotype may be responsible for
564 determining *Blastocystis* colonization, such as bacterial metabolic by-

565 products. There is no doubt that studies of *Blastocystis* in the context of
566 intestinal bacterial communities and host physiology and immunity are likely to
567 advance our understanding of the clinical significance of *Blastocystis*. The
568 apparent impact of the gut flora on *Blastocystis* colonization may also mean
569 that standard animal models may be of limited use in exploring the effects of
570 *Blastocystis* on the human gut.

571

572 Comparing both bacterial and eukaryotic microbial communities in samples
573 from 23 individuals from agrarian communities in Malawi following traditional
574 lifestyles and from 13 individuals residing in Pennsylvania and Colorado,
575 USA, following a modern lifestyle, Parfrey et al. [80] recently showed that the
576 Malawi population harbored a diverse community of protists, including
577 *Blastocystis*, when compared to the North American populations, and that the
578 overall organismal diversity in the Malawian human gut is comparable to that
579 in other mammals. These, and other, data could indicate that the declining
580 diversity of the human bacterial microbiota identified in the West compared
581 with populations with traditional agrarian lifestyle has led to a reduced
582 prevalence of *Blastocystis* in Western populations [81].

583

584 It is also clear that geographical differences in subtype distributions may result
585 in geographical differences in the clinical significance of the parasite. There is
586 precedent in *Entamoeba* for cryptic genetic differences underlying differences
587 in the clinical outcome of infection (the *E. histolytica*/*E. dispar* story; [60]). So
588 a working hypothesis over the past few years has been that differences
589 between the clinical outcome of *Blastocystis* infection may reflect genetic

590 differences in the organism. Hence, dozens of studies from all over the world
591 have sought to identify *Blastocystis* STs in both healthy and symptomatic
592 individuals (summarized in [11]). The distribution of subtypes across the major
593 geographical regions is depicted in Figure 3. So far, no particular subtype has
594 been linked consistently to disease. However, such a finding might not be
595 unexpected if the distribution of subtypes is uneven. While ST1, ST2, and ST3
596 appear to have a global distribution, current data suggest that ST4 is confined
597 mainly to Europe. ST4 was the only subtype identified in Danish patients with
598 acute diarrhea, but the overall prevalence of the parasite was also lower in
599 this group of patients than in others that have been studied in Denmark [82].
600 ST4 also dominated in symptomatic patients in Spain [83].

601

602 A significant gap in clinical *Blastocystis* research is the lack of large
603 randomized controlled clinical treatment trials [84-87]. To date these have
604 produced inconsistent and indeed contradictory results. It appears that no
605 single drug or drug combination currently in use consistently results in reliable
606 *Blastocystis* eradication [88-90]. Metronidazole has traditionally been used to
607 treat anaerobic microorganisms, including *Entamoeba* and *Giardia*; however,
608 its effect on *Blastocystis* has in some studies been minimal, with an
609 eradication rate as low as 0%. Even the use of combinations such as
610 diloxanide furoate, secnidazole, and trimethoprim/sulfamethoxazole or
611 nitazoxanide may not result in consistent eradication [90].

612

613

614 6. GENOMICS

615

616 With the advances in sequencing technology in recent years it has become
617 possible to sequence eukaryotic genomes quickly and relatively inexpensively
618 compared with even a few years ago. Perhaps surprisingly, the published
619 *Blastocystis* nuclear genome sequences at the time of writing are for ST7,
620 obtained by 'traditional' Sanger sequencing [57], and ST4, obtained by next
621 generation sequencing [91]. Others have not yet appeared in print despite
622 anecdotal evidence that suggests a flood of new data is about to arrive.

623

624 However, *Blastocystis* has two genomes. In addition to the nuclear genome it
625 also contains an organelle genome. In contrast to most anaerobic eukaryotes,
626 *Blastocystis* has mitochondrion-like organelles that have a quite normal
627 appearance under the transmission electron microscope (see [4]). It was
628 known for many years that these organelles contained DNA, based on
629 staining properties, but it was not until 2007 that the coding potential of these
630 molecules was uncovered. Two groups published sequences of the genomes
631 present in the mitochondrion-like organelle in three subtypes – STs 1, 4 and 7
632 [92-93]. The gene content and gene order of the 27-29 kilobasepair circular
633 molecules was identical, although the sequence divergence was
634 considerable. Subsequently, mitochondrion-like organelle genomes from
635 additional subtypes have been obtained (unpublished data) and these initial
636 observations have been upheld.

637

638 The gene content of the genome of the mitochondrion-like organelle is distinct
639 from the more familiar ones from mammals and yeast. Particularly notable is

640 the absence of any genes encoding cytochrome and ATPase subunits and
641 the presence of a number of ribosomal protein genes. In common are the
642 genes encoding ribosomal RNAs and several tRNAs plus NADH
643 dehydrogenase (Complex I) subunits. The nuclear genomes and expressed
644 sequence tag (EST) surveys that are available confirm that the *Blastocystis*
645 mitochondrion-like organelle has only retained complexes I and II of the
646 electron transport chain, a characteristic shared with certain other anaerobic
647 eukaryotes. However, many other features of mitochondrial metabolism are
648 also present [31, 57]. This is in contrast to the situation in, for example,
649 *Giardia* and *Entamoeba* where the genome has been lost completely and the
650 function of the resulting organelles (known as mitosomes) has become highly
651 reduced. Whether the *Blastocystis* organelle would follow a similar path given
652 enough time is impossible to predict.

653

654 The only published nuclear genomes at this time are for ST4 and ST7.
655 However, a recently published report on polyadenylation in *Blastocystis* also
656 includes data on a ST1 genome, suggesting its publication is imminent. The
657 polyadenylation report uncovered a unique situation in *Blastocystis*, where
658 around 15% of the stop codons in messenger RNAs are created through the
659 cleavage of a precursor and addition of the poly A tail to the mRNA [94]. This
660 is unprecedented outside of mitochondria. Given the degree of genetic
661 divergence between subtypes, comparative genomics may well reveal
662 significant differences between features of their nuclear genomes as well as
663 confirming genus-wide peculiarities, as in this case.

