Review

Novel *Entamoeba* findings in non-human primates

Hany M. Elsheikha,¹ Carl S. Regan,¹,² and C. Graham Clark ³,*

¹ Faculty of Medicine and Health Sciences, School of Veterinary Medicine and Science, University of Nottingham, Sutton Bonington Campus, Loughborough, LE12 5RD, UK

² Current address: Vets4pets Dover Whitfield, White Cliffs Retail Park, Whitfield, Dover, CT16 3PS, UK

³ Faculty of Infectious and Tropical Diseases, London School of Hygiene and Tropical Medicine, London, UK

*Correspondence: graham.clark@lshtm.ac.uk

(C. Graham Clark)

**Keywords:** *Entamoeba*, Evolution, Non-human primates, Phylogeny, Species delineation
Abstract

In addition to well-known human-infecting species, *Entamoeba* species not found in humans have been identified recently in non-human primates (NHPs). Importantly, it has become clear that the organism identified as *Entamoeba histolytica* in NHPs is usually a distinct species, *Entamoeba nuttalli*. Many DNA-based stool surveys use species-specific detection methods and so may miss the full range of *Entamoeba* species present. In addition, authors may be using the same species name to describe distinct organisms. These various shortcomings may not be obvious to readers. In this review, we clarify the relationships between *Entamoeba* species’ names based on morphological and molecular data, and highlight gaps in recently published data on *Entamoeba* species in wild NHPs resulting from the use of variable methodology.

Humans and NHPs are both primates, but how similar are their *Entamoeba* species?

Humans are primates, and therefore it would be logical to assume that the parasite fauna of humans and non-human primates (NHPs; see Glossary) is likely to be similar. However, this simplistic view ignores the huge range of life-styles, diets and ecological specialisations exhibited by NHPs, and the millions of years of independent evolution that separate us from even our closest NHP relatives, the great apes. Nevertheless, humans and NHPs do appear to have many parasites in common, at least when identified via microscopy. Over recent decades, molecular tools have allowed us to re-examine these similarities and to challenge the assumption that apparent morphological identity equates to species identity. This review discusses how molecular tools provide a clearer picture of the relationships between intestinal amoebae of the genus *Entamoeba* in humans and NHPs and where gaps in our understanding remain.
What is causing invasive amoebiasis in humans and NHPs?

The focus on *Entamoeba* is largely due to *Entamoeba histolytica* being a significant cause of morbidity and mortality in humans. Published estimates suggest this organism is responsible for millions of cases of disease and over 50,000 deaths in humans annually [1]. Although these numbers are extrapolated from a limited number of studies, *E. histolytica* is certainly responsible for a significant amount of disease in some locations. Captive NHPs occasionally die from a disease that is, superficially, indistinguishable from that caused by *E. histolytica* in humans (e.g. [2]). Several other *Entamoeba* species that resemble *E. histolytica* morphologically have been described in both humans and NHPs, making microscopic diagnosis problematic. Morphologically distinct, non-pathogenic species of *Entamoeba* also appear to be shared by humans and NHPs, further complicating diagnosis (see below).

The morphology era

The existence of species of *Entamoeba* in humans and NHPs that appear identical by microscopy has been known for over a century. At that time, organisms in new hosts were often given new species names, whether morphologically distinguishable or not. A major work by Dobell [3] concluded that all named intestinal species of *Entamoeba* in humans could be assigned to either *E. histolytica* or *Entamoeba coli*, but he equivocated about *Entamoeba* from NHPs on the grounds of insufficient data; he later concluded that intestinal *Entamoeba* species in NHPs were also *E. histolytica* and *E. coli* [4]. His two-species nomenclature stayed essentially intact for 35 years.
In the mid-1950s, Burrows [5] resurrected the name *Entamoeba hartmanni* for an organism that parasitologists were referring to as ‘small race *E. histolytica*’. Dobell [3] had viewed *E. hartmanni* as a synonym of *E. histolytica*; however, Burrows showed that the sizes of *E. histolytica* ‘large race’ and ‘small race’ cysts were not a continuum but had a clear bimodal distribution. This first ‘break’ with the Dobell nomenclature was quickly adopted, because parasitologists were already primed to accept it.

**The molecular era**

*Entamoeba hartmanni* was the last change to Dobell’s nomenclature scheme based on morphology alone. Additional changes followed but not for many years, as the changes were primarily dependent on **small subunit ribosomal RNA gene** (SSU-rDNA) analyses. Emile Brumpt [6] proposed the existence of *Entamoeba dispar*, a non-pathogenic species morphologically identical to *E. histolytica*. This proposal was rejected by most parasitologists at the time (see discussion following [7]) and the name *E. dispar* virtually disappeared from the literature. Suspicion that Brumpt had been correct followed on from studies based on both lectin agglutination [8] and **isoenzyme** patterns [9], in which two groups within *E. histolytica* were identified, only one of which was found in patients with invasive disease. Subsequently, studies (cited in [10]) using monoclonal antibodies, DNA hybridization, SSU-rDNA restriction fragment length polymorphism, and eventually DNA sequencing all identified the same two groups of strains, and this led to the formal redescription of *E. dispar* as a species distinct from *E. histolytica* [10].

