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1	Subtype distribution of Blastocystis isolates from synanthropic and zoo
2	animals and identification of a new subtype \bigstar
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4	C. Rune Stensvold ^{a,*} , Mohammed A. Alfellani ^b , Sara Nørskov-Lauritsen ^a , Katrine Prip ^a ,
5	Emma L. Victory ^b , Charlotte Maddox ^c , Henrik V. Nielsen ^a , C. Graham Clark ^b
6	
7	^a Department of Bacteriology, Mycology and Parasitology, Statens Serum Institut,
8	Artillerivej 5, DK-2300 Copenhagen S, Denmark
9	^b Department of Infectious and Tropical Diseases, London School of Hygiene and
10	Tropical Medicine, Keppel Street, London WC1E 7HT, United Kingdom
11	^c Department for Veterinary Diagnostics and Research, National Veterinary Institute, Technical
12	University of Denmark, Bülowsvej 27, Copenhagen V, Denmark
13	
14	*Corresponding author. Tel.: +45 32 68 36 04; fax: +45 32 68 30 33.
15	E-mail address: <u>RUN@ssi.dk</u>
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18	*Nucleotide sequence data reported in this paper are available in Genbank under the
19	accession numbers: <u>FM164412</u> and <u>FM164413.</u>
20	

21 Abstract

22	Blastocystis isolates from 56 Danish synanthropic and zoo animals, 62 primates
23	primarily from United Kingdom (UK) collections, and 16 UK primate handlers were
24	subtyped by PCR, sequencing and phylogenetic analysis. A new subtype (ST) from
25	primates and artiodactyls was identified and designated as Blastocystis sp. ST 10. STs
26	isolated from non-human primates ($n = 70$) included ST3 (33%), ST8 (21%), ST2 (16%),
27	ST5 (13%), ST1 (10%), ST4 (4%) and ST10 (3%). A high prevalence of ST8 was seen
28	among primate handlers (25%). This ST is normally very rare in humans, suggesting that
29	acquisition of Blastocystis ST8 infections from primates by their handlers had occurred in
30	these cases. Data from published studies of non-human primates, other mammals and
31	birds were collected and interpreted to generate a comprehensive overview on the ST
32	distribution in such animals. On the basis of information on 438 samples, it was found
33	that <i>Blastocystis</i> from primates belong mainly to ST1, ST2, ST3, ST5 and ST8, ungulates
34	and dogs mainly ST1, ST2, ST3, ST5 and ST10, rodents ST4, and birds mainly ST6 and
35	ST7. The data indicate moderate host specificity, most clearly exemplified by the fact that
36	STs isolated from avian and non-avian hosts rarely overlap.

Keywords: Blastocystis; PCR; Subtypes; Phylogeny; Epidemiology

40 **1. Introduction**

41	Blastocystis is a common single-celled parasite of humans, non-human primates,
42	other mammals, birds, amphibians, reptiles, fish, arthropods and annelids (Stenzel and
43	Boreham, 1996; König and Müller, 1997; Belova and Krylov, 1998; Yoshikawa et al.,
44	2004b, 2007). The parasite exhibits extensive genetic diversity, and on the basis of
45	molecular analysis of the ssrRNA gene, nine distinct subtypes (ST1-ST9) have been
46	identified from humans, non-human primates, other mammals and birds (Noël et al.,
47	2005; Stensvold et al., 2007). Blastocystis from non-human sources also comprise
48	isolates that appear to fall outside the genetic range of these nine subtypes, eg. reptilian,
49	amphibian and cockroach isolates (Yoshikawa et al., 2004b, 2007; Stensvold et al.,
50	2007), although they are clearly closely related.
51	It has been suggested by many authors that some human infections may result
52	from zoonotic transmission of the parasite, but at present this remains unproven. Humans
53	most frequently host ST3 but are also regularly found to carry ST1, ST2 and ST4 (Özyurt
54	et al. 2008). The five other STs (ST5-9) have been isolated only sporadically from
55	humans. Except for this information on humans, little is known about the potential host
56	specificity of Blastocystis. Such knowledge is necessary for epidemiological studies
57	aimed at identifying routes of transmission and zoonotic significance, which in turn are
58	important for strategies to control the spread and to increase our understanding of the
59	clinical impact of the parasite.
60	In recent years, molecular studies have produced a growing body of data on STs