664

665 Overall, the *Blastocystis* nuclear genome is quite small (under 19 Mb) with
666 relatively few genes (just over 6,000), quite a few of which appear to have
667 been acquired by horizontal gene transfer. Introns are numerous and small,
668 but repetitive DNA is rare. Of note is the fact that individual ribosomal RNA
669 cistrons are sometimes present in subtelomeric regions of the genome rather
670 than being exclusively found in long tandem arrays as in many other
671 eukaryotes [57].

672

673 7. CONCLUSION

674

675 *Blastocystis* is one of the most successful intestinal eukaryotes identified to
676 date, being able to infect a wide range of host species. It may reside in the gut
677 for years on end and appears to show remarkably little susceptibility to
678 standard chemotherapeutic interventions, although analysis of biochemical
679 pathways identified through genome sequencing may generate some new
680 directions for drug interventions. However, the recognition of a high
681 prevalence of *Blastocystis* in healthy populations, identified using sensitive
682 molecular diagnostic tools, has heralded a paradigm shift in clinical
683 *Blastocystis* research. Studies of the gut microbiota in people with and without
684 *Blastocystis* are likely to provide valuable - if not critical - information to help
685 determine the role of *Blastocystis* in human health and disease.

686

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690

691 9. REFERENCES

692

693 [1] A. Alexieff, Sur la nature des formations dites "Kystes de *Trichomonas*
694 *intestinalis*", C. R. Soc. Biol. 71 (1911) 296-298.

695 [2] A.A. Vdovenko, *Blastocystis hominis*: origin and significance of
696 vacuolar and granular forms, Parasitol. Res. 86 (2000) 8-10.

697 [3] C.H. Zierdt, *Blastocystis hominis*: past and future, Clin. Microbiol. Rev.
698 4 (1991) 61-79.

699 [4] D.J. Stenzel, P.F. Boreham, *Blastocystis hominis* revisited, Clin.
700 Microbiol. Rev. 9 (1996) 563-584

701 [5] K.S.W. Tan, New insights on classification, identification, and clinical
702 relevance of *Blastocystis* spp., Clin. Microbiol. Rev. 21 (2008) 639-
703 665.

704 [6] J.D. Silberman, M.L. Sogin, D.D. Leipe, C.G. Clark, Human parasite
705 finds taxonomic home, Nature 380 (1996) 398.

706 [7] S.M. Adl, A.G. Simpson, C.E. Lane, J. Lukeš, D. Bass, S.S. Bowser,
707 M.W. Brown, F. Burki, M. Dunthorn, V. Hampl, A. Heiss, M.
708 Hoppenrath, E. Lara, L. Le Gall, D.H. Lynn, H. McManus, E.A.
709 Mitchell, S.E. Mozley-Stanridge, L.W. Parfrey, J. Pawlowski, S.
710 Rueckert, R.S. Shadwick, C.L. Schoch, A. Smirnov, F.W. Spiegel, The
711 revised classification of eukaryotes, J. Eukaryot. Microbiol. 59 (2012)
712 429-493.

- 713 [8] M Kostka, I Čepička, V Hampl, J Flegr, Phylogenetic position of
714 *Karotomorpha* and paraphyly of Proteromonadidae, Mol. Phylogenet.
715 Evol. 43 (2007) 1167-1170.
- 716 [9] C.R. Stensvold, G.K. Suresh, K.S.W. Tan, R.C.A. Thompson, R.J.
717 Traub, E. Viscogliosi, H. Yoshikawa, C.G. Clark, Terminology for
718 *Blastocystis* subtypes: a consensus, Trends Parasitol. 23 (2007) 93-
719 96.
- 720 [10] M.A. Alfellani, D. Taner-Mulla, A.S. Jacob, C.A. Imeede, H.
721 Yoshikawa, C.R. Stensvold, C.G. Clark, Genetic diversity of
722 *Blastocystis* in livestock and zoo animals, Protist 164 (2013) 497-509.
- 723 [11] C.G. Clark, M. van der Giezen, M.A. Alfellani, C.R. Stensvold, Recent
724 developments in *Blastocystis* research, Adv. Parasitol. 82 (2013) 1-32.
- 725 [12] M.A. Alfellani, C.R. Stensvold, A. Vidal-Lapiedra, E.S.U. Onuoha, A.F.
726 Fagbenro-Beyioku, C.G. Clark, Variable geographic distribution of
727 *Blastocystis* subtypes and its potential implications, Acta Trop. 126
728 (2013) 11-18.
- 729 [13] M.A. Alfellani, A.S. Jacob, N. Ortíz Perea, R.C. Krecek, D. Taner-
730 Mulla, J.J. Verweij, B. Levecke, E. Tannich, C.G. Clark, C.R.
731 Stensvold, Diversity and distribution of *Blastocystis* sp. subtypes in
732 non-human primates, Parasitology 140 (2013) 966-971.
- 733 [14] W. Wang, H. Owen, R.J. Traub, L. Cuttell, T. Inpankaew, H. Bielefeldt-
734 Ohmann, Molecular epidemiology of *Blastocystis* in pigs and their in-

- 735 contact humans in Southeast Queensland, Australia, and Cambodia,
736 Vet. Parasitol. 203 (2014) 264-269.
- 737 [15] C.R. Stensvold, M.A. Alfellani, C.G. Clark, Levels of genetic diversity
738 vary dramatically between *Blastocystis* subtypes, Infect. Genet. Evol.
739 12 (2011) 263-273.
- 740 [16] Z. Wu, H. Mirza, K.S.W. Tan, Intra-subtype variation in enteroadhesion
741 accounts for differences in epithelial barrier disruption and is
742 associated with metronidazole resistance in *Blastocystis* subtype 7,
743 PLoS Negl. Trop. Dis. 8 (2014) e2885.
- 744 [17] C.R. Stensvold, M.A. Alfellani, S. Nørskov-Lauritsen, K. Prip, E.L.
745 Victory, C. Maddox, H.V. Nielsen, C.G. Clark, Subtype distribution of
746 *Blastocystis* isolates from synanthropic and zoo animals and
747 identification of a new subtype, Int. J. Parasitol. 39 (2009) 473-479.
- 748 [18] J.J. Verweij, C.R. Stensvold, Molecular testing for clinical diagnosis
749 and epidemiological investigations of intestinal parasitic infections,
750 Clin. Microbiol. Rev. 27 (2014) 371-418.
- 751 [19] C.H. Zierdt, W.S. Zierdt, B. Nagy, Enzyme-linked immunosorbent
752 assay for detection of serum antibody to *Blastocystis hominis* in
753 symptomatic infections, J. Parasitol. 81 (1995) 127-129.
- 754 [20] R. Nagel, R.J. Traub, M.M. Kwan, H. Bielefeldt-Ohmann, *Blastocystis*
755 specific serum immunoglobulin in patients with irritable bowel