Other SSU-rDNA-based changes to the nomenclature of human *Entamoeba* species include the reassignment of ‘*E. histolytica*-like’ amoebae to the species *Entamoeba moshkovskii* [11]...
and the recognition that uninucleate cysts occasionally seen in humans were not always immature *E. histolytica* but were in fact *Entamoeba polecki* [12]. Most recently, *Entamoeba bangladeshi* was described as a new human species [13]; if it were not for SSU-rDNA sequences this organism would have been identified as *E. moshkovskii* despite it being genetically quite distinct.

The nomenclature for *Entamoeba* species in NHPs has followed suit, for the most part. *Entamoeba hartmanni* is commonly found in NHPs. *Entamoeba dispers* is also widespread in NHPs. *Entamoeba chattoni* had long been accepted as a NHP-specific species of *Entamoeba* with uninucleate cysts. It was designated a subtype of *E. polecki* a few years ago [14], but this change of nomenclature for *E. chattoni* has not been universally accepted; this will be discussed further below.

Thus, for the most part, the NHP *Entamoeba* nomenclature changes simply mirrored those in humans without any investigations to evaluate whether they were in fact the same organisms. This was understandable initially because there was no reason to suspect there were differences and the investigative tools were not readily available to many researchers. However, now that molecular techniques are routine in most research laboratories and some diagnostic laboratories, investigations into the diversity and identity of *Entamoeba* in NHPs have become more common and are revealing some surprising and important findings.

The evidence for *Entamoeba* genetic diversity in NHPs is based almost exclusively on SSU-rDNA analyses. Analyses of other markers are rarely possible because most studies use DNA extracted directly from stool samples, but when available they show the same species relationships. SSU-rDNA is a multicopy gene, which makes it relatively easy to amplify from
stool samples. In addition, and in contrast to some eukaryotes, the *Entamoeba* SSU-rDNA is relatively fast evolving (as evidenced by long branches in phylogenetic trees) meaning that sufficient resolution is obtained to allow differentiation of *Entamoeba* taxa using this gene alone.

*Entamoeba nuttalli*

*Entamoeba histolytica* causes disease of two main types: 1. amoebic dysentery/colitis, resulting from trophozoite invasion of the colonic mucosa and leading to ulceration, bleeding and the production of loose stool with blood and mucus; 2. amoebic liver abscess, resulting from haematogenous spread of trophozoites from the colon via the portal system to the liver, where tissue lysis leads to formation of a sterile pus-filled abscess [15]. Both types of disease have been reported in NHPs, and there have been a number of reports over the years of spontaneous invasive disease occurring in captive NHPs. Histologically, the diseases in humans and NHPs appear identical, as do the amoebae under the microscope [e.g. 2]. *Entamoeba histolytica* of human origin has been shown experimentally to be capable of infecting NHPs, where it can cause indistinguishable pathology [e.g. 16]. The organism responsible was therefore presumed to be *E. histolytica* in all cases of disease in NHPs.

In the last 10 years, however, molecular studies have been performed on amoebae from cases of invasive amoebiasis occurring spontaneously in NHPs. The amoebae in NHPs are consistently distinguishable from *E. histolytica* using a variety of DNA and protein markers: isoenzymes, SSU-rDNA and short tandem-repeat-containing loci [17-20]. Although closely related to *E. histolytica* – indeed it has been called “*E. histolytica*-like variant” [19] and “*E. histolytica* NHP variant” [21] by some – this is clearly a distinct organism and the name *E. nuttalli* has been revived for this amoeba [17]. *Entamoeba nuttalli* was originally described by
Castellani [22] in the liver abscess of a toque macaque (Macaca sinica) in Sri Lanka and is one of the species considered synonymous with *E. histolytica* by Dobell [3, 23]. Although we cannot prove after 110 years that the amoeba observed by Castellani is the same as the one now being called *E. nuttalli*, this seems quite likely. A recent survey of wild toque macaques in Sri Lanka detected asymptomatic carriage of *E. nuttalli* in 18.5% of the 227 animals studied [24]. *Entamoeba histolytica* was not detected in the population. *Entamoeba nuttalli* has been found in a variety of other NHPs – guenon, baboon, colobus and chimpanzee – in addition to other species of both captive and wild macaques [19, 25, 26].

