

Running title: *Entamoeba histolytica* genome

Structure and Content of the *Entamoeba histolytica* Genome

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1 **ABSTRACT**

2 The intestinal parasite *Entamoeba histolytica* is one of the first protists for
3 which a draft genome sequence has been published. Although the genome is
4 still incomplete, it is unlikely that many genes are missing from the list of those
5 already identified. In this review we summarise the features of the genome as
6 they are currently understood and provide previously unpublished analyses of
7 many of the genes.

8

9 **1. INTRODUCTION**

10 *Entamoeba histolytica* is one of the most widespread and clinically important
11 parasites, causing both serious intestinal (amoebic colitis) and extraintestinal
12 (amoebic liver abscess) diseases throughout the world. A recent WHO estimate
13 (WHO, 1998) places *E. histolytica* second after *Plasmodium falciparum* as
14 causing the most deaths annually (70,000) among protistan parasites.

15

16 Recently a draft of the complete genome of *E. histolytica* was published (Loftus
17 *et al.*, 2005) making it one of the first protist genomes to be sequenced. The *E.*
18 *histolytica* genome project was initiated in 2000 with funding from the
19 Wellcome Trust and the National Institute of Allergy and Infectious Diseases to
20 the Wellcome Trust Sanger Institute and The Institute for Genomic Research
21 (TIGR) in the UK and the USA, respectively. The publication describing the
22 draft sequence concentrated on the expanded gene families, metabolism and the
23 role of horizontal gene transfer in the evolution of *E. histolytica*. In this review
24 we summarise the structure and content of the *E. histolytica* genome in
25 comparison to other sequenced parasitic eukaryotes, provide a description of
26 the current assembly and annotation, place the inferred gene content in the
27 context of what is known about the biology of the organism, and discuss plans
28 for completing the *E. histolytica* genome project and extending genome
29 sequencing to other species of *Entamoeba*.

30

1 The fact that the genome sequence is still a draft has several important
2 consequences. The first is that a few genes may be missing from the sequence
3 data we have at present, although the number is likely to be small. For example,
4 at least one gene (amoebapore B) is not present in the genome data despite it
5 having been cloned, sequenced and the protein extensively characterised well
6 before the start of the genome project. The second consequence is that the
7 assembly contains a number of large duplicated regions that may be assembly
8 artifacts, meaning that the number of gene copies is over-estimated in several
9 cases. These problems cannot as yet be resolved but should be eventually as
10 more data becomes available. Nevertheless, it is important to remember these
11 issues when reading the rest of this article.

12

13 As the number of genes in *E. histolytica* runs into several thousands it is not
14 possible to discuss all of them. However, we have generated a number of tables
15 that identify many genes and link them to their entries in GenBank using the
16 relevant protein identifier. Only a few tables are included in the text of this
17 review, but the others are available on line as supplementary material -
18 http://pathema.tigr.org/pathema/entamoeba_resources.shtml. The *E. histolytica*
19 genome project data are being 'curated' at TIGR and it is on that site that the
20 most current version of the assembled genome will be found. The 'Pathema'
21 database will hold the data and the annotation (<http://pathema.tigr.org/>). The
22 gene tables are also linked to the appropriate entry in the Pathema database and
23 the links will be maintained as the genome structure is refined over time.

24

25 Reference is made throughout the text to other species of *Entamoeba* where
26 data are available. *Entamoeba dispar* is the sister species to *E. histolytica* and
27 infects humans without causing symptoms. *Entamoeba invadens* is a reptilian
28 parasite that causes invasive disease, primarily in snakes and lizards, and is
29 widely used as a model for *E. histolytica* in the study of encystation although
30 the two species are not very closely related (Clark *et al.*, 2006b). Genome
31 projects for both these species are underway at TIGR and it is anticipated that

1 high quality draft sequences will be produced for both in the near future. It is
2 hoped that the *E. dispar* sequence will prove useful in identifying genomic
3 differences linked to disease causation while that of *E. invadens* will be used to
4 study patterns of gene expression during encystation. Small-scale genome
5 surveys have been performed for two other species: *E. moshkovskii*, which is
6 primarily a free-living species although it occasionally infects humans, and *E.*
7 *terrapinae*, a reptilian commensal species
8 (http://www.sanger.ac.uk/Projects/Comp_Entamoeba/).

9

10 **2. GENOME STRUCTURE**

11 **2.1 The *E. histolytica* Genome Sequencing, Assembly and** 12 **Annotation Process**

13 The first choice to be made in the genome project was perhaps the easiest - the
14 identity of the strain to be used for sequencing. A significant majority of the
15 existing sequence data prior to the genome project was derived from one strain:
16 HM-1:IMSS. This culture was established in 1967 from a rectal biopsy of a
17 Mexican man with amoebic dysentery and axenised shortly thereafter. It has
18 been used widely for virulence, immunology, cell biology and biochemistry in
19 addition to genetic studies. In an attempt to minimise the effects of long-term
20 culture cryopreserved cells that had been frozen in the early 1970s were revived
21 and this uncloned culture used to generate the DNA for sequencing.

22

23 Before undertaking a genome scale analysis it is important to understand the
24 quality and provenance of the underlying data. The *E. histolytica* genome was
25 sequenced by whole genome shotgun approach with each center generating
26 roughly half of the reads. Several different DNA libraries containing inserts of
27 different sizes were produced using DNA that had been randomly sheared and
28 sequences were obtained from both ends of each cloned fragment. The Phusion
29 assembler (Mullikin and Ning, 2003) was used to assemble the 450,000 short
30 reads into larger contigs (contiguous sequences), resulting in 1819 genome
31 fragments that were approximately 12X deep, which means that each base has

1 been sequenced 12 times, on average. While the genome shotgun sequence
2 provides high coverage of each base it is inevitable that there will be
3 misassemblies and sequencing errors in the final consensus particularly towards
4 each end of the contigs. Another problem with draft sequence is that it contains
5 gaps, and while most of these will be small and will mostly contain repetitive
6 non-coding “junk” sequence, some of the gaps will probably contain genes.
7 This makes it impossible to be absolutely certain of the absence of particular
8 genes in *E. histolytica* and, in some cases, the presence or absence of particular
9 biological pathways. Due to the high repeat content and low GC content
10 (24.1%) of the *E. histolytica* genome, closure of the remaining gaps is likely to
11 be a lengthy process. Therefore it was decided to undertake and publish an
12 analysis of the genome draft following assembly of the shotgun reads.

13

14 Annotation of the protein coding regions of the genome was initially carried out
15 using two genefinders (GlimmerHMM (Majoros *et al.*, 2004) and Phat (Cawley
16 *et al.*, 2001)) previously used successfully on another low G+C genome, that of
17 *P. falciparum*. The software was re-trained specifically for analysis of the *E.*
18 *histolytica* genome. The training process involved preparing a set of 600
19 manually edited genes to be used as models with the subsequent genefinding
20 then being carried out on all of the assembled contigs to generate a 'complete'
21 gene set. Predicted gene functions were generated automatically by homology
22 searches using public protein and protein-domain databases, with subsequent
23 refinement of identifications being carried out by manual inspection. For
24 particular genes and gene families of special interest, members of the
25 *Entamoeba* scientific community were involved throughout this process as
26 expert curators with each individual assisting in the analysis and annotation of
27 their genes of interest. Therefore although the manual curation of the genome
28 has not been systematic, those areas of biology that are of primary interest to
29 the *Entamoeba* community have been annotated most thoroughly. The
30 publication of the genome by Loftus *et al* therefore represents a “first draft” of
31 the complete genome sequence and the level of annotation is similar to the

1 initial publications of other genomes such as *Drosophila* (Adams *et al.*, 2000;
2 Myers *et al.*, 2000) and human (Lander *et al.*, 2001).