In recent years, molecular studies have produced a growing body of data on STs
of *Blastocystis* isolated from various non-human hosts. The aim of the present study was
to identify STs of *Blastocystis* in synanthropic and zoo animals and to generate

63	hypotheses regarding the distribution and degree of host specificity of STs among non-
64	human Blastocystis.
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67	2. Materials and methods
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69	2.1. Danish samples: origin of isolates and PCR
70	DNA was extracted from faecal samples from a variety of synanthropic and zoo
71	animals at the National Veterinary Institute, Technical University of Denmark (Table 1).
72	All samples were from animals positive for Giardia and/or Cryptosporidium. None of the
73	samples was examined by in vitro culture for the specific detection of <i>Blastocystis</i> . Faecal
74	DNA extraction was performed using the QIAamp DNA Stool Mini Kit (QIAGEN,
75	Hilden, Germany) according to the manufacturer's recommendations.
76	Samples were screened for Blastocystis by PCR at the Statens Serum Institut as
77	previously described (Stensvold et al., 2006) using the primers bl1400ForC and
78	bl1710RevC, which amplify a 310 bp ssrRNA gene fragment, and Extract-N-Amp PCR
79	ReadyMix (Sigma-Aldrich Danmark, Brøndby, Denmark). PCR-positive samples were
80	sequenced as described previously (Stensvold et al., 2006). In cases where sequences
81	indicated the presence of a potential new ST, the primers RD5 and BhRDr (Scicluna et
82	al., 2006) were employed in order to obtain additional ssrRNA gene sequence
83	information for inclusion in phylogenetic analyses.

84	Two nucleotide sequences amplified by the two different primer sets (Scicluna et
85	al., 2006; Stensvold et al., 2006) representing a novel subtype obtained from a Danish
86	cow (RL056) were submitted to GenBank (accession nos. FM164412 and FM164413).
87	
88	2.2. United Kingdom samples: origin of isolates and PCR
89	Non-human primate and monkey handler material was received by the Diagnostic
90	Parasitology Laboratory of the London School of Hygiene and Tropical Medicine from
91	animal facilities and collections for routine parasitological investigation (Table 1). Faecal
92	samples were cultured in Robinson's medium (Clark and Diamond, 2002). Blastocystis
93	was harvested from positive cultures and stored in lysis buffer before discontinuing the
94	cultures. Culture lysate DNA was extracted using either a CTAB-based method (Ali et
95	al., 2005) or, more recently, the Gentra Puregene Cell kit (QIAGEN Ltd., Crawley, UK).
96	DNA samples were amplified using primers RD5, BhRDr and BioTaq polymerase
97	(Bioline Ltd., London, UK). PCR products were gel purified and sequenced using the
98	BhRDr primer as previously described (Scicluna et al., 2006).
99	
100	2.3. Phylogenetic analysis of isolates
101	Nucleotide sequences were aligned with a selection of 29 previously sequenced
102	Blastocystis ssrRNA genes, representing all nine established STs from mammals and
103	birds, and phylogenetic analysis was performed using Bayesian (MrBayes), Maximum
104	Likelihood and Neighbour-joining (PHYLIP) methods as described previously (Scicluna

105 et al., 2006). Pair-wise genetic distances within ST10 and between ST10 and other STs

106 were generated from the 'uncorrected "p" distance matrix' calculated using PAUP*

107	v.4.0b10 (Swofford, D.L., 2000. PAUP*. Phylogenetic analysis using parsimony (*and
108	other methods). Version 4. Sinauer Associates. Sunderland, Massachusetts, USA).
109	
110	2.4. Data collection, interpretation and terminology
111	The references used in the data collection process are listed in Table 2. Different
112	research groups have used distinct molecular methods and terminologies for analysing
113	isolates genetically, hence complicating comparison of results. Original data generated
114	from various studies using different molecular methodologies were standardised to meet
115	the proposed consensus terminology using a recently described algorithm (Stensvold et
116	al., 2007). Not all animal isolates described in the literature have been sequenced and
117	some were characterised only by PCR-restriction fragment length polymorphism (RFLP)
118	or PCR using sequence-tagged-site primers (PCR-STS). However, sequence data and
119	supplementary information published by Abe (2004) enabled interpretation of PCR-RFLP
120	data from some previous studies (Abe et al., 2003a, 2003b, 2003c) and PCR-STS data
121	can also be linked to most of the currently recognised STs. Where some STs were not
122	known at the time of publication, subsequent sequence analyses of such isolates and cross
123	referencing using data from Arisue et al. (2003), Noël et al. (2005) and Stensvold et al.
124	(2007) made it possible to identify STs originally described as 'ND' (not determined).
125	Blastocystis has also been reported in reptiles, amphibians, arthropods and annelids, but
126	only a small number of isolates have been sequenced. These were not included in the
127	present study as they appear to represent distinct lineages and are unlikely to represent a
128	zoonotic infection risk for humans.
129	