The host and geographic ranges of *E. nuttalli* seem to be quite large, but so far it seems to be found primarily in primates of the Old World. Invasive disease has been reported in captive spider monkeys [25], but whether it infects wild New World NHPs is unknown. Only one human infection with *E. nuttalli* has been reported to date, in a zookeeper [27]. This is despite analyses of human samples that would have revealed its presence if it had been there. Isoenzyme analysis, which was used widely for *Entamoeba* species differentiation in the 1980s and early 1990s [e.g. 28], would have distinguished *E. nuttalli* from *E. histolytica* [17, 19], but although many thousands of human samples were studied in order to differentiate *E. dispar* and *E. histolytica*, no evidence of what is now being called *E. nuttalli* was reported. A second human infection has apparently been identified in Iraq, based on a sequence in GenBank stated to be of human origin [55; GenBank accession number: KP233837].

Note that most DNA-based diagnostic tools cannot distinguish *E. nuttalli* from *E. histolytica*, unless combined with sequencing, and neither can some commercial antigen-based diagnostic kits and monoclonal antibodies [17]. Therefore, although it seems unlikely that significant numbers of humans will be found to be infected with *E. nuttalli*, such infections may occur
occasionally among those who have close contact with NHPs, and may go unrecognized depending on the diagnostic method used. Primer pairs specific for *E. nuttalli* do now exist [17, 25] so that positive identification of this species without sequencing is possible.

NHPs can be infected experimentally with *E. histolytica* cysts of human origin [23, 29], although no invasive disease has resulted from such experiments. Captive NHP infections involving *E. histolytica* have been confirmed by DNA sequencing [30]. Therefore, it cannot be ruled out that some natural *E. histolytica* infections will occur in wild NHPs – most likely among those that come into contact regularly with humans or human waste – although there are no sequence-confirmed infections to date [e.g. 56]. It is impossible retrospectively to know which organism was responsible for the invasive amoebiasis cases in NHPs reported in the literature. Indeed, it is not possible to be certain that the amoeba observed was responsible for the disease in some cases – the presence of an *Entamoeba* and dysentery in the same host does not necessarily imply cause and effect.

*Entamoeba polecki*

*Entamoeba polecki* produces cysts with one nucleus, as does *E. chattoni*. Sequencing of their SSU-rDNAs revealed them to be closely related organisms [31]. The former species is traditionally associated with pigs and the latter with NHPs. Despite sporadic reports of *E. polecki* infections in humans for many years [32], when uninucleated cysts were seen in humans it was generally assumed that they represented immature cysts of *E. histolytica* rather than of *E. polecki* or *E. chattoni*. Verweij et al. [12] studied human *Entamoeba* infections where only uninucleated cysts were seen and found four distinct SSU-rDNA sequences. Two of these sequences were essentially identical to those of *E. polecki* and *E. chattoni* isolated from a pig and a monkey, respectively, while the other two sequences were related but distinct. This meant
that there were four closely-related organisms with two names between them and that *E. polecki* and *E. chattoni* were not host-specific since all four organisms were found in humans. Verweij et al. proposed [12] that the four should be viewed as variants of the same organism and called ‘*E. polecki*-like’, as the name *E. polecki* has precedence. Later, Stensvold et al. [14] proposed that they should be considered subtypes and numbered ST1-ST4, with the former *E. polecki* becoming *E. polecki* ST1 and the former *E. chattoni* becoming *E. polecki* ST2. The rationale for this approach is that there is no host specificity and no known difference except for small amounts of sequence divergence. This subtype nomenclature has not been fully accepted. One of the two ‘unnamed’ subtypes was in the interim named *Entamoeba struthionis* [33] as it was isolated from an ostrich, but this subtype (ST3) has subsequently been found in pigs [34] as well as humans. The fourth subtype has never had a species name and for a long time was only known from humans, where it is the most common subtype. Recently, however, ST4 was found to be the only *E. polecki* subtype in wild Celebes crested macaques (*Macaca nigra*) [35], proving that *E. polecki* ST2 (*E. chattoni*) is not the only subtype found in NHPs. It is possible that *E. polecki* ST1 and ST3 will also eventually be identified in NHP hosts. In the absence of host-specificity, use of the ‘*E. polecki* subtype’ nomenclature seems appropriate.