3

4 **2.2 Karyotype and Chromosome Structure.**

5 The current *E. histolytica* genome assembly is approximately 23.7 million
6 basepairs (Mbp) in size (Table 1). This figure is not likely to be a very accurate
7 measure. In part this is due to misassembly of repetitive regions, which will
8 cause the genome to appear smaller, and in part because of the possibility of
9 aneuploidy in some regions of the genome, which would cause them to appear
10 more than once in the assembly. Overall, however, this size is not inconsistent
11 with data from pulse-field gels (Willhoeft and Tannich, 1999) and kinetic
12 experiments (Gelderman *et al.*, 1971a,b) making the *E. histolytica* genome
13 comparable in size (24 Mbp) to that of *Plasmodium falciparum* (23 Mbp)
14 (Gardner *et al.*, 2002), *Trypanosoma brucei* (26 Mbp) (Berriman *et al.*, 2005),
15 and the free living amoeba *Dictyostelium discoideum* (34 Mbp) (Eichinger *et*
16 *al.*, 2005).

17

18 The current assembly does not represent complete chromosomes. Analysis of
19 pulse-field gels predicts 14 chromosomes ranging in sizes from 0.3 to 2.2 Mb
20 and possibly a ploidy of four (Willhoeft and Tannich, 1999). There is no current
21 information regarding the size and nature of the centromeres and there are no
22 contigs that appear to contain likely centromeric regions based on comparisons
23 with other organisms. A search for signature telomeric repeats within the data
24 indicates that these are either not present in the genome, not present in our
25 contigs, or are diverged enough to be unidentifiable. However, there is
26 circumstantial evidence that the chromosome ends may contain arrays of tRNA
27 genes (see 2.4 below).

28

29 **2.3 Ribosomal RNA Genes**

30 The organisation of the structural RNA genes in *E. histolytica* is unusual with
31 the rRNA genes carried exclusively on 24 kb circular episomes (Bhattacharya

1 *et al.*, 1998) that have two transcription units in an inverted repeat. These
2 episomes are believed to make up about 20% of the total cellular DNA; indeed,
3 roughly 15% of all of the sequencing reads generated in the genome project
4 were derived from this molecule with the exception of certain libraries where
5 attempts were made to exclude it. There are thought to be numerous other
6 circular DNA molecules of varying sizes present with unknown functions (Dhar
7 *et al.*, 1995; Lioutas *et al.*, 1995)but unfortunately they have not yet been
8 identified in the genome shotgun sequence data. The exact reasons for this are
9 unknown but the small size of the DNA may have prevented proper shearing
10 during the library construction process. These molecules represent an intriguing
11 unsolved aspect of the *E. histolytica* genome.

12

13 **2.4 tRNA Genes**

14 Perhaps the most unusual structural feature identified in the *E. histolytica*
15 genome is the unprecedented number and organisation of its tRNA genes (Clark
16 *et al.*, 2006a). Over 10% of the sequence reads contained tRNA genes and these
17 are (with a few exceptions) organised in linear arrays. The array organisation of
18 the tRNAs was immediately obvious in some cases from the presence of more
19 than one repeat unit in individual sequence reads and in other cases from their
20 presence in both reads from the two ends of the same clone. However because
21 of the near complete identity of the array units they were impossible to
22 assemble by the software used and therefore the size of the arrays cannot be
23 estimated accurately.

24

25 By manual assembly of tRNA gene-containing reads, 25 distinct arrays with
26 unit sizes ranging from under 500 bp to over 1750 bp were identified (Clark *et*
27 *al.*, 2006a). The arrayed genes are predicted to be functional because of the 42
28 acceptor types found in arrays none has been found elsewhere in the genome.
29 These array units encoded between one and five tRNAs and a few tRNA genes
30 are found in more than one unit. Three arrays also encode the 5S RNA and one
31 encodes what is thought to be a small nuclear RNA. Experimental quantitative

1 hybridisations suggest a copy number of between about 70 and 250 for various
2 array units. In total it is estimated that there are about 4500 tRNA genes in the
3 genome. The frequency of a particular tRNA isoacceptor appears to be
4 independent of the codon usage in *E. histolytica* protein-coding genes.

5

6 Between the genes in the array units are complex, non-coding, short tandem
7 repeats ranging in size from 5 to over 36 bp. Some variation in short tandem
8 repeat number is observed between copies of the same array unit but this
9 variation is usually minor and not visible when inter-tRNA PCR amplification
10 is performed. However, these regions often exhibit substantial variation when
11 different isolates of *E. histolytica* are compared and this is the basis of a
12 recently described genotyping method for this organism (Ali *et al.*, 2005).

13

14 There is indirect evidence to suggest that the tRNA arrays are present at the
15 ends of chromosomes. Although allelic *E. histolytica* chromosomes often differ
16 substantially in size in pulse-field gels, a central protein-encoding region
17 appears to be conserved as DNA digested with rare cutting enzymes gives only
18 a single band in Southern blots when most protein-coding genes are used as
19 probes. In contrast, when some tRNA arrays are used as probes on such blots,
20 the same number of bands is seen in digested and undigested DNA. It is
21 therefore tempting to conclude that the tRNA genes are at the ends of the
22 chromosomes and to speculate that these repeat units may perform a structural
23 role. In *D. discoideum* it is thought that rDNA may function as a telomere in
24 some cases (Eichinger *et al.*, 2005) and the tRNA arrays in *E. histolytica* may
25 perform a similar role.

26

27 The chromosomal regions flanking the tRNA arrays are generally devoid of
28 protein coding genes but often contain incomplete transposable elements (see
29 next section) and other repetitive sequences (Clark *et al.*, 2006a). This is also
30 consistent with a telomeric location.

31

1 2.5 LINEs

2 The *E. histolytica* genome is littered with transposable elements. There are two
3 major types autonomous LINEs (Long Interspersed Elements) of which there
4 are three subtypes (EhLINE 1, 2 and 3) and there are two types of SINEs (Short
5 Interspersed Elements) (Eh SINE1 and 2) (Table 2a). The classification of these
6 elements and their organisation has been reviewed recently (Bakre *et al.*, 2005).
7 Phylogenetic analysis of the EhLINEs places them in the R4 clade of non- Long
8 Terminal Repeat (LTR) elements, a mixed clade of elements that includes
9 members from nematodes, insects, and vertebrates (Van Dellen *et al.*, 2002a).
10 Analysis of the *E. histolytica* genome shows no evidence for the presence of
11 LTR retrotransposons and very few DNA transposons (of the *Mutator* family)
12 (Pritham *et al.*, 2005).