- 756 syndrome (IBS) versus healthy controls, *Parasit. Vectors.* 8 (2015)
757 453.
- 758 [21] C.R. Stensvold, H.V. Nielsen, K. Mølbak, H.V. Smith, Pursuing the
759 clinical significance of *Blastocystis*: diagnostic limitations, *Trends*
760 *Parasitol.* 25 (2009) 23-29.
- 761 [22] C.R. Stensvold, *Blastocystis*: Genetic diversity and molecular methods
762 for diagnosis and epidemiology, *Trop. Parasitol.* 3 (2013) 26-34.
- 763 [23] P.D. Scanlan, C.R. Stensvold, *Blastocystis*: getting to grips with our
764 guileful guest, *Trends Parasitol.* 29 (2013) 523-529.
- 765 [24] C.R. Stensvold, M.C. Arendrup, C. Jespersgaard, K. Mølbak, H.V.
766 Nielsen, Detecting *Blastocystis* using parasitologic and DNA-based
767 methods: a comparative study, *Diagn. Microbiol. Infect. Dis.* 59 (2007)
768 303-307.
- 769 [25] C.R. Stensvold, A. Brillowska-Dabrowska, H.V. Nielsen, M.C.
770 Arendrup, Detection of *Blastocystis hominis* in unpreserved stool
771 specimens by using polymerase chain reaction, *J. Parasitol.* 92 (2006)
772 1081-1087.
- 773 [26] P. Poirier, I. Wawrzyniak, A. Albert, H. El Alaoui, F. Delbac, V. Livrelli,
774 Development and evaluation of a real-time PCR assay for detection
775 and quantification of *Blastocystis* parasites in human stool samples:
776 prospective study of patients with hematological malignancies, *J. Clin.*
777 *Microbiol.* 49 (2011) 975-983.

- 778 [27] T. Roberts, J. Barratt, J. Harkness, J. Ellis, D. Stark, Comparison of
779 microscopy, culture, and conventional polymerase chain reaction for
780 detection of *Blastocystis* sp. in clinical stool samples, Am. J. Trop. Med.
781 Hyg. 84 (2011) 308-312.
- 782 [28] Á.L. Londoño-Franco, J. Loaiza-Herrera, F.M. Lora-Suárez, J.E.
783 Gómez-Marín, [*Blastocystis* sp. frequency and sources among children
784 from 0 to 5 years of age attending public day care centers in Calarcá,
785 Colombia (in Spanish)], Biomedica 34 (2014) 218-227.
- 786 [29] C.G. Clark, L.S. Diamond, Methods for cultivation of luminal parasitic
787 protists of clinical importance, Clin. Microbiol. Rev. 15 (2002) 329-341.
- 788 [30] S. Leelayoova, P. Taamasri, R. Rangsin, T. Naaglor, U. Thathaisong,
789 M. Mungthin, In-vitro cultivation: a sensitive method for detecting
790 *Blastocystis hominis*, Ann. Trop. Med. Parasitol. 96 (2002)803-807.
- 791 [31] A. Stechmann, K. Hamblin, V. Pérez-Brocal, D. Gaston, G.S.
792 Richmond, M. van der Giezen, C.G. Clark, A.J. Roger, Organelles in
793 *Blastocystis* that blur the distinction between mitochondria and
794 hydrogenosomes, Curr. Biol. 18 (2008) 580-585.
- 795 [32] C.R. Stensvold, U.N. Ahmed, L.O. Andersen, H.V. Nielsen,
796 Development and evaluation of a genus-specific, probe-based, internal
797 process controlled real-time PCR assay for sensitive and specific
798 detection of *Blastocystis*, J. Clin. Microbiol. 50 (2012) 1847-1851.

- 799 [33] R. Fayer, M. Santin, D. Macarisin, Detection of concurrent infection of
800 dairy cattle with *Blastocystis*, *Cryptosporidium*, *Giardia*, and
801 *Enterocytozoon* by molecular and microscopic methods, *Parasitol. Res.*
802 111 (2012) 1349-1355.
- 803 [34] F. Dogruman-AI, Z. Simsek, K. Boorum, E. Ekici, M. Sahin, C. Tuncer,
804 S. Kustimur, A. Altinbas, Comparison of methods for detection of
805 *Blastocystis* infection in routinely submitted stool samples, and also in
806 IBS/IBD Patients in Ankara, Turkey. *PLoS One* 5 (2010) e15484.
- 807 [35] S.M. El-Marhoumy, K. Abd EL-Nouby, Z.S. Shoheib, A.M. Salama,
808 Prevalence and diagnostic approach for a neglected protozoon
809 *Blastocystis hominis*, *Asian Pac. J. Trop. Med.* 5 (2015) 51-59.
- 810 [36] R. Gould, K. Boorum, *Blastocystis* surface antigen is stable in
811 chemically preserved stool samples for at least 1 year, *Parasitol. Res.*
812 112 (2013) 2469-2471.
- 813 [37] F. Dogruman-AI, S. Turk, G. Adiyaman-Korkmaz, A. Hananel, L. Levi,
814 J. Kopelowitz, O. Babai, S. Gross, Z. Greenberg, Y. Herschkovitz, I.
815 Mumcuoglu, A novel ELISA test for laboratory diagnosis of *Blastocystis*
816 spp. in human stool specimens, *Parasitol. Res.* 114 (2015) 495-500.
- 817 [38] M.S. Jones, R.D. Ganac, G. Hiser, N.R. Hudson, A. Le, C.M. Whipps,
818 Detection of *Blastocystis* from stool samples using real-time PCR,
819 *Parasitol. Res.* 103 (2008) 551-557.