*Entamoeba dispar, Entamoeba hartmanni and Entamoeba coli*

For the most part, these three species meet the original expectation that human and NHP *Entamoeba* species are the same. *Entamoeba dispar* is quite a homogeneous species and there is no indication to date that *E. dispar* from humans is in any way distinct from that in NHPs. Although *E. hartmanni* shows a greater degree of SSU-rDNA variation than *E. dispar*, there is no obvious clustering of sequences that reflects human or NHP origin [14, 36], suggesting it is a discrete species with moderate intraspecific variation.
The situation in *E. coli* is more complex and less clear-cut. *Entamoeba coli* samples from humans group into two clusters, which have been named ST1 and ST2 [14]; ST1 appears to be slightly more common than ST2 in humans. When NHP *E. coli* samples are examined, the same two STs are identified, with ST2 being slightly more common, although this is based on relatively few samples. ST2 was recently identified in wild mountain gorillas (*Gorilla beringei*) [36]. The degree of divergence between the SSU-rDNAs of the two subtypes is substantial and distinct species names could be justified. However, other than this sequence divergence, there are no known differences between the two subtypes to date. *Entamoeba coli* cysts can vary quite dramatically in size [37, 38]. Whether this size variation is a morphological reflection of the underlying sequence divergence remains to be established.

Another *Entamoeba* that has been detected in NHPs is *Entamoeba* RL7 [14]. No species name has been assigned to this organism – it is simply known by its *ribosomal lineage* (RL) number [14]. *Entamoeba* RL7 was originally identified in a sample from a Phayre’s leaf monkey (*Trachypithecus phayrei*) [14], but it has subsequently been detected in humans in West Africa [34]. Uniquely, this *Entamoeba* is most closely related to *Entamoeba muris* (Figure 1), which, like *E. coli*, produces cysts with eight nuclei. Based on morphology, this organism previously would have been reported as *E. coli*.

**NHP-restricted Entamoeba Species**

There are several NHP-restricted *Entamoeba* sequences worthy of discussion here. The first is *Entamoeba* RL3, which to date has only been detected in langurs of various species and produces cysts with a single nucleus. In the past it would likely have been reported as *E.*
chattoni based on microscopy. No infections with this organism have been reported in humans, or indeed in any other NHP. It is closely related to, but distinct from, Entamoeba bovis and related lineages that are confined to ungulates [14]. RL3 has only been found in a few samples but it is notable that two lineages of Entamoeba (RL3 and RL7) have to date been detected exclusively in langurs. Whether this is linked to their unusual foregut fermentative digestion is unclear.

Villanueva-García et al. [39] recently reported SSU-rDNA sequences of an apparently novel Entamoeba in two species of Howler monkey. Because these were only partial sequences they were given a conditional lineage identifier [34] rather than a RL number. Entamoeba CL8 is clearly distinct from previously sequenced Entamoeba SSU-rDNAs and, interestingly, the CL8 sequence branches within a cluster of sequences obtained from reptilian Entamoeba strains. Villanueva-García et al. found a second Entamoeba sequence in their samples that is virtually identical to Entamoeba RL6, which was originally described from the green iguana (Iguana iguana) [14, 40]. The complete SSU-rDNA sequence of both these organisms would be helpful in order to confirm their phylogenetic tree placement.

Finally, there has been one report of Entamoeba suis from a gorilla (Gorilla gorilla) [14], but whether this is a natural host for this Entamoeba species remains to be established. This species also produces cysts with a single nucleus.

**Missing Entamoeba Species?**

Perhaps surprisingly, there are to date no reports of E. moshkovskii from NHPs. This organism is actually a species complex with substantial intra-specific sequence variation [40] and is
being reported from humans with increasing frequency now that PCR-based detection is being employed [e.g. 41-43]. *Entamoeba moshkovskii* has also been detected in cattle, elephants [34], reptiles [44] and insects [Silberman JD, personal communication], so it is likely only a matter of time before it is also found in NHPs. Not all published molecular studies have tested for this species and in those that did it is not clear whether the primers used would detect all variants of this genetically diverse species complex. The most recently described *Entamoeba* of humans, *E. bangladeshi* [13], is also yet to be reported from NHPs.

*Entamoeba gingivalis*, which colonises the gingival pockets in the mouth of humans, is listed as having been found in NHPs [e.g. in 45]. No molecular data are available to know whether the organisms reported in NHPs differ from those in humans. This may be important, as there are at least two SSU-rDNA variants of *E. gingivalis* in humans [40] and additional diversity could exist in other hosts.

A summary of the relationships between species names and identifiers can be found in Table 1 and an outline phylogenetic tree depicting the relationships between *Entamoeba* SSU-rDNA sequences is depicted in Figure 1.