13

14 All copies of EhLINEs examined encode non-conservative amino acid changes,
15 frame shifts, and/or stop codons and no copy with a continuous open reading
16 frame (ORF) has yet been found. This suggests that the majority of these
17 elements are inactive. However, a large number of EhLINE1 copies do contain
18 long ORFs without mutations in the conserved protein motifs of the RT and EN
19 domains, suggesting that inactivity is quite recent. ESTs corresponding to
20 EhLINEs have been found suggesting that transcription of these elements still
21 occurs. Although most R4 elements insert in a site-specific manner, EhLINEs
22 do not show strict site-specificity and are widely dispersed in the genome. They
23 are quite frequently found close to protein-coding genes and inserted near T-
24 rich stretches (Bakre *et al.*, 2005).

25

26 All three EhLINE subtypes are of approximately equal size ranging from 4715
27 to 4811 bp in length. Individual members within an EhLINE family typically
28 share >85% identity, while between families they are <60% identical. By
29 aligning the available sequences, each EhLINE can be interpreted to encode a
30 single predicted ORF that spans almost the entire element (EhLINE1, 1589 aa;
31 EhLINE2, 1567 aa; EhLINE3, 1587 aa). However, a precise 5bp duplication at

1 nucleotide position 1442 in about 80% of the copies of EhLINE1 creates a stop
2 codon, dividing the single ORF in two. Similarly in 92% of EhLINE2 copies,
3 the single ORF contains a precise deletion of two nucleotides at position 1272,
4 resulting in two ORFs. Very few intact copies of EhLINE3 are found. The
5 location of the stop codon leading to two ORFs appears to be conserved since
6 in both EhLINE1 and EhLINE2 the size of ORF1 is about half that of ORF2
7 (Bakre *et al.*, 2005). Among the identifiable domains in the predicted proteins
8 are reverse transcriptase (RT) and a restriction enzyme-like endonuclease (EN).
9 The putative 5' and 3' untranslated regions are very short (3-44 bp).

10

11 EhLINEs 1 and 2 appear to be capable of mobilising partner SINEs (see next
12 section) for which abundant transcripts have been detected in *E. histolytica*.
13 Putative LINE/SINE partners can be assigned on the basis of conserved
14 sequences at the 3' -ends of certain pairs, which otherwise showed no sequence
15 similarity. The relevance of this assignment for the EhLINE1/SINE1 pair has
16 recently been demonstrated (Mandal *et al.*, 2004).

17

18 **2.6 SINEs**

19 The two EhSINEs are clearly related to the EhLINEs as they have a conserved
20 3' sequence. They are nonautonomous, non-LTR retrotransposons
21 (nonautonomous SINEs). The genetic elements encoding the abundant
22 polyadenylated but untranslatable transcripts found in *E. histolytica* cDNA
23 libraries (initially designated IE elements (Cruz-Reyes and Ackers, 1992;
24 Cruz-Reyes *et al.*, 1995) or *ehapt2* (Willhoeft *et al.*, 2002)) have now been
25 designated EhSINE1 (Van Dellen *et al.*, 2002a; Willhoeft *et al.*, 2002). BLAST
26 searching with representative examples of the first 44 EhSINE1s detected has
27 identified 90 full-length (= 99% complete) copies and at least a further 120
28 partial (= 50% of full length) copies in the genome. Length variation is
29 observed among EhSINE1s and is largely due to variable numbers of internal
30 26-27 bp repeats (Ackers, unpublished). The majority contain two internal
31 repeats and cluster closely around 546 bp in length.

1
2 A second *E. histolytica* SINE (EhSINE2) has recently been described (Van
3 Dellen *et al.*, 2002a; Willhoeft *et al.*, 2002). Examination of the four published
4 sequences again suggests the presence of variable numbers of short (20 bp)
5 imperfect repeats. BLAST searching identified a total of 47 full-length (= 99%)
6 and at least 60 partial copies in the genome. The 3'-end of EhSINE2 shows high
7 similarity (76%) to the 3' end of EhLINE2.

8
9 A polyadenylated transcript designated UEE1 found commonly in cDNA
10 libraries from *E. dispar* (Sharma *et al.*, 1999) is also a non-LTR
11 retrotransposon. A single copy of a UEE1-like element has been identified in
12 the *E. histolytica* genome and is here designated EhSINE3. There is no
13 significant sequence identity between EhSINE3 and EhLINE3 but the 3' end of
14 EhSINE3 is very similar to that of EhLINE1.

15
16 Analysis of an *E. histolytica* EST library identified over 500 significant hits to
17 both EhSINE1 and EhSINE2. No convincing transcript from EhSINE3 could
18 be identified although the nearly identical *E. dispar* UEE elements (EdSINE1;
19 Shire and Ackers, submitted) are abundantly transcribed.

20
21 A very abundant polyadenylated transcript, *ehapt1*, was described by Willhoeft
22 *et al.* (1999) in a cDNA library. However, only a small number of partial
23 matches could be found in the current *E. histolytica* assembly and only 10-20
24 strong hits in the much larger *E. histolytica* EST library now available. *ehapt1*
25 does not appear to be a SINE element and its nature is currently unclear. The
26 lack of matches in the genome suggests either that it is encoded in regions
27 missing from the current assembly or that it contains numerous introns.

28

29 **2.7 Other Repeats**

30 The *E. histolytica* genome contains a number of other repetitive elements
31 whose functions are not always clear. There are over 75 genes encoding

1 leucine-rich tandem repeats (LRR) of the type found in BspA-like proteins of
2 the *Treponema pallidum* LRR (TpLRR) subfamily, which has a consensus
3 sequence of LxxIxIxxVxxIgxxAFxxCxx (Davis *et al.*, 2006). These proteins
4 generally have a surface location and may be involved in cell-cell interaction.
5 Genes encoding such proteins are found mainly in Bacteria and some Archaea;
6 so far they have been identified in only one other eukaryote, *Trichomonas*
7 *vaginalis* (Hirt *et al.*, 2002). An extensive description of the BspA-like proteins
8 of *E. histolytica* has recently been published (Davis *et al.*, 2006) and one of
9 them has been shown to be surface exposed (Davis *et al.*, 2006).

10

11 *E. histolytica* stress sensitive protein (Ehssp) 1 is a dispersed, polymorphic and
12 multicopy gene family (Satish *et al.*, 2003) and is present in ca. 300 copies per
13 haploid genome as determined by hybridisation (Table 2a). The average Ehssp1
14 ORF is 1 kb in length with a centrally-located acidic-basic region (ABR) that is
15 highly polymorphic. Unlike other such domains no clear repetitive motifs are
16 present. The protein has, on average, 21% acidic (aspartate and glutamate) and
17 17% basic (arginine and lysine) amino acids, most of which are located in the
18 ABR. The ABR varies in size from 5 to 104 amino acids among the various
19 copies. No size polymorphism is seen outside the central ABR domain. The
20 genes have an unusually long 5' untranslated region (UTR; 280 nucleotides).
21 Only one or a few copies of the gene are transcribed during normal growth, but
22 many are turned on under stress conditions. Homologues of this gene are
23 present in *E. dispar*, but there is very little size polymorphism in the *E. dispar*
24 gene family.