- 820 [39] T.C. Tan, G.S. Kumar, Predominance of amoeboid forms of
821 *Blastocystis hominis* in isolates from symptomatic patients, Parasitol.
822 Res. 98 (2006) 189-193.
- 823 [40] D.J. Stenzel, P.F. Boreham, A cyst-like stage of *Blastocystis hominis*,
824 Int. J. Parasitol. 21 (1991) 613-615.
- 825
- 826 [41] K.T. Moe, M. Singh, J. Howe, L.C. Ho, S.W. Tan, G.C. Ng, X.Q. Chen,
827 E.H. Yap, Observations on the ultrastructure and viability of the cystic
828 stage of *Blastocystis hominis* from human feces, Parasitol. Res. 82
829 (1996) 439-444.
- 830 [42] I.F. Abou El Naga, A.Y. Negm, Morphology, histochemistry and
831 infectivity of *Blastocystis hominis* cyst, J. Egypt. Soc. Parasitol. 31
832 (2001) 627-635.
- 833 [43] D. El Safadi, L. Gaayeb, D. Meloni, A. Cian, P. Poirier, I. Wawrzyniak,
834 F. Delbac, F. Dabboussi, L. Delhaes, M. Seck, M. Hamze, G. Riveau,
835 E. Viscogliosi, Children of Senegal River Basin show the highest
836 prevalence of *Blastocystis* sp. ever observed worldwide, BMC Infect.
837 Dis. 14 (2014) 164.
- 838 [44] P.D. Scanlan, C.R. Stensvold, M. Rajilić-Stojanović, H.G. Heilig, W.M.
839 De Vos, P.W. O'Toole, P.D. Cotter, The microbial eukaryote
840 *Blastocystis* is a prevalent and diverse member of the healthy human
841 gut microbiota, FEMS Microbiol. Ecol. 90 (2014) 326-330.

- 842 [45] G.D. Vennila, G.S. Kumar, A.K. Anuar, S. Rajah, R. Saminathan, S.
843 Sivanandan, K. Ramakrishnan, Irregular shedding of *Blastocystis*
844 *hominis*, Parasitol. Res. 85 (1999) 162-164.
- 845 [46] S.M. Scicluna, B. Tawari, C.G. Clark, DNA barcoding of *Blastocystis*,
846 Protist 157 (2006) 77-85.
- 847 [47] C.R. Stensvold, Comparison of sequencing (barcode region) and
848 sequence-tagged-site PCR for *Blastocystis* subtyping, J. Clin.
849 Microbiol. 51 (2013) 190-194.
- 850 [48] C.G. Clark, Extensive genetic diversity in *Blastocystis hominis*, Mol.
851 Biochem. Parasitol. 87 (1997) 79-83.
- 852 [49] H. Yoshikawa, Z. Wu, I. Kimata, M. Iseki, I.K.M. Ali, M.B. Hossain, V.
853 Zaman, R. Haque, Y. Takahashi, Polymerase chain reaction-based
854 genotype classification among human *Blastocystis hominis* populations
855 isolated from different countries, Parasitol. Res. 92 (2004) 22-29.
- 856 [50] C.R. Stensvold, M. Lebbad, J.J. Verweij, C. Jespersgaard, G. von
857 Samson-Himmelstjerna, S.S. Nielsen, H.V. Nielsen, Identification and
858 delineation of members of the *Entamoeba* complex by pyrosequencing,
859 Mol. Cell. Probes 24 (2010) 403-406.
- 860 [51] M. Özyurt, Ö. Kurt, K. Mølbak, H.V. Nielsen, T. Haznedaroglu, C.R.
861 Stensvold, Molecular epidemiology of *Blastocystis* infections in Turkey,
862 Parasitol. Int. 57 (2008) 300-306.

- 863 [52] C.R. Stensvold, H.C. Lewis, H.M. Hammerum, L.J. Porsbo, S.S.
864 Nielsen, K.E. Olsen, M.C. Arendrup, H.V. Nielsen, K. Mølbak K,
865 *Blastocystis*: unravelling potential risk factors and clinical significance
866 of a common but neglected parasite, *Epidemiol. Infect.* 137 (2009)
867 1655-1663.
- 868 [53] U. Parkar, R.J. Traub, S. Kumar, M. Mungthin, S. Vitali, S. Leelayoova,
869 K. Morris, R.C.A. Thompson, Direct characterization of *Blastocystis*
870 from faeces by PCR and evidence of zoonotic potential, *Parasitology*
871 134 (2007) 359-367.
- 872 [54] U. Parkar, R.J. Traub, S. Vitali, A. Elliot, B. Levecke, I. Robertson, T.
873 Geurden, J. Steele, B. Drake, R.C.A. Thompson, Molecular
874 characterization of *Blastocystis* isolates from zoo animals and their
875 animal-keepers, *Vet. Parasitol.* 169 (2010) 8-17.
- 876 [55] M. Santín, M.T. Gómez-Muñoz, G. Solano-Aguilar, R. Fayer,
877 Development of a new PCR protocol to detect and subtype *Blastocystis*
878 spp. from humans and animals, *Parasitol. Res.* 109 (2011) 205-212.
- 879 [56] K.H. Wong, G.C. Ng, R.T. Lin, H. Yoshikawa, M.B. Taylor, K.S.W. Tan,
880 Predominance of subtype 3 among *Blastocystis* isolates from a major
881 hospital in Singapore, *Parasitol. Res.* 102 (2008) 663-670.
- 882 [57] F. Denoëud, M. Roussel, B. Noel, I. Wawrzyniak, C. Da Silva, M.
883 Diogon, E. Viscogliosi, C. Brochier-Armanet, A. Couloux, J. Poulain, B.
884 Segurens, V. Anthouard, C. Texier, N. Blot, P. Poirier, G.C. Ng, K.S.W.
885 Tan, F. Artiguenave, O. Jaillon, J.M. Aury, F. Delbac, P. Wincker, C.P.