**Captive vs. Wild NHPs**

Data on the presence and prevalence of *Entamoeba* species in NHPs is patchy at best, and most reports are based on animals in zoological parks. This is a problem when it comes to interpreting the data. The first issue is how to interpret the presence of parasites in captive NHPs. Animals in captivity may be exposed to organisms they would never encounter in the wild. Therefore, the data only indicate that the NHP species is capable of becoming colonised
by the parasite identified, not that it is a natural host for this parasite. A second issue is the
impact of captivity on prevalence. It is likely that animals come in contact with faeces and
faecal contamination of food and water more frequently in captivity than they would in the
wild; this is especially true of species that are primarily or exclusively arboreal. Only by
studying wild NHPs can ‘natural’ infections be identified, although in the case of peri-urban
and urban NHPs the possibility of infection through contact with human faeces cannot be
excluded. It is, of course, also likely that wild NHPs will ingest faeces from other hosts,
accidentally or on purpose. If the ingested faeces contains *Entamoeba* cysts it is possible that
DNA of these organisms will be detected when the NHP faeces is screened by PCR. However,
unless the NHP species ingests faeces frequently and in significant amounts it would be
unlucky if the small amount of NHP faeces analysed contained detectable DNA of *Entamoeba*
cysts that were just passing through.

Relatively few studies of *Entamoeba* in wild NHPs have employed molecular diagnostics to
date, and microscopy does not differentiate most of the known *Entamoeba* species: only *E.
histolytica*, *E. coli* and *E. chattoni* are regularly reported in publications reliant on microscopy.
Each of these names actually represents a mixture of distinct organisms united only by the
number of nuclei in their mature cyst. *Entamoeba hartmanni* is the only additional species that
can be identified by morphology, but only if cyst diameters are measured; often this is either
not the case or the information is not given. As a result, only studies employing sequence-based
identification will be discussed below. We recognise that this excludes the vast majority of
studies, but if the data are not interpretable we feel they are better omitted.

Molecular studies in wild NHPs published to date (Table 2) are few in number, mostly involve
Old World NHPs, and vary in the methodology used. In some studies, species-specific PCR
has been used, but often not all known species were tested for despite primers being available, leaving gaps in the data (Table 2, notes). When species-specific PCR has been used, this often means subtypes were not identified and potentially interesting data on sequence variation and host range have been lost. Several studies did not test for *E. hartmanni*, leading to a false impression of the distribution of this *Entamoeba* species in NHPs. It is notable that *E. histolytica* was not detected in any of these studies.

The use of only species-specific primers can mean that novel *Entamoeba* species are missed. For example, if Villanueva-Garcia et al. [39] had used species-specific primers for *Entamoeba*, the two novel *Entamoeba* species found in Howler monkeys (CL8 and RL6) would not have been identified – the samples would have appeared negative even though *Entamoeba* organisms were present. Sequencing of products amplified using genus-specific primers may seem the best way forward, but there is a catch. NHPs are often carriers of multiple *Entamoeba* species and mixed PCR products give unreadable sequences with the standard DNA sequencing. The approach of Jirků-Pomajbíková et al. [46] could be a good compromise – genus-specific amplification coupled with nested species-specific PCR. This allows identification of species in mixed infections yet does not miss mono-infections with novel *Entamoeba* species, as these would be positive with genus-specific but negative with all the species-specific primers used. Jirků-Pomajbíková et al. [46] did not initially test for *E. hartmanni* but through sequencing discovered that it was the *Entamoeba* present in the samples positive with the genus-specific primers but negative with the species-specific primer pairs used. However, this method will only identify the presence of novel *Entamoeba* species if they are present as a single infection unless it is combined with cloning of the PCR products.
It seems likely that identification of *Entamoeba* in NHPs in the future will be through microbiome data, whether from targeted amplification and sequencing of a portion of eukaryotic SSU-rDNA or by extraction of such sequences from metagenomic data. Both approaches are in use in humans and have identified *Entamoeba* when present, but to date have rarely been applied to NHP samples. In one example, Wegener Parfrey et al. [47] identified *E. hartmanni* (among many other eukaryotes) in captive NHPs through eukaryote-targeted SSU-rDNA amplification and 454 sequencing. Similarly, random sequencing of faecal DNA has the potential to identify not only all the species present, but could enable assembly of partial or complete genomes for the organisms identified [e.g. 48]. While such approaches are expensive and likely to be available only to a few at present, the holistic information on the eukaryome of NHPs likely to be obtained by such approaches makes them very attractive and we look forward to seeing the data emerge in the next few years.

**Concluding Remarks**

Currently, at least six *Entamoeba* species with valid published names have been confirmed by molecular analysis in NHPs: *E. coli*, *E. polecki*, *E. histolytica*, *E. nuttalli*, *E. dispar* and *E. hartmanni*. However, in addition there are multiple subtypes within *E. coli* and *E. polecki*, plus organisms with no name but distinct gene sequences (*Entamoeba* RL3, RL6, RL7 and CL8). This remarkable expansion in known diversity has been driven largely by the use of molecular techniques that have facilitated the identification of many novel and previously unrecognised *Entamoeba* species in NHPs.