25

26 Eukaryotic genomes usually contain numerous microsatellite loci with repeat
27 sizes of 2-3 basepairs. With the exception of di- and tri-nucleotides made up
28 entirely of A+T such sequences are rare in the *E. histolytica* genome. In
29 contrast, two dispersed repeated sequences of unknown function occur far more
30 frequently than would be expected at random. Family 16 has a 42 base
31 consensus sequence and occurs approximately 38 times in the genome while

1 family 17 has a 27 base consensus sequence and occurs 35 time in the genome
2 (Table 2b). The significance of these sequences remains to be determined.

3

4 **2.8 Gene Number**

5 The current assembly predicts that the genome contains around 10,000 genes,
6 almost twice as many as seen in *P. falciparum* (Gardner *et al.*, 2002) or
7 *Saccharomyces cerevisiae* (Goffeau *et al.*, 1996) but closer to that of the free
8 living protist *Dictyostelium discoideum* (ca. 12,500; Eichinger *et al.*, 2005). It
9 should be remembered that this number will change as the assembly improves,
10 and is likely to decrease somewhat. Nevertheless, the comparatively large gene
11 number when compared to some other parasitic organisms reflects both the
12 relative complexity of *E. histolytica* and the presence of large gene families,
13 despite the loss of certain genes as a consequence of parasitism. Gene loss and
14 gain can both represent an adaptive response to life in the human host. Gene
15 loss is most evident in the reconstruction of metabolic pathways of *E.*
16 *histolytica*, which show a consistent pattern of loss of synthetic capacity as a
17 consequence of life in an environment rich in complex nutrient sources.
18 Similarly, analyses of expanded gene families with identifiable functions
19 indicate that many are directly associated with the ability to sense and adapt to
20 the environment within the human host and the ability to ingest and assimilate
21 the nutrients present. One consequence of these gene family expansions being
22 linked to phagocytosis of bacteria and other cells may be an association
23 between many of these gene families and pathogenicity.

24

25 **2.9 Gene Structure**

26 Most *E. histolytica* genes comprise only a single exon; however as many as
27 25% may be spliced and 6% contain two or more introns. Therefore mRNA
28 splicing is far less common than in the related protist *D. discoideum* or the
29 malaria parasite *P. falciparum*. The genome contains all of the essential
30 machinery for splicing (section 2.14) and a comparison of intron positions
31 suggests that *D. discoideum* and *E. histolytica* have both lost introns since their

1 shared common ancestor with *P. falciparum*, although many more have been
2 lost in the *E. histolytica* lineage. A good example of this intron loss is the
3 vacuolar ATP synthase subunit D gene (Figure 1). This protein is highly
4 conserved but the number of introns in each gene varies. *P. falciparum* has 5
5 introns, *D. discoideum* has two and *E. histolytica* has one. The positions of
6 three of the five *P. falciparum* introns are conserved in one of the other species
7 which suggests that these three (at least) were present in the common ancestor
8 and that intron loss has led to the lower number seen in *E. histolytica* today.
9 This loss is consistent with reverse transcriptase mediated 3' intron loss (Roy
10 and Gilbert, 2005) as the 5' -most introns are retained. It would appear that this
11 process has been more active in the *E. histolytica* and *D. discoideum* lineages
12 than in *P. falciparum*, possibly because *Plasmodium* lacks a reverse
13 transcriptase.

14

15 **2.10 Gene Size**

16 Genes in *E. histolytica* are surprisingly short, not only due to the loss of introns
17 but also in the predicted lengths of the proteins they code for. On average the
18 predicted length of a protein in *E. histolytica* is 389 amino acids (aa) which is
19 129 aa and 372 aa shorter than in *D. discoideum* and *P. falciparum* respectively.
20 In fact the protein length distribution is most similar to that of the
21 microsporidian *Encephalitozoon cuniculi* (Figure 2) which has a very compact
22 genome of 3Mb and less than 2000 genes. Direct comparison of orthologous
23 genes between *E. histolytica* and its closest sequenced relative *D. discoideum*
24 demonstrates this phenomenon quite well, with the majority of *E. histolytica*
25 proteins being shorter than the *D. discoideum* counterpart (Hall, unpublished).
26 Protein length is normally very well conserved among eukaryotes so the reason
27 for protein shortening is unclear. It has been postulated that in bacteria reduced
28 protein lengths reflects a reduced capacity for signaling (Zhang, 2000). This
29 would not seem to be the case here as the number of genes identified as having
30 a role in signaling suggests quite the opposite. An alternative theory is that as *E.*

1 *histolytica* has reduced organelles it is possible that its proteins contain fewer or
2 simpler targeting signals.

3

4 **2.11 Protein Domain Content**

5 The most common protein family (Pfam) domains of *E. histolytica* are shown in
6 Table 3. The domains that are unusually common in *E. histolytica* reflect some
7 of the more unusual aspects of the biology of this protist. For example, the Rab
8 and Rho families that are involved in signaling and vesicle trafficking are
9 among the most common domains in *E. histolytica* while in other species they
10 are not often among the top 50 families. This could well be due to the fact that
11 *E. histolytica* has a 'predatory' life style and these domains are intimately
12 involved in environmental sensing, endocytosis and delivery of lysosomes to
13 the phagosome. There are also a number of domains involved in actin
14 dynamics and cytoskeletal rearrangement that are not common in non-
15 phagocytic species, such as the gelsolin and SH3 domains. Myb domains are
16 the most common transcription regulatory domains in *E. histolytica*; this
17 domain is also common in plants where the proteins regulate many plant-
18 specific pathways (Ito, 2005). An important finding from an initial analysis
19 was the presence of unusual multidomain proteins, including five proteins
20 containing both RhoGEF and Arf-GAP domains, suggesting a mechanism for
21 direct communication between the regulators of vesicle budding and
22 cytoskeletal rearrangement. Over 80 receptor kinases were identified (section
23 7.2.2), each containing a kinase domain and a C rich extracellular domain.
24 These kinases fall into distinct classes depending on the presence of CXC or
25 CXXC repeats. There are also domains that are common in most other
26 sequenced genomes but rare or missing from *E. histolytica*. For example, most
27 mitochondrial carrier domain proteins are not needed in *E. histolytica* as it lacks
28 a normal mitochondrion (section 8).

29

30 **2.12 Translation-Related Proteins**

1 Two of the predicted tRNAs (Ile^{TAT} and Tyr) need to be spliced due to the
2 presence of an intron. tRNA introns are distinct in structure from those in
3 protein-coding genes and require a distinct splicing machinery. The expected
4 enzymes required for this splicing are present as are a number of tRNA
5 modification enzymes (including those for synthesising queuine and
6 pseudouridine) and rRNA methylases that act on specific bases in their
7 respective RNA molecules. The expected panel of tRNA synthetases necessary
8 for aminoacylating the tRNAs is also present, with one or two gene copies for
9 each type.