3. Results

132	PCR-positive samples from the present study included material from 16 primate
133	handlers, 70 non-human primates, 20 pigs, 25 cattle, two sheep, one deer and one dog
134	(Table 1). The ST distribution of isolates from primate handlers and animals identified in
135	the study is displayed in Table 1. Sequences obtained from isolates from 22 cattle, two
136	lemurs, one deer and one sheep showed relatively low similarity to existing STs when
137	percent identities were examined. These sequences formed a distinct group that clustered
138	together as a separate lineage emerging at the base of the ST4+ST8 clade (Fig. 1) when
139	the 310 bp region was used in phylogenetic analysis; this lineage was interpreted as a
140	novel ST and is here designated as ST10 (Table 1). When the longer sequences obtained
141	using the primers described by Scicluna et al. (2006) were used, ST10 emerged as a sister
142	group to ST8 (Fig. 2). The maximum likelihood bootstrap support for both of these
143	potential relationships was low and the affinities of ST10 must remain unresolved at
144	present. This ST has hitherto not been reported from human infections. Table 3 shows the
145	pair-wise genetic distances within ST10 and between ST10 and other STs.
146	Table 2 displays the STs of <i>Blastocystis</i> infecting 438 animals, including the data
147	from analysis of over 100 isolates in the present study and identifiable ST data from all
148	previously published studies. For comparison, Table 2 also includes the distribution of
149	STs isolated from humans based on 16 major studies (Alfellani, unpublished data).
150	It can be seen from Table 2 that ST3 is more common in humans than all other
151	STs combined. However, ST1, ST2 and ST4 also occur fairly frequently, whereas ST5
152	through ST9 occur only sporadically. Hence, the ST distribution among primate handlers

153 included in the present study was atypical: as expected, ST3 was the predominant ST, 154 seen in 9/16 (56%) of the monkey handlers, but ST8 was the next most common subtype, 155 being seen in four individuals (25%). Two of seven handler samples described previously 156 were also ST8 (Scicluna et al., 2006). Although the numbers are small, since ST8 is very rare in other humans but common in non-human primates, particularly woolly monkeys, 157 158 and given that the handlers would regularly come into contact with primate faeces in the 159 course of their work, the most likely explanation for this observation is that the handlers 160 acquired the ST8 infections from their charges. 161 To date, ST4 is the only ST to be isolated from rodents, but the total number of 162 samples (seven) and host species (two) studied is small. This apparent ST restriction in 163 rodents should be viewed with caution until larger studies have been performed. In 164 contrast, ST6 and ST7 predominate in birds, where more samples (35) and host species (eight) have been studied, giving a much stronger indication that a link exists between 165 these STs and avian hosts. 166 167

168

169 **4. Discussion**

We believe this is the first study to publish data on *Blastocystis* ST occurrences in
non-human hosts in Scandinavia and provides new data from molecular characterisation
of 119 animals and 16 primate handlers, adding substantially to the knowledge of *Blastocystis* host specificity. We present a comprehensive and systematic overview of the *Blastocystis* ST distribution in non-human hosts as it is known at the present time. Such

data are essential for an understanding of the host specificity and epidemiology of distinct *Blastocystis* sp. STs.