However, many points remain to be clarified (see “Outstanding Questions”). It is unclear whether *E. moshkovskii*, *E. bangladeshi* and *E. gingivalis* colonise NHPs as well as humans.
Novel sequences with no linked species name are likely to continue to be detected in NHPs around the world. This search for new types of *Entamoeba* in NHPs is essential as it remains to be proven whether only *E. nuttalli* is responsible for morbidity and mortality in these hosts. However, unless the correct approaches are used, such organisms will remain undiscovered.

We now know that NHPs are infected by both NHP-restricted and human-infective *Entamoeba* species. Morphological diagnosis of *Entamoeba* species will always be problematic, but most molecular approaches used to date may also be considerably underestimating the prevalence, diversity, and distribution of *Entamoeba* in NHPs. At the same time, insufficient taxon sampling and the heavy focus on humans may well have led us to inaccurate conclusions about *Entamoeba* evolution. Fortunately, interest in the eukaryotic microbiome is growing in parallel with improvements in technology, and it is likely that within the next few years a better understanding of the evolution and host ranges of *Entamoeba* in NHPs will emerge.

Metagenomic analyses could allow the use of genes other than SSU-rDNA for phylogenetic analyses. Obtaining sequence data for other genes is difficult - if not impossible - using traditional molecular approaches and DNA from faecal samples. Multigene phylogenies may well provide greater resolution that could confirm or refute our current views of relationships within *Entamoeba*. Greater resolution is essential for evaluating the relative importance of cospeciation and host-switching in the evolution of primate *Entamoeba* species. It seems likely that these data will start to become available in the near future.

A recent study showed a significant reduction in the gut microbiome diversity of captive NHPs, with a shift occurring from a wild NHP microbiome state toward a modern human microbiome state [49]. Whether alterations in the lifestyle and diet of captive NHPs or the disruption of
normal hierarchical social behavior [50] has led to this perturbation of their gut microbiome, the change may predispose captive NHPs to infection with certain Entamoeba spp, normally confined to humans. Comparison of gut microbiomes across NHPs living in the wild, semicaptivity and captivity using sequencing of both bacteria and Entamoeba SSU rDNA, is already possible. Such data will allow us to investigate the correlation between microbiota signatures and prevalence of specific Entamoeba species in NHPs.

There is much more to learn regarding both the microbiome and the eukaryome of NHPs, especially those in the wild. There has been a strong focus on Old World primates, in particular macaques, while New World primates are significantly underrepresented and prosimians have not been studied. It is hoped that the range of species sampled will broaden, otherwise we will continue to have a rather limited view of Entamoeba diversity in NHPs.

Acknowledgments

We would like to thank all the scientists whose research discoveries have advanced our knowledge of the epidemiology of Entamoeba in non-human primates and enabled us to write this review.


53. Debenham, J.J. et al. (2017) Occurrence of Giardia, Cryptosporidium, and Entamoeba in wild rhesus macaques (Macaca mulatta) living in urban and semi-rural North-West India. Int. J. Parasitol. Parasites Wildl. 6, 29–34


Glossary

Conditional lineage (CL): an Entamoeba identified as likely to be distinct based on sequencing of partial SSU-rDNA, but for which sufficient data are not yet available. See RL, below.

Isoenzymes: each of two or more sequence variants of an enzyme that exhibit different migration in electrophoresis gels due to charge differences.

Non-human primates (NHPs): all members of the order Primates other than humans; NHPs share many similarities with humans in terms of physiology, anatomy, immunology, and neurology, but are very diverse in their ecology, diet, etc. The split between humans and NHPs is an artificial one, as humans are much more closely related to some NHPs than others.

Ribosomal lineage (RL): an Entamoeba taxon identified as distinct by sequencing of its complete SSU-rDNA gene. Often no corresponding morphological data are available. In other groups of organisms these are often called operational taxonomic units (OTUs) but in this case, it is clear that they belong to the genus Entamoeba.

Small subunit ribosomal RNA gene (SSU-rDNA): the gene encoding the smaller of the two major RNA components of the ribosome, also known as 18S rDNA. This gene is the most widely used single locus for phylogenetic analyses in eukaryotes and bacteria. In Entamoeba, the gene size generally falls between 1800 and 2200 bases.

Subtype: a discrete genetic clade within a named species.
Figure 1: Phylogenetic relationships among *Entamoeba* species. The phylogenetic tree shown is modified from Figure 1 in Jacob et al. [34]. Names in bold lettering are those that have been identified by sequencing of SSU-rDNA in NHPs. Adjacent to the *Entamoeba* names are those of the NHP species (wild or captive) in which the *Entamoeba* has been identified.