- 886 Vivarès, H. El Alaoui, Genome sequence of the stramenopile
887 *Blastocystis*, a human anaerobic parasite, *Genome Biol.* 12 (2011)
888 R29.
- 889 [58] D. Meloni, P. Poirier, C. Mantini, C. Noël, N. Gantois, I. Wawrzyniak, F.
890 Delbac, M. Chabé, L. Delhaes, E. Dei-Cas, P.L. Fiori, H. El Alaoui, E.
891 Viscogliosi, Mixed human intra- and inter-subtype infections with the
892 parasite *Blastocystis* sp., *Parasitol. Int.* 61 (2012) 719-722.
- 893 [59] M.G. Tellevik, S.J. Moyo, B. Blomberg, T. Hjøllø, S.Y. Maselle, N.
894 Langeland, K. Hanevik, Prevalence of *Cryptosporidium*
895 *parvum/hominis*, *Entamoeba histolytica* and *Giardia lamblia* among
896 young children with and without diarrhea in Dar es Salaam, Tanzania,
897 *PLoS Negl. Trop. Dis.* 9 (2015) e0004125
- 898 [60] B.S. Pritt, C.G. Clark, Amebiasis, *Mayo Clin. Proc.* 83 (2008) 1154-
899 1159.
- 900
- 901 [61] L.A. Dunn, P.F. Boreham, D.J. Stenzel, Ultrastructural variation of
902 *Blastocystis hominis* stocks in culture, *Int. J. Parasitol.* 19 (1989) 43-56.
- 903
- 904 [62] R. Fayer, T. Elsasser, R. Gould, G. Solano, J. Urban Jr, M. Santin,
905 *Blastocystis* tropism in the pig intestine, *Parasitol. Res.* 113 (2014)
906 1465-1472

907

908 [63] W. Wang, H. Bielefeldt-Ohmann, R.J. Traub, L. Cuttell, H. Owen,
909 Location and pathogenic potential of *Blastocystis* in the porcine
910 intestine, PLoS One. 9 (2014) e103962

911

912 [64] C. Vogelberg, C.R. Stensvold, S. Monecke, A. Ditzen, K. Stopsack, U.
913 Heinrich-Gräfe, C. Pöhlmann, *Blastocystis* sp. subtype 2 detection
914 during recurrence of gastrointestinal and urticarial symptoms, Parasitol.
915 Int. 59 (2010) 469-471.

916

917 [65] H.L. Santos, F.C. Sodré, H.W. de Macedo, *Blastocystis* sp. in splenic
918 cysts: causative agent or accidental association? A unique case report,
919 Parasit. Vectors. 7 (2014) 207.

920

921 [66] T.V. Prodeus, O.P. Zelia, T.A. Khlebnikova, D.A. Pikul', [Extraenteric
922 infection caused by *Blastocystis* spp. in a female patient with liver
923 abscess (in Russian)], Med. Parazitol. (Mosk.) Apr-Jun (2014) 6-9.

924

925 [67] W.D. Patino, D. Cavuoti, S.K. Banerjee, K. Swartz, R. Ashfaq, T.
926 Gokaslan, Cytologic diagnosis of *Blastocystis hominis* in peritoneal
927 fluid: a case report, Acta Cytol. 52 (2008) 718-720.

928

929 [68] R. Silard, M. Petrovici, D. Panaitescu, V. Stoicescu, *Blastocystis*
930 *hominis* in the liver of *Cricetus auratus*, Arch. Roum. Pathol. Exp.
931 Microbiol. 36 (1977) 55-60.

932

933 [69] S.M. Collins, A role for the gut microbiota in IBS, Nat. Rev.
934 Gastroenterol. Hepatol. 11 (2014) 497-505.

935

936 [70] P.J. Kennedy, J.F. Cryan, T.G. Dinan, G. Clarke, Irritable bowel
937 syndrome: A microbiome-gut-brain axis disorder?, World J.
938 Gastroenterol. 20 (2014)14105-14125.

939

940 [71] J.K. Beatty, A. Bhargava, A.G. Buret, Post-infectious irritable bowel
941 syndrome: mechanistic insights into chronic disturbances following
942 enteric infection, World J. Gastroenterol. 20 (2014) 3976-3985

943

944 [72] P. Poirier, I. Wawrzyniak, C.P. Vivarès, F. Delbac, H. El Alaoui, New
945 insights into *Blastocystis* spp.: a potential link with irritable bowel
946 syndrome, PLoS Pathog. 8 (2012) e1002545.

- 947 [73] L.R. Krogsgaard, A.L. Engsbro, C.R. Stensvold, H.V. Nielsen, P.
948 Bytzer, The prevalence of intestinal parasites is not greater among
949 individuals with Irritable Bowel Syndrome: a population-based case-
950 control study, *Clin Gastroenterol Hepatol.* 13 (2014) 507-513.
- 951 [74] A.M. Petersen, H.V. Nielsen, C.R. Stensvold, J.H. Engberg, A. Friis-
952 Møller, I. Nordgaard-Lassen, S. Wildt, K.A. Krogh, *Blastocystis* and
953 *Dientamoeba fragilis* in active and inactive Inflammatory Bowel
954 Disease, *Gastroenterology* 140 (2011) S329-S330.
- 955 [75] A.M. Petersen, C.R. Stensvold, H. Mirsepasi, J. Engberg, A. Friis-
956 Møller, L.J. Porsbo, A.M. Hammerum, I. Nordgaard-Lassen, H.V.
957 Nielsen, K.A. Krogh, Active ulcerative colitis associated with low
958 prevalence of *Blastocystis* and *Dientamoeba fragilis* infection, *Scand.*
959 *J. Gastroenterol.* 48 (2013) 638-639.
- 960 [76] A.D. Kostic, R.J. Xavier, D. Gevers, The microbiome in inflammatory
961 bowel disease: current status and the future ahead, *Gastroenterology.*
962 146 (2014) 1489-1499.
- 963 [77] M. Arumugam, J. Raes, E. Pelletier, D. Le Paslier, T. Yamada, D.R.
964 Mende, G.R. Fernandes, J. Tap, T. Bruls, J.M. Batto, M. Bertalan, N.
965 Borruel, F. Casellas, L. Fernandez, L. Gautier, T. Hansen, M. Hattori,
966 T. Hayashi, M. Kleerebezem, K. Kurokawa, M. Leclerc, F. Levenez, C.
967 Manichanh, H.B. Nielsen, T. Nielsen, N. Pons, J. Poulain, J. Qin, T.
968 Sicheritz-Ponten, S. Tims, D. Torrents, E. Ugarte, E.G. Zoetendal, J.
969 Wang, F. Guarner, O. Pedersen, W.M. de Vos, S. Brunak, J. Doré, M.