10

11 The majority of ribosomal protein genes are well-conserved in *E. histolytica*
12 and only the gene for large subunit protein L41 could not be identified. The
13 missing protein is only 25 amino acids in length, 17 of which are arginines or
14 lysines, which would make it difficult to identify in this A+T-rich genome, but
15 it is highly conserved, having been reported from Archaea to mammals.
16 However, it also appears to be dispensable, as *S. cerevisiae* can grow relatively
17 normally after deletion of both its copies (Yu and Warner, 2001). Nevertheless,
18 deletion of L41 in *S. cerevisiae* reduces the level of 80S ribosomes, suggesting
19 that it is involved in ribosomal subunit association, reduces peptidyl transferase
20 activity, and increases translocation (Dresios *et al.*, 2003). In addition, L41 has
21 been shown to interact with the beta subunit of protein kinase CKII and to
22 stimulate phosphorylation of DNA topoisomerase II alpha by CKII (Lee *et al.*,
23 1997b). If this gene is truly absent from *E. histolytica* it may have important
24 consequences for the cell.

25

26 No genes for mitochondrial ribosomal proteins were found. Their absence is not
27 surprising since *E. histolytica* lacks typical mitochondria (see section 8 below).

28

29 In eukaryotic translation, elongation factor EF-1 is activated upon GTP binding
30 and forms a ternary complex with aminoacyl tRNAs and ribosomes. EF-1 beta
31 and delta subunits work as GDP-GTP exchange factors to cycle EF-1 alpha

1 between two forms while EF-1 gamma provides structural support for the
2 formation of this multimeric complex. EF2 assists in the translocation of tRNAs
3 on the mRNA by exactly one codon. *E. histolytica* has most of the expected
4 factors except for EF-1 delta, a protein involved in exchanging GDP with GTP.
5 This is also absent from *S. cerevisiae* and *P. falciparum*. It is likely that EF-1
6 beta carries out this activity. It is thought that the EF-1 complex can exist in
7 two forms, EF-1-alpha/beta/gamma and EF-1-alpha/delta/gamma. In *E.*
8 *histolytica*, probably only the former complex exists.

9

10 Eukaryotes typically have two polypeptide release factors, eRF1 and eRF3.
11 Both of these factors have been found in *E. histolytica*.

12

13 **2.13 Analysis of Cell Cycle Genes**

14 Alternation of DNA duplication and chromosome segregation is a hallmark in
15 the cell cycle of most eukaryotes. Carefully orchestrated processes coordinate
16 an ensemble of cell cycle regulating ‘checkpoint’ proteins ensure that progeny
17 cells receive an exact copy of the parental genetic material (Hartwell and
18 Weinert, 1989). Unlike most eukaryotes, *Entamoeba histolytica* cells can
19 reduplicate their genome several times before cell division occurs
20 (Gangopadhyay *et al.*, 1997). Approximately 5-20% of the trophozoites
21 (depending on the growth phase) of axenic culture are multi-nucleated.
22 Additionally, DNA reduplication may occur without nuclear division so that
23 single nuclei contain 1X -6X or more genome contents (Das and Lohia, 2002).
24 Thus axenically cultured *E. histolytica* trophozoites display heterogeneity in
25 their genome content suggesting that eukaryotic cell cycle checkpoints are
26 either absent or altered in this organism. Around 200 genes have been identified
27 in yeast that play a direct role in cell cycle progression.

28

29 *2.13.1 DNA replication initiation and DNA duplication*

30 The DNA replication licensing system is one of the crucial mechanisms that
31 ensures the alternation of S-phase with mitosis in most cells (Tye, 1999).

1 Initiation of DNA replication involves binding of the replicative helicases to
2 DNA replication origins in late mitosis. Loading of the replicative helicase
3 Mcm2-7 proteins is preceded by formation of the pre-replicative complex (pre-
4 RC) and its subsequent activation. Formation of pre-RC requires the ordered
5 assembly of the origin recognition complex (ORC), Cdc6, Cdt1 and the Mcm2-
6 7 proteins. The pre-RC is activated by the protein kinase Cdc7p and its
7 regulatory subunit Dbf4 (Masai and Arai, 2002). Other factors that regulate the
8 transition from pre-RC to replication initiation are Mcm10p, Cdc45p, TopBP1,
9 RecQL4 and the GINS complex (Gregan *et al.*, 2003; Machida *et al.*, 2005;
10 Merchant *et al.*, 1997; Wohlschlegel *et al.*, 2002). Two other Mcm proteins –
11 Mcm8 and Mcm9 - have been identified in metazoan systems and are believed
12 to be part of the replicative helicase (Maiorano *et al.*, 2006). Replication origin
13 licensing is inactivated during S-phase but Mcm2-9p may function as a
14 helicase that unwinds DNA ahead of the replication fork during S-phase
15 (Maiorano *et al.*, 2006). Once S-phase has begun, the formation of new pre-RC
16 is kept in check by high CDK activity and by the activity of the protein geminin
17 (Bell and Dutta, 2002).

18

19 A detailed analysis of the *E. histolytica* genome shows that homologues of
20 several proteins required for DNA replication initiation are absent. These
21 include ORC (Origin Recognition Complex) 2-6, Cdt1, geminin, Cdc7/Dbf4
22 and Mcm10. A single gene encoding a homologue of the archaeal and human
23 Cdc6/Orc1p (Capaldi and Berger, 2004) was identified. This suggests that DNA
24 replication initiation in *E. histolytica* is likely similar to archaeal replication
25 initiation where a single Cdc6p/ORC1p replaces the hetero-hexameric ORC
26 complex (Kelman and Kelman, 2004). Several proteins described from
27 metazoa, such as Cdt1, geminin, Mcm8 and Mcm9, have not been found in
28 yeast. Surprisingly, Mcm8 and Mcm9 were identified in the *E. histolytica*
29 genome.

30

1 Of the four known checkpoint genes that regulate DNA replication in *S.*
2 *cerevisiae* only Mec1 and Mrc1 have homologues in *E. histolytica*. *E.*
3 *histolytica* homologues of several proteins involved in G1-S transitions are
4 absent, such as Sic1, Chk1. The S-phase checkpoint genes p21, p27, p53 and
5 retinoblastoma (RB) required for transition from G1 to S-phase in humans were
6 absent in *E. histolytica*. Chk1 and Chk2 genes encode kinases that act
7 downstream from the ATM and ATR kinases (intra-S phase checkpoint genes).
8 The Chk1 homologue is absent but a Chk2 homologue has been identified in *E.*
9 *histolytica* and partially characterised (Iwashita *et al.*, 2005).

10

11 2.13.2 Chromosome segregation and cell division

12 A large number of genes are known to regulate different events during the
13 transition from G2-Mitosis - spindle formation checkpoint, chromosome
14 segregation, mitosis, exit from mitosis, and cytokinesis - in *S.cerevisiae*. Many
15 of the proteins required by yeast for kinetochore formation have no obvious
16 homologues in *E. histolytica* suggesting that amoeba kinetochores may have an
17 altered composition and structure. Proteins of the Anaphase Promoting
18 Complex (APC) regulate transition from metaphase to anaphase. With the
19 exception of APC11, none of the APC proteins could be identified in
20 *E.histolytica*. In contrast two genes encoding CDC20 homologues, which are
21 known to activate the APC complex, were identified in *E.histolytica* along with
22 ubiquitin and related proteins (Wöstmann *et al.*, 1992), indicating that although
23 most APC subunit homologues were absent the pathway of proteasomal
24 degradation for regulation of cell cycle proteins may still be functional in
25 *E.histolytica*. Effectors of the apoptotic pathway and meiosis were also largely
26 absent.