![Phylogenetic Tree](image-url)
<table>
<thead>
<tr>
<th>Dobell nomenclature&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Current species names</th>
<th>Identified in primates (incl. humans)</th>
<th>Molecular identification in NHPs</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>E. histolytica</em></td>
<td><em>E. histolytica</em></td>
<td><em>E. histolytica</em></td>
<td>Y&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td><em>E. dispar</em></td>
<td><em>E. dispar</em></td>
<td></td>
<td>Y</td>
</tr>
<tr>
<td><em>E. hartmanni</em></td>
<td><em>E. hartmanni</em></td>
<td></td>
<td>Y</td>
</tr>
<tr>
<td><em>E. nuttalli</em></td>
<td><em>E. nuttalli</em></td>
<td></td>
<td>Y</td>
</tr>
<tr>
<td><em>E. moshkovskii</em></td>
<td><em>E. moshkovskii</em> (complex)</td>
<td></td>
<td>N</td>
</tr>
<tr>
<td><em>E. polecki</em></td>
<td><em>E. polecki</em> ST1&lt;sup&gt;c&lt;/sup&gt;</td>
<td></td>
<td>N</td>
</tr>
<tr>
<td></td>
<td><em>E. polecki</em> ST4</td>
<td></td>
<td>Y</td>
</tr>
<tr>
<td><em>E. chattoni</em></td>
<td><em>E. polecki</em> ST2</td>
<td></td>
<td>Y</td>
</tr>
<tr>
<td><em>E. struthionis</em></td>
<td><em>E. polecki</em> ST3</td>
<td></td>
<td>N</td>
</tr>
<tr>
<td><em>E. bangladeshi</em></td>
<td><em>E. bangladeshi</em></td>
<td></td>
<td>N</td>
</tr>
<tr>
<td><em>E. suis</em></td>
<td><em>E. suis</em></td>
<td></td>
<td>Y&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td><em>E. coli</em></td>
<td><em>E. coli</em></td>
<td><em>E. coli</em> ST1</td>
<td>Y&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td><em>E. coli</em> ST2</td>
<td></td>
<td>Y</td>
</tr>
<tr>
<td><em>E. gingivalis</em></td>
<td><em>E. gingivalis</em></td>
<td><em>E. gingivalis</em> ribodeme 1&lt;sup&gt;d&lt;/sup&gt;</td>
<td>N</td>
</tr>
<tr>
<td></td>
<td><em>E. gingivalis</em> ribodeme 2</td>
<td></td>
<td>N</td>
</tr>
<tr>
<td>None</td>
<td><em>Entamoeba</em> RL3&lt;sup&gt;e&lt;/sup&gt;</td>
<td></td>
<td>Y</td>
</tr>
<tr>
<td></td>
<td><em>Entamoeba</em> RL6</td>
<td></td>
<td>Y</td>
</tr>
<tr>
<td></td>
<td><em>Entamoeba</em> RL7</td>
<td></td>
<td>Y</td>
</tr>
<tr>
<td></td>
<td><em>Entamoeba</em> CL8&lt;sup&gt;f&lt;/sup&gt;</td>
<td></td>
<td>Y</td>
</tr>
</tbody>
</table>

**Table 1.** Correspondence between historic, binomial, and sequence-based nomenclature for *Entamoeba* species in primates.

<sup>a</sup>Dobell’s nomenclature is that proposed in his 1919 monograph [3].