177 A novel ST (ST10) was isolated from both primates and ungulates in Denmark. 178 The reason why ST10 has not been identified previously could be due to a geographically 179 restricted distribution. However, given the high frequency of isolation in the present 180 study and the fact that it was isolated from different types of primates and other 181 mammals, it is more likely that some of the primers hitherto employed for *Blastocystis* 182 ST characterisation are unsuitable for the detection of this particular ST. For instance, the 183 R1 primer developed by Böhm-Gloning et al. (1997), which has been used in several 184 studies, anneals to a region of the ssrRNA gene that exhibits sequence variation, and 185 recently it was shown that this primer might preferentially amplify some STs over others 186 (Wong et al., 2008). Indeed, the ssrRNA gene of *Blastocystis* is relatively poorly 187 conserved, causing difficulties in designing sensitive genus-specific primers. In the study 188 by Thathaisong and colleagues (2003), 186 (mainly human) isolates were positive by 189 culture but were negative by PCR, and the isolates may have represented STs that were 190 not amplifiable by the R1 primer. Moreover, many studies (including this one) have used 191 in vitro culture for screening with subsequent extraction of DNA from the cultured 192 isolates. It is not known, however, whether all STs grow equally well in culture; recently, 193 data obtained by Parkar et al. (2007) suggested preferential in vitro amplification of ST2 194 over ST1. It is possible that ST10 does not grow under culture conditions commonly used 195 and this is why it has not been identified previously.

196 Since mixed ST infections (MSI) are quite common in humans (Stensvold et al.,197 unpublished data), it is possible that other animals also host MSI, which are not readily

198	identified by conventional PCR and sequencing. Indeed cultures from several primate
199	samples examined in the UK appeared from the sequence traces to be MSI and were
200	excluded from further analysis. It is suggested that, where possible, DNA should be
201	extracted directly from faeces and that ST-specific primers be developed for PCR
202	analysis to complement the genus-specific primers already in use. Sequencing of genus-
203	specific products will generate data regarding the extent of MSI in non-human hosts. The
204	ST-specific primers might not enable the detection of novel STs, but in combination with
205	the genus-specific data they should identify samples worthy of further investigation.
206	ST8 has been isolated from humans only rarely (Scicluna et al., 2006; Motazedian
207	et al., 2008; Stensvold et al., 2008) but is common in primate handlers, suggesting that
208	zoonotic spread from primates to primate handlers is responsible for the unexpectedly
209	high prevalence of this ST among these individuals. Zoonotic transmission of
210	Blastocystis has been suggested by a plethora of research groups (Snowden et al., 2000;
211	Abe et al., 2003c; Arisue et al., 2003; Thathaisong et al., 2003; Yoshikawa et al., 2003,
212	2004a; Abe, 2004; Noël et al., 2005; Parkar et al., 2007; Yan et al., 2007; Navarro et al.,
213	2008), yet the extent and nature of this phenomenon remains unclear as the published
214	evidence is equivocal. Given the ubiquity and the host range of <i>Blastocystis</i> , our ability to
215	assess the zoonotic potential of <i>Blastocystis</i> is dependent on our ability to i) correctly and
216	unambiguously identify STs, ii) detect and differentiate MSI, and iii) understand and
217	analyse possible factors involved in transmission such as transmission sources,
218	transmission vehicles, infectivity of cysts and other stages, contact with faeces or faecally
219	contaminated soil, water and food, coprophagy, and the possibility of animals shedding
220	ingested cysts that are simply passing through the host. In the future, identifying variable

molecular markers (eg. mini- or microsatellites) that differentiate strains within STs will
likely prove necessary in positively identifying links between potential animal sources
and specific human infections.

224 Comparing the data in Table 2 with the summary of the data from humans it 225 appears clear that birds usually host ST6 and ST7, but that these are rarely found in 226 mammals, having only been isolated from humans occasionally (Yan et al., 2007; 227 Alfellani, unpublished data; Stensvold, unpublished data). Interestingly, ST9 has so far 228 only been isolated from humans and on very few occasions. ST9 clusters with 'avian' 229 ST6 and ST7, so it is possible that birds are also the normal hosts of this ST. Given their 230 apparent host specificity, it is highly likely that human infections due to such avian STs 231 are of zoonotic origin as was previously suggested by Noël et al. (2005).

The situation in pigs is unclear. Studies seem to fall into two groups – those that find predominantly ST1 (Thathaisong et al., 2003; Navarro et al., 2008) and those that find predominantly ST5 (Abe et al., 2003c; Yoshikawa et al., 2004a; Yan et al., 2007; present study). There appears to be no geographic component to this difference, which at present remains a mystery.

To date, ST4 is the only ST found among rodents and marsupials. This ST has only infrequently been isolated from non-human primates and has not so far been isolated from other mammals; however, in humans ST4 represents approximately 5% of the isolates characterised to date (Table 2). It remains to be established whether contact with rodents poses a risk of transmission to humans of this particular subtype. The high prevalence of ST1-ST3 in humans and other mammals means that differentiating human origins from zoonotic origins of such human infections is not possible at present.