27

28 2.13.3 CDKs and cyclins

29 The CDC28 gene encodes the single cyclin dependant kinase (CDK) in *S.*
30 *cerevisiae* and regulates cell cycle progression by binding to different cyclins at
31 the G1/S or G2/M boundaries (Reed, 1992; Surana *et al.*, 1991; Wittenberg *et*

1 *al.*, 1990). Similarly, *Schizosaccharomyces pombe* also encodes a single CDK
2 (*cdc2*) (Simanis and Nurse, 1986). Mammals and plants can encode multiple
3 CDKs and an equally large number of cyclins (Morgan, 1995; Vandepoele *et*
4 *al.*, 2002). Association of different CDKs with specific cyclins regulates the
5 cell cycle in different developmental stages as well as in specific tissues. CDKs
6 belong to the serine/threonine family of kinases with a conserved PSTAIRE
7 domain where cyclins are believed to bind (Jeffrey *et al.*, 1995; Morgan, 1996)
8 although some mammalian and plant CDKs have been shown to have divergent
9 PSTAIRE motifs. This heterogeneity may or may not affect cyclin binding
10 (Poon *et al.*, 1997). The *E. histolytica* genome encodes at least 9 different
11 CDKs among which not even one has the conserved PSTAIRE motif. The
12 closest homologue of the CDC28/*cdc2* gene, which shows only conservative
13 substitutions in the PSTAIRE motif (PVSTVRE), was cloned previously (Lohia
14 and Samuelson, 1993). The remaining 8 CDK homologues exhibit even greater
15 divergence in this motif. Eleven putative cyclin homologues with a high degree
16 of divergence have been found. Identifying their CDK/cyclin partner along with
17 their roles in the cell cycle is a major task that lies ahead. Some of the CDKs
18 may not function by associating with their functional cyclin partners but may
19 play a role in regulating global gene expression, either by activation from non-
20 cyclin proteins or by other mechanisms (Nebreda, 2006).

21

22 *E. histolytica* presents a novel situation where the eukaryotic paradigm of a
23 strictly alternating S-phase and mitosis is absent. Discrete G1, S and G2
24 populations of cells are not routinely found in axenic cultures. Instead cells in
25 S-phase show greater than 2x genome contents, suggesting that the G2 phase is
26 extremely short and irregular. This observation together with the absence of a
27 large number of checkpoint genes suggests that regulation of genome
28 partitioning and cell division in *E. histolytica* may be additionally dependant on
29 extracellular signals. *E. histolytica* must however contain regulatory
30 mechanisms to ensure that its genome is maintained and transmitted with
31 precision even in the absence of the expected checkpoint controls. The

1 discovery of these mechanisms will be crucial to our understanding of how the
2 *E. histolytica* cell divides.

3

4 **2.14 Transcription**

5 RNA polymerase II transcription in *E. histolytica* is known to be α -amanitin -
6 resistant (Lioutas and Tannich, 1995). The F homology block of the RNA
7 polymerase II large subunit has been identified as the putative α -amanitin
8 binding site. This block is highly divergent in the α -amanitin resistant
9 *Trichomonas vaginalis* RNA polymerase II (Quon *et al.*, 1996). The *E.*
10 *histolytica* RPB1 homologue also diverges from the consensus in this region
11 but, interestingly, it is also quite dissimilar to the *T. vaginalis* sequence.

12

13 The heptapeptide repeat (TSPTSPS) common to other eukaryotic RNA
14 polymerase II large subunit C terminal domains (CTD) is not present in the *E.*
15 *histolytica* protein. Indeed, the *E. histolytica* CTD is not similar to any other
16 RNA polymerase II domain in the current database. However, the CTD of the
17 *E. histolytica* enzyme does remain proline/serine-rich (these amino acids
18 constitute 40% of the CTD sequence). The *E. histolytica* CTD also retains the
19 potential to be highly phosphorylated: of the 24 serines, 6 threonines and 3
20 tyrosines within the CTD, 9 serines, 3 threonines and 1 tyrosine are predicted to
21 be within potential phosphorylation sites. It is therefore possible that, despite
22 its divergence, modification of the CTD by kinases and phosphatases could
23 modulate protein-protein interactions as is postulated to occur in other RNA
24 polymerases (Yeo *et al.*, 2003). In yeast, phosphorylation of the CTD regulates
25 association with the mediator protein (Davis *et al.*, 2002; Kang *et al.*, 2001;
26 Kornberg, 2001). The yeast mediator protein complex consists of 20 subunits.
27 However, perhaps due to the divergence of the CTD, only two of these proteins
28 have been identified in *E. histolytica* (Med7 and Med10). Homologues of the
29 Spt4 and Spt5 elongation factors, also thought to interact with the CTD, have
30 been identified.

31

1 The RNA polymerase core is composed of 12 putative subunits in *S. cerevisiae*
2 (Young, 1991), while *S. pombe* contains a subset of 10 of these proteins,
3 lacking the equivalents of subunits 4 and 9 (Yasui *et al.*, 1998). In *E.*
4 *histolytica* only 10 of the RNA polymerase subunits have been identified,
5 identifiable homologues of subunits 4 and 12 being absent. While the
6 homologue to subunit 9 was present it lacks the first of the two characteristic
7 zinc binding motifs of this protein and the DPTLPR motif in the C terminal
8 region. A similar sequence, DPTYPK, is however present and a homologue of
9 the TFIIE large subunit Tfa1, which is proposed to interact with this region of
10 the protein, has been identified (Hemming and Edwards, 2000; Van Mullem *et*
11 *al.*, 2002). The conserved N terminal portion (residues 1-52) of Rpb9 is
12 thought to interact with both Rpb1 and Rpb2 in *S. cerevisiae* (Hemming and
13 Edwards, 2000) and homologues of these have been identified.

14

15 The core promoter of *E. histolytica* has an unusual tripartite structure consisting
16 of the three conserved elements TATA, GAAC and INR (Purdy *et al.*, 1996;
17 Singh and Rogers, 1998; Singh *et al.*, 2002; Singh *et al.*, 1997). Singh and
18 Rogers (1998) have speculated that the GAAC motif may be the binding site of
19 a second or alternative *E. histolytica* DNA binding protein in the preinitiation
20 complex. It is therefore of interest that, in addition to the *E. histolytica* TATA-
21 binding protein (TBP), two other proteins contain the TATA-binding motif
22 (Hernandez *et al.*, 1997). TBP is a subunit of the TFIID general transcription
23 factor (GTF) which in other organisms is required for the recognition of the
24 core promoter. In light of the variation in the core promoter previously
25 mentioned, and the divergence in proteins that bind to the core promoter in
26 other parasitic protists, it is not surprising that only six of the 14 evolutionary
27 conserved subunits of TFIID, TBP Associated Factors (TAFs) 1, 5, 6, 10, 12
28 and 13 were identified. Homologues of some of the global regulatory subunits
29 of the Ccr4/Not complex, which interacts with TBP and TAFs 1 and 13, have
30 also been identified.