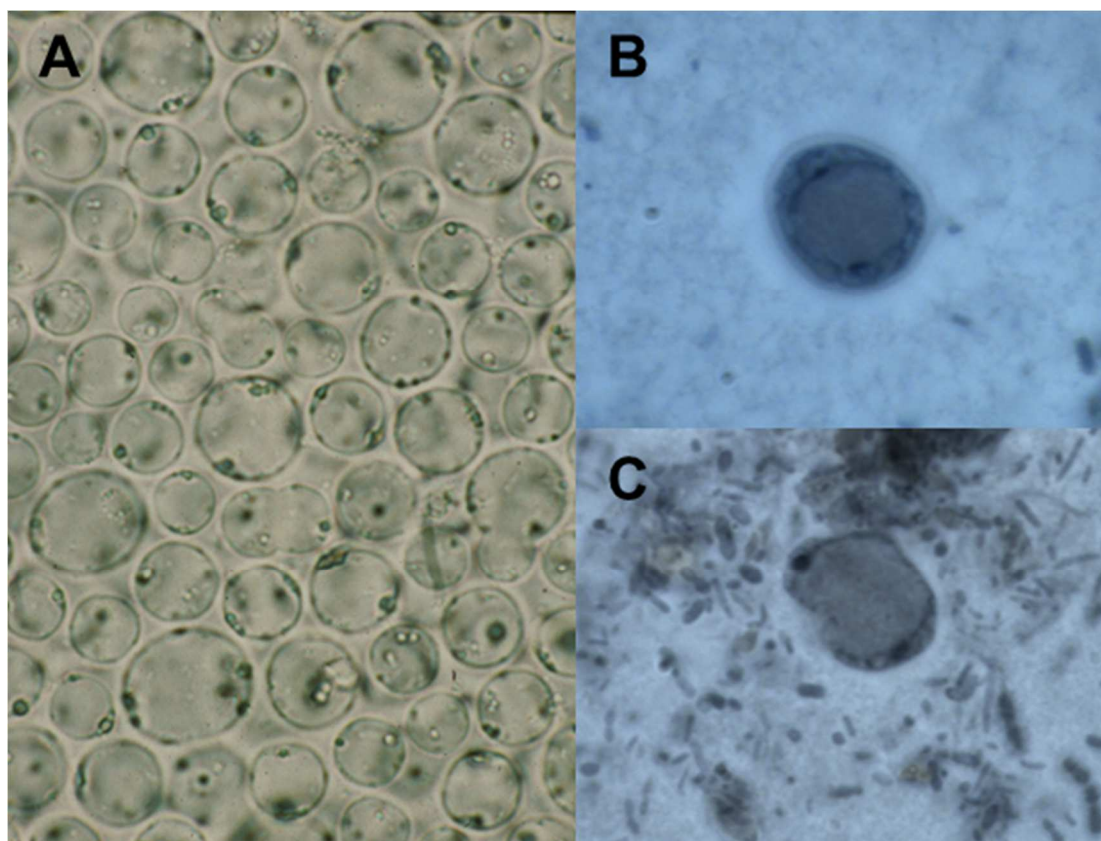
- 970 Antolín, F. Artiguenave, H.M. Blottiere, M. Almeida, C. Brechot, C.
971 Cara, C. Chervaux, A. Cultrone, C. Delorme, G. Denariáz, R. Dervyn,
972 K.U. Foerstner, C. Friss, M. van de Guchte, E. Guedon, F. Haimet, W.
973 Huber, J. van Hylckama-Vlieg, A. Jamet, C. Juste, G. Kaci, J. Knol, O.
974 Lakhdari, S. Layec, K. Le Roux, E. Maguin, A. Mérieux, R. Melo
975 Minardi, C. M'rini, J. Muller, R. Oozeer, J. Parkhill, P. Renault, M.
976 Rescigno, N. Sanchez, S. Sunagawa, A. Torrejon, K. Turner, G.
977 Vandemeulebrouck, E. Varela, Y. Winogradsky, G. Zeller, MetaHIT
978 Consortium, J. Weissenbach, S.D. Ehrlich, P. Bork, Enterotypes of the
979 human gut microbiome, *Nature* 473 (2011) 174-180.
- 980 [78] L.O. Andersen, I. Bonde, H.B. Nielsen, C.R. Stensvold, A retrospective
981 metagenomics approach to studying *Blastocystis*, *FEMS Microbiol.*
982 *Ecol.* 91 (2015) fiv072.
- 983 [79] A.H. Cekin, Y. Cekin, Y. Adakan, E. Tasdemir, F.G. Koclar, B.O.
984 Yolcular, Blastocystosis in patients with gastrointestinal symptoms: a
985 case-control study, *BMC Gastroenterol.* 12 (2012) 122.
- 986 [80] L.W. Parfrey, W.A. Walters, C.L. Lauber, J.C. Clemente, D. Berg-
987 Lyons, C. Teiling, C. Kodira, M. Mohiuddin, J. Brunelle, M. Driscoll, N.
988 Fierer, J.A. Gilbert, R. Knight, Communities of microbial eukaryotes in
989 the mammalian gut within the context of environmental eukaryotic
990 diversity, *Front. Microbiol.* 5 (2014) 298.
- 991