244	In conclusion, moderate host specificity seems to prevail among Blastocystis STs,
245	and the present data corroborate trends from other studies suggesting possible zoonotic
246	transmission of Blastocystis, at least of some STs. Future studies should aim to develop
247	high resolution molecular markers for analysing isolates in order to further elucidate the
248	zoonotic potential of the parasite.
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250	
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259	ACCEPT

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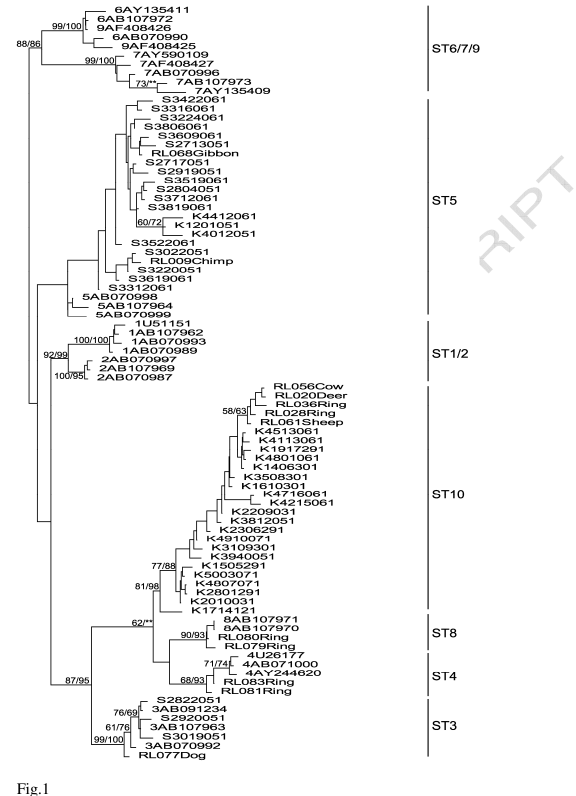
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387	585-594.
388	
	N N N N N N N N N N N N N N N N N N N

389 Figure legends

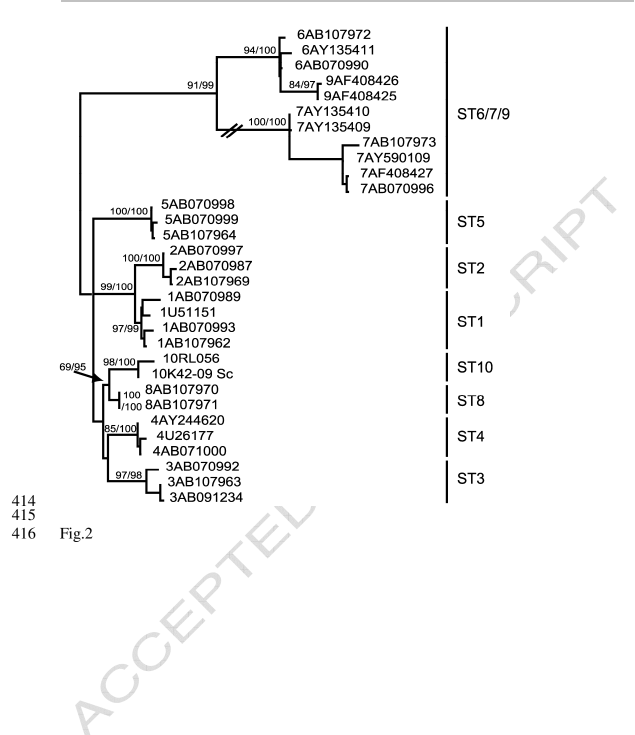
390

391	Fig. 1. Phylogenetic tree of the Danish Blastocystis sample sequences. The analysis was
392	performed using the 310 bp sequences, which are identified by their sample code. Those
393	starting with S are from pigs, those with K from cattle and most starting with R are from
394	other animals, with the species identity appended (Ring = Ring-tailed lemur). Reference
395	sequences from GenBank have the accession number preceded by the subtype
396	identification. The clade consisting of subtypes 6, 7 and 9 was used as an outgroup. The
397	tree shown is that obtained from the Bayesian analysis with the bootstrap proportions
398	shown being from the Maximum Likelihood analysis (100 replicates) on the left and
399	Neighbor-Joining analysis (1,000 replicates) on the right. The posterior probabilities
400	obtained in the Bayesian analysis were all 1.0. Bootstrap values of less than 50% in both
401	analyses are not shown. Where one analysis gave a value over 50% and the other below
402	50% the latter is indicated by two asterisks.
403	
404	Fig. 2. Phylogenetic relationships of <i>Blastocystis</i> subtype 10. The analysis was performed
405	using the 'barcode' region (Scicluna et al., 2006). Both samples sequenced (RL056 and
406	K42-09) are cattle. Reference sequences from GenBank have the accession number
407	preceded by the subtype identification. Subtypes 6, 7 and 9 were used as an outgroup.
408	The tree shown is that obtained from the Bayesian analysis with the bootstrap proportions
409	labelled as in Fig. 1. Bootstrap values of less than 50% are not shown. The branch leading