31

1 TAFs 5, 6, 10 and 12 are also components of the histone acetyltransferase
2 (HAT) complexes in other organisms as is SPT6 and 16 (Carrozza *et al.*, 2003).
3 While all known components of the HAT complexes have by no means been
4 identified or the role of the previously unknown bromodomain containing
5 proteins encoded in the *E. histolytica* genome, histone acetylation complexes
6 are known to be active in *E. histolytica* (Ramakrishnan *et al.*, 2004). Other
7 potential members of chromatin remodeling complexes of *E. histolytica* include
8 the TBP interacting helicase (RVB1 & 2) and the SNF2 subunit of the
9 SWI/SNF complex.

10

11 Homologues of some of the other GTFs (TFII E, F and H) but not the large or
12 small subunits were identified. In contrast to the difficulty identifying some of
13 the GTFs, the *E. histolytica* spliceosomes components U1, U2, U4/6, U5 and
14 the Prp19 complex have all been identified. In fact homologues of ten of the
15 fourteen “core” snRNP proteins, two of the U1 specific snRNPs, seven of the
16 ten U2 specific snRNPs, five of the six U5 specific snRNPs, three of the U4/6
17 specific snRNPs, and four of the nine subunits of the Prp19 complex have been
18 found. In fact *E. histolytica* has homologues of approximately 80% of the *S.*
19 *cerevisiae* splicing machinery (Jurica and Moore, 2003).

20

21 Like *G. intestinalis*, *E. histolytica* has short 5' untranslated regions on its
22 mRNAs. However, unlike those of *G. intestinalis*, *E. histolytica* mRNA has
23 been shown to be capped (Ramos *et al.*, 1997; Vanacová *et al.*, 2003).

24 Identification of homologues of the Ceg1 RNA guanylyltransferase - an
25 enzyme which adds an unmethylated GpppRNA cap to new transcripts - and of
26 Abd1 - which methylates the cap to form m7GpppRNA - gives new insight into
27 the probable cap structure in *E. histolytica* (Hausmann *et al.*, 2001; Pillutla *et*
28 *al.*, 1998). It has been proposed that the capping enzymes interact with the
29 phosphorylated CTD of RNA polymerase (Schroeder *et al.*, 2000). The CTD of
30 *E. histolytica* large subunit is, as discussed earlier, not well conserved but
31 contains several probable phosphorylation sites.

1
2 mRNAs in *E. histolytica* are polyadenylated and the polyadenylation signal is
3 found within the short 3' untranslated region (Bruchhaus *et al.*, 1993; Li *et al.*,
4 2001). However only eight of the eighteen yeast Cleavage and Polyadenylation
5 Specificity Factor (CPSF) subunits are identifiable in *E. histolytica*.

6

7 **3. VIRULENCE FACTORS**

8 **3.1 Gal/GalNAc Lectin**

9 One of the hallmarks of *E. histolytica* pathogenicity is contact-dependent killing
10 of host cells. *E. histolytica* is capable of killing a variety of cells types including
11 human intestinal epithelium, erythrocytes, neutrophils, and lymphocytes
12 (Burchard and Bilke, 1992; Burchard *et al.*, 1992a; Burchard *et al.*, 1992b;
13 Guerrant *et al.*, 1981; Ravdin and Guerrant, 1981). Cytolysis occurs as a step-
14 wise process that begins with adherence to target cells via galactose/N-acetyl
15 D-galactosamine-inhibitable (Gal/GalNAc) lectin (Petri *et al.*, 1987; Ravdin
16 and Guerrant, 1982). Adherence via the Gal/GalNAc lectin is a requirement for
17 cell killing because in the presence of galactose or GalNAc targets cells are not
18 killed by the amoebae. Target cell death occurs within 5 to 15 minutes and is
19 often followed by phagocytosis. Inhibition of the Gal/GalNAc lectin with
20 galactose or specific antibody also blocks phagocytosis (Bailey *et al.*, 1990).
21 Resistance to lysis by the complement system is also mediated in part by the
22 Gal/GalNAc lectin. The lectin contains a CD59-like domain that likely helps
23 protect the trophozoites from complement; CD59 is a surface antigen of many
24 blood cells known to have this property (Braga *et al.*, 1992).

25

26 The Gal/GalNAc lectin is a membrane complex that includes heavy (Hgl) 170
27 kDa, and light (Lgl) 30-35 kDa subunits linked by disulphide bonds, and a non-
28 covalently associated intermediate (Igl) 150 kDa subunit (Cheng *et al.*, 2001;
29 Petri *et al.*, 1989). The structure and function of the Gal/GalNAc lectin has
30 recently been reviewed (Petri *et al.*, 2002). The heavy subunit is a type1
31 transmembrane protein while the light and intermediate subunits have

1 glycosylphosphatidylinositol (GPI) anchors (Cheng *et al.*, 2001; McCoy *et al.*,
2 1993). Gal/GalNAc lectin subunits do not share any significant protein identity
3 or similarity to any other known proteins, though Hgl and Igl have some very
4 limited regions of similarity with known classes of proteins that will be
5 discussed below.

6

7 *3.1.1 The heavy (Hgl) subunit*

8 Based on pulse-field gel electrophoresis there are five loci in the genome with
9 similarity to the Hgl subunit. However, the current genome assembly only
10 identifies two complete genes, one of which corresponds to Hgl2 (Tannich *et*
11 *al.*, 1991b). The predicted proteins encoded by these loci are 92% identical. In
12 initial assemblies there were three other sequences with high similarity to the
13 Hgl subunit that were pseudogenes. These pseudogenes may account for the
14 additional loci detected by pulse-field gel electrophoresis. The large size of
15 these genes means that assembly problems may also be affecting our
16 interpretation.

17

18 Hgl subunit sequences can be divided in to domains based on amino acid
19 content and distribution (Figure 3). The amino-terminal domain of ca. 200
20 amino acids consist of 3.2% cysteine and 2.1% tryptophan residues. The next
21 domain, also ca. 200 amino acids, is completely devoid of these two amino
22 acids. The C-terminal domain of ca. 930 amino acids is cysteine-rich,
23 comprising 10.8 % cysteine. The number and spacing of all predicted
24 tryptophan and cysteine residues are 100% conserved in the two complete
25 genes. Although a portion of the C-terminal domain can be said to contain
26 cysteine-rich pseudo- repeats, there is no clear repetitive nature to the protein
27 (Tannich *et al.*, 1991b). The Hgl subunit has a single transmembrane domain
28 and a highly conserved 41 amino acid cytoplasmic domain. In addition to these
29 two *hgl* genes, the genome contains a newly identified divergent member of the
30 Hgl gene family (XP_650534). This ORF shares 43% similarity with the two

1 other Hgl isoforms, and is predicted to encode a protein with an almost
2 identical domain structure to that of Hgl described above.