- 992 [81] C. De Filippo, D. Cavalieri, M. Di Paola, M. Ramazzotti, J.B. Poullet, S.
993 Massart, S. Collini, G. Pieraccini, P. Lionetti, Impact of diet in shaping
994 gut microbiota revealed by a comparative study in children from Europe
995 and rural Africa, Proc. Natl. Acad. Sci. USA. 107 (2010) 14691-14696.
- 996 [82] C.R. Stensvold, D.B. Christiansen, K.E. Olsen, H.V. Nielsen,
997 *Blastocystis* sp. subtype 4 is common in Danish *Blastocystis*-positive
998 patients presenting with acute diarrhea, Am. J. Trop. Med. Hyg. 84
999 (2011) 883-885.
- 1000 [83] M.V. Domínguez-Márquez, R. Guna, C. Muñoz, M.T. Gómez-Muñoz,
1001 R. Borrás, High prevalence of subtype 4 among isolates of *Blastocystis*
1002 *hominis* from symptomatic patients of a health district of Valencia
1003 (Spain), Parasitol. Res. 105 (2009) 949-955.
- 1004 [84] K. Heyland, M. Friedt, P. Buehr, C.P. Braegge, No advantage for
1005 antibiotic treatment over placebo in *Blastocystis hominis*-positive
1006 children with recurrent abdominal pain, J. Pediatr. Gastroenterol. Nutr.
1007 54 (2012) 677-679.
- 1008 [85] B. Speich, H. Marti, S.M. Ame, S.M. Ali, I.I. Bogoch, J. Utzinger, M.
1009 Albonico, J. Keiser, Prevalence of intestinal protozoa infection among
1010 school-aged children on Pemba Island, Tanzania, and effect of single-
1011 dose albendazole, nitazoxanide and albendazole-nitazoxanide, Parasit.
1012 Vectors 6 (2013) 3.

- 1013 [86] J.F. Rossignol, S.M. Kabil, M. Said, H. Samir, A.M. Younis, Effect of
1014 nitazoxanide in persistent diarrhea and enteritis associated with
1015 *Blastocystis hominis*, Clin. Gastroenterol. Hepatol. 3: (2005) 987-991.
- 1016 [87] L. Nigro, L. Larocca, L. Massarelli, I. Patamia, S Minniti, F. Palermo, B.
1017 Cacopardo, A placebo-controlled treatment trial of *Blastocystis hominis*
1018 infection with metronidazole, J. Travel Med. 10 (2003) 128-130.
- 1019 [88] C.R. Stensvold, H.V. Smith, R. Nagel, K.E. Olsen, R.J. Traub,
1020 Eradication of *Blastocystis* carriage with antimicrobials: reality or
1021 delusion?, J. Clin. Gastroenterol. 44 (2010) 85-90.
- 1022 [89] T. Roberts, J. Ellis, J. Harkness, D. Marriott, D. Stark, Treatment failure
1023 in patients with chronic *Blastocystis* infection, J. Med. Microbiol. 63
1024 (2014) 252-257.
- 1025 [90] R. Nagel, H. Bielefeldt-Ohmann, R. Traub, Clinical pilot study: efficacy
1026 of triple antibiotic therapy in *Blastocystis* positive irritable bowel
1027 syndrome patients, Gut Pathog. 6 (2014) 34.
- 1028 [91] I. Wawrzyniak, D. Courtine, M. Osman, C. Hubans-Pierlot, A. Cian, C.
1029 Nourrisson, M. Chabe, P. Poirier, A. Bart, V. Polonais, P. Delgado-
1030 Viscogliosi, H. El Alaoui, A. Belkorchia, T. van Gool, K.S.W. Tan, S.
1031 Ferreira, E. Viscogliosi, F. Delbac, Draft genome sequence of the
1032 intestinal parasite *Blastocystis* subtype 4-isolate WR1, Genomics Data
1033 4 (2015) 22-23.

- 1034 [92] V. Pérez-Brocal, C.G. Clark, Analysis of two genomes from the
1035 mitochondrion-like organelle of the intestinal parasite *Blastocystis*:
1036 complete sequences, gene content and genome organization, Mol.
1037 Biol. Evol. 25 (2008) 2475-2482.
- 1038 [93] I. Wawrzyniak, M. Roussel, M. Diogon, A. Couloux, C. Texier, K.S.W.
1039 Tan, C.P. Vivarès, F. Delbac, P. Wincker, H. El Alaoui, Complete
1040 circular DNA in the mitochondria-like organelles of *Blastocystis*
1041 *hominis*, Int. J. Parasitol. 38 (2008) 1377-1382.
- 1042 [94] V. Klimeš, E. Gentekaki, A.J. Roger, M. Eliáš, A large number of
1043 nuclear genes in the human parasite *Blastocystis* require mRNA
1044 polyadenylation to create functional termination codons, Genome Biol.
1045 Evol. 6 (2014) 1956-1961.
- 1046