410 to ST7 has been shortened for convenience.







Host	Host	Country					Subt	ype (S	T)			
(common	(Latin name)	of	ST	ST	ST3	ST4	ST5	ST6	ST7	ST8	ST9	ST10
name)		Isolation	1	2					$\hat{\boldsymbol{\mathcal{O}}}$, ,		
Humans								2				_
(primate							\mathbf{C}					
handlers)	Homo sapiens	UK	2	-	9	1	<u>)</u> -	-	-	4	-	
)						
Non-human				5								
primates			6	r								
	Pan	UK	7									-
Chimpanzee	troglodytes	\sim	1	4	8	-	6	-	-	-	-	
		Denmark	-	-	-	-	1	-	-	-	-	-
	Pongo	UK										-
Orang Utan	pygmaeus		1	1	2	-	-	-	-	-	-	
	Gorilla	UK										-
Gorilla	gorilla		-	4	1	-	1	-	-	-	-	
Y	Hylobates	UK										-
Siamang	syndactylus		3	-	-	-	-	-	-	1	-	
Mueller's	Hylobates	UK										-
gibbon	muelleri		_	1		_	_	_	_	_		

Golden		UK										-
cheeked	Hylobates											
gibbon	gabriellae		1	-	1	-	-	-	-	-	-	
Lar gibbon	Hylobates lar	UK	1	-		-	-	-	-	1	-	-
Gibbon		Denmark								P		-
(unspecified)	Hylobates sp.		-	-	-	-	1	- <	2	-	-	
		UK	-	-	1	-	-	À	-	-	-	-
Woolly	Lagothrix	UK					\bigcirc					-
monkey	lagotricha		-	1	4	1) -	-	-	10	-	
Diana	Cercopithecus	UK			\mathcal{A})						-
monkey	diana		-	-5	1	-	-	-	-	-	-	
Barbary	Macaca	UK	A	r	2							-
macaque	sylvanus	^	-	-	1	-	-	-	-	-	-	
Stump-tailed	Macaca	UK										-
macaque	speciosa		-	-	1	-	-	-	-	-	-	
Common	Callithrix	UK										-
marmoset	jacchus		-	-	1	-	-	-	-	-	-	
Ring-tailed		Denmark										2
lemur	Lemur catta		-	-	-	2	-	-	-	2	-	
Unidentified		UK	-	-	2	-	-	-	-	1	-	-

Other

animals

Pig	Sus scrofa	Denmark	-	-	3	-	17	-	-	-	-	-
	domestica											
Cattle	Bos taurus	Denmark	-	-	-	-	3	-	-	-	-	22
Sheep	Ovis aries	Denmark	-	-	-	-	-	-	-	-	-	1
		UK	-	-	1	-	-	-	-	-	-	-
Roe Deer	Capreolus	Denmark	-	-	-	-	-	- <	ŻÌ	-	-	1
	capreolus						_	2				
Dog	Canis lupus	Denmark	-	-	1	-	(-)	<u> </u>	-	-	-	-
	familiaris					C						
) ~						

421

.mals 422 ^aAll Danish isolates were from animals that were also positive for *Giardia* and/or

Table 2. 424

Table 2. Blastocystis subtype distribution identified in non-human primates, other mammals and birds (n = 438). 425 426