<sup>b</sup>Identified in captive NHPs only, to date.
c Subtypes (ST) are distinct small-subunit ribosomal DNA sequence variants that clearly fall within a named species.
d Ribodemes are small-subunit ribosomal DNA variants detected by restriction enzymes.
e Ribosomal (RL) [14] lineages indicate complete small-subunit ribosomal DNA sequences that are clearly distinct from all named species.
f Conditional (CL) [34] lineages indicate partial small-subunit ribosomal DNA sequences that are clearly distinct from all named species.
<table>
<thead>
<tr>
<th>NHP species</th>
<th>Type of amplification</th>
<th>Total no. of samples</th>
<th>Species identified (no. of samples)</th>
<th>Reference</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rhesus macaques (<em>Macaca mulatta</em>)</td>
<td>Species-specific</td>
<td>715</td>
<td><em>E. nattali</em> (440), <em>E. dispar</em> (16), <em>E. coli</em> (574), <em>E. polecki ST2</em> (649)</td>
<td>51</td>
<td>a</td>
</tr>
<tr>
<td>Rhesus macaque (<em>Macaca mulatta</em>)</td>
<td>Species-specific</td>
<td>112</td>
<td><em>E. nattali</em> (57), <em>E. dispar</em> (13), <em>E. coli</em> (83), <em>E. polecki ST2</em> (96)</td>
<td>26</td>
<td>b</td>
</tr>
<tr>
<td>Tibetan macaque (<em>Macaca thibetana</em>)</td>
<td>Species-specific</td>
<td>89</td>
<td><em>E. nattali</em> (15), <em>E. coli</em> (37), <em>E. polecki ST2</em> (59)</td>
<td>52</td>
<td>c</td>
</tr>
<tr>
<td>Savannah woodland chimpanzee (<em>Pan troglodytes schweinfurthii</em>)</td>
<td>Genus- and species-specific</td>
<td>107</td>
<td><em>E. hartmanni</em> (32), <em>E. dispar</em> (10), <em>E. coli ST2</em> (33)</td>
<td>46</td>
<td>d</td>
</tr>
<tr>
<td>Celebes crested macaque (<em>Macaca nigra</em>)</td>
<td>Species/subtype-specific</td>
<td>77</td>
<td><em>E. polecki ST4</em> (75)</td>
<td>35</td>
<td>e</td>
</tr>
<tr>
<td>Toque macaque (<em>Macaca sinica</em>)</td>
<td>Species-specific</td>
<td>227</td>
<td><em>E. nattali</em> (42), <em>E. dispar</em> (1), <em>E. coli</em> (40), <em>E. polecki ST2</em> (197)</td>
<td>24</td>
<td>f</td>
</tr>
<tr>
<td>Rhesus macaque (<em>Macaca mulatta</em>)</td>
<td>Genus- and species-specific</td>
<td>128</td>
<td><em>E. coli</em> (63), unidentified <em>Entamoeba</em> (65)</td>
<td>53</td>
<td>g</td>
</tr>
<tr>
<td>Mountain gorilla (<em>Gorilla beringei beringei</em>)</td>
<td>Genus-specific</td>
<td>68</td>
<td><em>E. coli ST2</em> (4), <em>E. hartmanni</em> (33)</td>
<td>36</td>
<td>h</td>
</tr>
<tr>
<td>Howler monkeys (<em>Alouatta palliata</em> and <em>A. pigra</em>)</td>
<td>Genus-specific</td>
<td>155</td>
<td><em>Entamoeba CL8</em> (6 from <em>A. pigra</em>, 1 from <em>A. palliata</em>), <em>Entamoeba RL6</em> (1 from <em>A. pigra</em>)</td>
<td>39</td>
<td>i</td>
</tr>
</tbody>
</table>

**Table 2.** Summary of results from molecular screening of faecal samples from wild NHP populations*

* The publication by Dong et al. [54] includes data on several NHP species in China (mostly *Macaca mulatta* and *M. fascicularis*) but it is not possible to identify which results came from sampling wild populations. Samples were tested by species-specific amplification for *E. histolytica,*
E. nuttalli, E. dispar, E. moshkovskii, E. coli, and E. polecki ST2. Only E. coli and E. dispar were detected. No tests for E. hartmanni or other E. polecki subtypes were performed.

a: Authors also tested captive macaques; these are excluded from the table. Tested for E. histolytica, E. dispar, E. nuttalli, E. coli, and E. polecki ST2 only. No test for E. hartmanni, E. moshkovskii or other E. polecki subtypes.

b: Tested for E. histolytica, E. dispar, E. nuttalli, E. moshkovskii, E. coli, and E. polecki ST2 only. No test for E. hartmanni or other E. polecki subtypes.

c: Tested for E. histolytica, E. dispar, E. nuttalli, E. coli, and E. polecki ST2 only. No test for E. hartmanni, E. moshkovskii or other E. polecki subtypes.

d: Genus-PCR-positive samples were tested for E. histolytica, E. nuttalli, E. dispar, E. moshkovskii, E. coli, and E. polecki ST2. Genus-PCR positive, but species-specific PCR negative samples were sequenced and identified as E. hartmanni.

e: Tested for E. histolytica, E. dispar, E. nuttalli, E. moshkovskii, E. coli, and E. polecki ST1, ST2 and ST4. No test for E. hartmanni or E. polecki ST3.

f: Tested for E. histolytica, E. dispar, E. nuttalli, E. moshkovskii, E. coli, and E. polecki ST2. No test for E. hartmanni or other E. polecki subtypes.

g: Genus-PCR positive samples were tested for E. coli. Multiplex PCR for E. histolytica, E. dispar and E. moshkovskii on all samples. No test for E. nuttalli, E. polecki, or E. hartmanni.

h: Sequencing of Genus-PCR positive amplicons identified only these two species.

i: Sequencing of Genus-PCR positive amplicons identified only these two organisms.