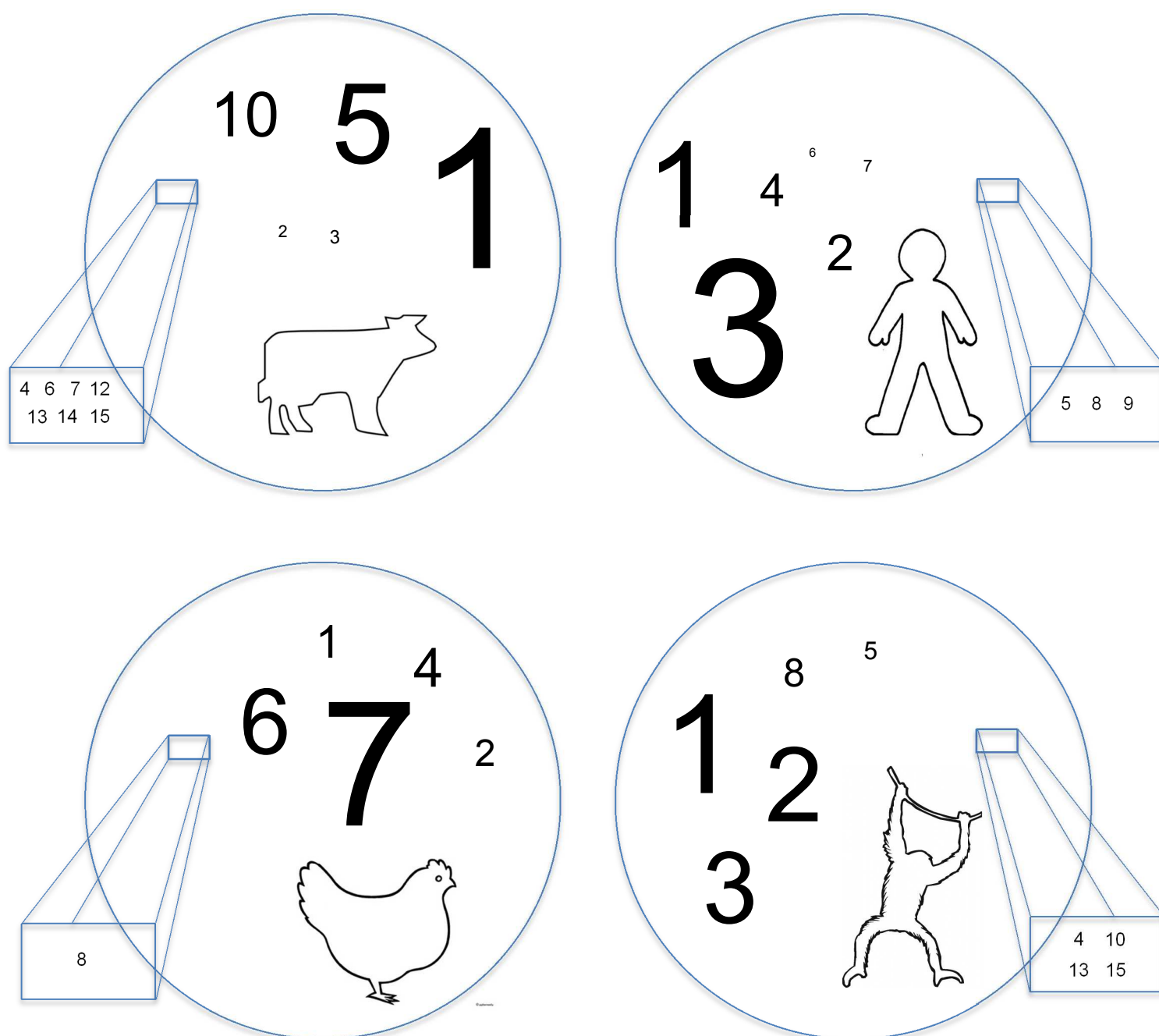
1047 Figure 1. Light microscopy images of *Blastocystis*. A. *Blastocystis* in culture.
1048 Using Robinson's and other media [29], *Blastocystis* often reaches high
1049 density in xenic culture. This stage is typically reported as 'vacuolar' due to
1050 the large central region of uncertain function. Organelles are seen as 'dots'
1051 along the periphery of the cell. B and C. *Blastocystis* in fecal smears, stained
1052 using iron-hematoxylin. Prominent nuclei are seen in the periphery of the cells
1053 as the most conspicuous morphological hallmark, along with the large central
1054 'void'. Other organelles can be discerned as smaller peripheral 'dots', which
1055 will include the mitochondrion-like organelles, etc. However, these can only be
1056 positively identified by transmission electron microscopy. Images courtesy of
1057 John Williams (A) and Claire Rogers (B, C), Diagnostic Parasitology
1058 Laboratory, London School of Hygiene and Tropical Medicine.
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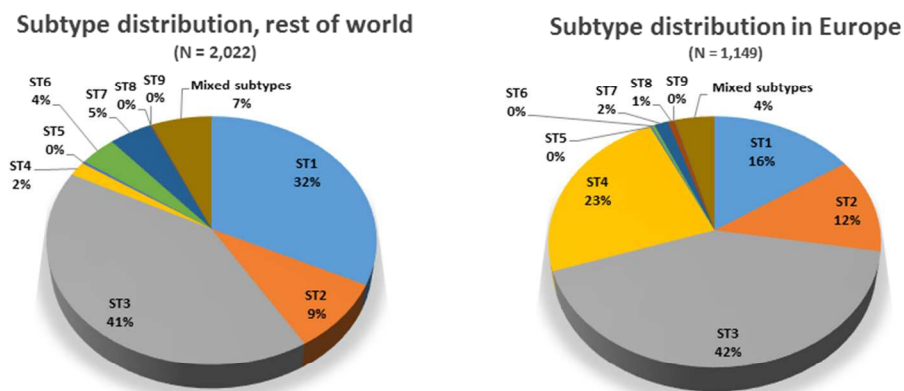
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1061 Figure 2. Host range and relative prevalence of *Blastocystis* subtypes. In this
 1062 schematic, the range of subtypes reported for four major host groups
 1063 (humans, non-human primates, ungulates and birds) is shown. In the circle,
 1064 the numbers are those of the most common subtypes found in the respective
 1065 host, with the integer font size proportional to its prevalence. Numbers in the
 1066 magnified boxes represent those subtypes that each constitute less than 5%
 1067 of the total samples subtyped to date. Derived from the numbers presented in
 1068 reference [10]. As an indication, prevalence figures for STs 1-4 in humans are
 1069 28.0%, 10.9%, 44.4% and 10.0% respectively.

1070



1071 Figure 3: Pie charts of human *Blastocystis* subtype distributions in Europe (A)
1072 and the rest of the world (B). These were produced from the data presented in
1073 Alfellani et al. [12]. Of note is the fact that although ST4 accounted for 10% of
1074 the samples across the world ($N = 318$), 87% of these (278) were from
1075 Europe, suggesting that ST4 is more or less geographically restricted to
1076 Europe.



1077