Host group					Bla	stocysti	s sp. su	btype (ST)		\sim	Reference
	ST1	ST2	ST3	ST4	ST5	ST6	ST7	ST8	ST9	ST10	ST unknown	
Chimpanzee	1	-	-	-	-	-	-	-	-	-		Abe et al. (2003b); Abe (200
	1	1	-	-	-	-	-	-	-	C	-	Yoshikawa et al. (2004a)
	1	4	8	-	7	-	-	-	-	5-	-	Present study
Gorilla	-	4	1	-	1	-	-	-		-	-	Present study
Orang Utan	1	-	-	-	-	-	-		-	-	-	Abe et al. (2003b); Abe (200
	1	-	-	-	-	-	<u></u>	<u> </u>	-	-	-	Parkar et al. (2007)
	1	-	-	-	-	-		-	-	-	-	Yoshikawa et al. (2004a)
	1	1	2	-	-	9	-	-	-	-	-	Present study
Gibbons	-	1	-	-		-	-	-	-	-	-	Abe et al. (2003b); Abe (200
	2	-	-		-	-	-	-	-	-	-	Parkar et al. (2007)
	-	1	-	-	-	-	-	-	-	-	-	Yoshikawa et al. (2004a)
	5	1	2	-	1	-	-	2	-	-	-	Present study
Baboon	2	Q	-	-	-	-	-	-	-	-	-	Parkar et al. (2007)
Mandrill/Drill	2		_	_	_	_	-	-	-	_	_	Abe et al. (2003b); Abe (2004

	-	-	-	-	-	-	-	-	-	-	1	Yoshikawa et al. (2004a)
Macaques	1	2	-	-	-	-	-	-	-	-	-	Abe et al. (2003b); Abe (2004)
	1	-	-	-	-	-	-	-	-	-		Parkar et al. (2007)
	-	-	1	-	-	-	-	-	-	-		Scicluna et al. (2006)
	-	2	-	-	-	-	-	-	-	-) -	Yoshikawa et al. (2004a)
	-	-	2	-	-	-	-	-	-	6	-	Present study
Vervet monkey	1	-	-	-	-	-	-	-)-	-	Abe et al. (2003b); Abe (2004)
	1	1	-	-	-	-	-	- <		-	-	Parkar et al. (2007)
De Brazza's monkey	1	-	-	-	-	-	-		_	-	-	Abe et al. (2003b); Abe (2004)
	-	-	-	-	-	-	A	-	-	-	1	Yoshikawa et al. (2004a)
Diana monkey	-	-	1	-	-	Ā		-	-	-	-	Present study
Leaf monkey	1	-	-	-		\mathcal{A}	-	-	-	-	-	Abe et al. (2003b); Abe (2004)
	-	-	-	-/		-	-	-	-	-	1	Yoshikawa et al. (2004a)
'Japanese monkey'	-	1	-	<u>ó</u> -``	-	-	-	-	-	-	-	Yoshikawa et al. (1998)
	-	1	-	-	-	-	-	-	-	-	-	Yoshikawa et al. (2003)
Woolly monkey	2	2	1	-	-	-	-	3	-	-	-	Scicluna et al. (2006)
	C		4	1	-	-	-	10	-	-	-	Present study
Common marmoset	N	-	1	-	-	-	-	-	-	-	-	Present study
	Y											

Lemurs	-	-	-	-	-	-	-	1	-	-	-	Abe et al. (2003b); Abe (2
	3	1	-	-	-	-	-	-	-	-	-	Parkar et al. (2007)
	-	-	-	2	-	-	-	2	-	2		Present study
Unidentified primate	-	1	5	-	-	-	-	1	-	-		Scicluna et al. (2006)
	-	-	2	-	-	-	-	1	-	-	O^{-}	Present study
Primates Total	29	25	30	3	9	-	-	20	-	2	3	
Pigs	3	-	1	-	8	-	-	-)-	-	Abe et al. (2003c)
	-	-	-	-	1	-	-	- <		-	-	Arisue et al. (2003)
	122	7	-	-	-	-	-		-	-	-	Navarro et al. (2008)
	1	-	-	-	-	-	-	<u> </u>	-	-	-	Noël et al. (2003)
	-	-	-	-	1	_	_	-	-	-	-	Scicluna et al. (2006)
	20	-	-	-			-	-	-	-	-	Thathaisong et al. (2003)
	-	-	-	-/	16	-	-	-	-	-	-	Yan et al. (2007)
	-	-	-	<u>)</u> -	1	-	-	-	-	-	-	Yoshikawa et al. (1998)
	-	-	-	- \	1	-	-	-	-	-	-	Yoshikawa et al. (2003)
	4	Ē	2	-	14	-	-	-	-	-	-	Yoshikawa et al. (2004a)
	F	Y	3	-	17	-	-	-	-	-	-	Present study

Cattle	1	-	2	-	7	-	-	-	-	-	-	Abe et al. (2003c)
	1	-	1	-	6	-	-	-	-	-	-	Yoshikawa et al. (2004a)
	-	-	-	-	3	-	-	-	-	22	-	Present study
Cattle Total	2	-	3	-	16	-	-	-	-	22	-	
Horse	1	-	-	-	-	-	-	-	-	-) -	Thathaisong et al. (2003)
Deer	-	-	-	-	-	-	-	-	-	1	-	Present study
Sheep	-	-	1	-	-	-	-	-	~	1	-	Present study
Dog	1	3	-	-	-	-	-	- <		-	-	Parkar et al. (2007)
	-	-	1	-	-	-	-	P	-	-	-	Present study
Horse/Deer/Sheep /Dog Total	2	3	2	-	-	-	-	-	-	2	-	
Rat	-	-	-	1	-	~	-	-	-	-	-	Noël et al. (2003)
	-	-	-	3	1	2	-	-	-	-	-	Noël et al. (2005)
	-	-	-	1		-	-	-	-	-	-	Yoshikawa et al. (1998)
Guinea pig	-	-	-)1	-	-	-	-	-	-	-	Leipe et al. (1996)
	-	-	-	1	-	-	-	-	-	-	-	Silberman et al. (1996)
Opossum	-	6	<u> </u>	1	-	-	-	-	-	-	-	Parkar et al. (2007)
	-			8						-	-	

Non-primate mammals Total	154	10	11	8	75	-	-	-	-	24	-	
Duck	-	-	-	-	-	-	1	-	-	-	- /	Noël et al. (2003)
Goose	-	-	-	-	-	-	1	-	-	-	-	Abe (2004)
Chicken	-	-	-	-	-	1	-	-	-	-		Arisue et al. (2003)
	-	-	-	-	-	-	1	-	-	- (_	Noël et al. (2003)
	-	1	-	-	-	1	-	-	-	6	-	Yoshikawa et al. (2003)
	2	-	-	-	-	-	1	-)-	-	Yoshikawa et al. (2004a)
Quail	-	-	-	-	-	-	1	- <	7	-	-	Arisue et al. (2003)
	-	-	-	-	-	1	-		-	-	-	Yoshikawa et al. (1998)
	-	-	-	-	-	-	1	-	-	-	-	Yoshikawa et al. (2003)
	-	-	-	-	-	4	4	-	-	-	-	Yoshikawa et al. (2004a)
Pheasant	-	-	-	-			-	1	-	-	2	Abe et al. (2003a)
	1	-	-	-/		-	1	-	-	-	5	Yoshikawa et al. (2004a)
Guineafowl	-	-	-	<u>)</u> -	<u> </u>	1	-	-	-	-	-	Abe et al. (2003a)
Partridge	-	-		-	-	-	1	-	-	-	-	Abe et al. (2003a)
Turkey	-	-	<u>_</u>	-	-	-	1	-	-	-	-	Hess et al. (2006)
	C	Y	-	-	-	1	-	-	-	-	-	Noël et al. (2003)
Birds Total	3	1	-	-	-	10	13	1	-	-	7	

Mammals and birds total	186	36	41	11	84	10	13	21	0	26	10	
Total all subtypes = 438												
Humans total	316	71	577	54	-	28	18	3	2	-	17	Alfellani (unpublished)
Total all subtypes =1,086												
								5				
							0	P				
						-	9					
				$\boldsymbol{\wedge}$								
				2	•							
		()										

- 428 Table 3.
- 429 Average pair-wise distances within ST10 and between ST10 and other subtypes based on
- 430 sequences of the ssrRNA gene region amplified by the primers of (A) Scicluna et al.
- 431 (2006) or (B) Stensvold et al. (2006).
- 432

-	Blastocystis sp. subtypes (ST)	(A) Pairwise distance (%)	(B) Pairwise distance (%)
-	ST10/ST10	2.4	0.4
	ST10/ST8	5.0	4.1
	ST10/ST4	6.6	4.4
	ST10/ST3	8.8	6.0
	ST10/ST1	9.1	8.2
	ST10/ST2	10.1	7.4
	ST10/ST5	10.2	6.7
	ST10/ST9	12.0	10.3
	ST10/ST6	12.0	10.5
_	ST10/ST7	14.8	11.9
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