3

4

5 3.1.2 *The light (Lgl) subunit*

6 The Lgl subunit is encoded by five genes (*lgl1-5*) that share 74-85% amino acid
7 identity. A sequence corresponding to Lgl2 is missing from the current genome
8 assembly. The light subunits range from 270 to 294 amino acids in length. Each
9 isoform has a 12 amino acid signal peptide, 5 conserved cysteine residues, and
10 a GPI-anchor addition site. Lgl1 has two potential glycosylation sites. Lgl2 has
11 one of these sites, Lgl3 has one different site, and Lgl4 and Lgl5 have none.

12

13 3.1.3 *The intermediate (Igl) subunit*

14 The Igl subunit was first identified by a monoclonal antibody that blocked
15 amoebic adherence to and cytotoxicity for mammalian cells (Cheng *et al.*,
16 1998). Co-purification of the Hgl, Lgl, and Igl suggests that these three subunits
17 form a complex (Cheng *et al.*, 1998; Cheng *et al.*, 2001). The Igl subunit also
18 has galactose-binding activity (Cheng *et al.*, 1998) and can serve as protective
19 antigen in vaccine trials (Cheng and Tachibana, 2001). There are two loci that
20 encode Igl subunits (Cheng *et al.*, 2001) and the predicted amino acid
21 sequences are 81% identical. The Igl subunit, like the Hgl subunit, does not
22 have any recognisable carbohydrate-binding domain.

23

24 3.1.4 *Conservation of Gal/GalNAc lectin subunits in other species of*

25 *Entamoeba*

26 There are clearly identifiable orthologues of the Hgl and Lgl subunits among
27 the limited sequences of *E. dispar*, *E. invadens*, *E. moshkovskii*, and *E.*
28 *terrapinae* available at present (Dodson *et al.*, 1997; Pillai *et al.*, 1997; Wang *et*
29 *al.*, 2003). Because these genomes are incomplete it is possible that as yet
30 unidentified family members will show greater similarity to the *E. histolytica*
31 sequences. Nevertheless, the Lgl subunit is quite conserved among the five

1 *Entamoeba* species. For instance, the *E. terrapinae* gene is 56% identical and
2 62 % similar to *E. histolytica* Lgl1 over a span of 201 amino acids. The Hgl
3 subunits are more diverse. The *E. dispar* Hgl orthologue is highly similar to the
4 *E. histolytica* subunit (86%) but the other species show more diversity,
5 including the region that corresponds to the CRD. However, the number and
6 positions of the cysteine residues are highly conserved, as is the sequence of the
7 cytoplasmic domain, showing only a few changes. It is difficult to put precise
8 numbers to these similarities because the complete sequences of Hgl subunits
9 from the other species are not present in the database. The character of the
10 conservation of the Hgl subunits suggests that the ligand specificity is different
11 for the Hgl subunits of each species but the signaling functions of the
12 cytoplasmic domains are similar, if not perhaps identical. Only *E. dispar* has an
13 identifiable Igl subunit. The other three species clearly have paralogues of the
14 CXXC repeat family to which Igl belongs, but their similarity to Igl is mostly
15 restricted to the CXXC and CXC repeat motifs.

16

17 **3.2 Cysteine endopeptidases**

18 *Entamoeba histolytica* is characterised by its extraordinary capacity to invade
19 and destroy human tissues. The main lytic activity has been attributed to
20 cysteine endopeptidases. This class of enzymes, which is found in all
21 organisms, plays a major role in the pathogenicity of *E. histolytica* as
22 demonstrated in a large number of in vitro and in vivo studies (Ankri *et al.*,
23 1999; Gadasi and Kessler, 1983; Keene *et al.*, 1990; Li *et al.*, 1995; Luaces and
24 Barrett, 1988; Lushbaugh *et al.*, 1985; Reed *et al.*, 1989; Schulte and Scholze,
25 1989; Stanley *et al.*, 1995). Most striking are results from laboratory animal
26 infections showing that *E. histolytica* trophozoites with reduced cysteine
27 proteinase activity are greatly impaired in their ability to induce amoebic
28 disease (Ankri *et al.*, 1999; Stanley *et al.*, 1995). In addition, the discovery that
29 *E. histolytica* cysteine proteinases possess interleukin-1 β convertase activity
30 suggests that these enzymes use a mechanism that is novel in microbial
31 pathogenicity (Zhang *et al.*, 2000).

1
2 Thiol-dependent proteolytic activity in *E. histolytica* was first attributed to a
3 neutral sulphhydryl proteinase (McLaughlin and Faubert, 1977) and later to a
4 cytotoxic proteinase (Lushbaugh *et al.*, 1984). Other terms that have been used
5 to describe closely related or identical enzymes are cathepsin B (Lushbaugh *et*
6 *al.*, 1985), neutral proteinase (Keene *et al.*, 1990), histolysin (Luaces and
7 Barrett, 1988) (later changed to histolysain; Luaces *et al.*, 1992)), and
8 amoebapain (Scholze *et al.*, 1992). *E. histolytica* cysteine endopeptidases were
9 found to be secreted (Leippe *et al.*, 1995) and localised in lysosome-like
10 vesicles or at the surface of the cell (Garcia-Rivera *et al.*, 1999; Jacobs *et al.*,
11 1998). Molecular cloning has revealed a large number of cysteine
12 endopeptidase genes in the *E. histolytica* genome (Bruchhaus *et al.*, 2003;
13 Garcia-Rivera *et al.*, 1999; Reed *et al.*, 1993; Tannich *et al.*, 1991c; Tannich *et*
14 *al.*, 1992). Interestingly, most of these genes are not expressed during in vitro
15 cultivation (Bruchhaus *et al.*, 2003). As our current knowledge of *E. histolytica*
16 biology and pathogenicity is mostly based on analysis of cultured cells, the
17 function of most of the cysteine endopeptidases and their precise role in *E.*
18 *histolytica* virulence is largely unknown.

19
20 Homology searches using conserved active site regions revealed that the *E.*
21 *histolytica* genome contains at least 44 genes coding for cysteine
22 endopeptidases. Of these, the largest group is structurally related to the C1
23 papain superfamily (Table 4), whereas a few others are more similar to family
24 C2 (calpain-like cysteine proteinases), C19 (ubiquitinyl hydrolase), C54
25 (autophagin), and C65 (otubain), respectively (Table 5).

26
27 Phylogenetic analyses of the 36 C1-family members revealed that they
28 represent 3 distinct clades (A, B, C), consisting of 12, 11 and 13 members,
29 respectively. Clade A and B members correspond to the two previously
30 described subfamilies of *E. histolytica* cysteine proteinases, designated EhCP-A
31 and EhCP-B (Bruchhaus *et al.*, 2003). In contrast, clade C represents a new

1 group of *E. histolytica* cysteine endopeptidases that has not been described
2 before. EhCP-A and EhCP-B subfamily members are classical pre-pro enzymes
3 with an overall cathepsin L-like structure (Barrett 1998) as indicated by the
4 presence of an ERFNIN motif in the pro region of at least 21 of the 23 EhCP-A
5 and EhCP-B enzymes (Figure 4). Interestingly, biochemical studies with
6 purified EhCP-A indicated a cathepsin B-like substrate specificity (Scholze and
7 Schulte, 1988). This is likely due to the substitution of an alanine residue by
8 acidic or charged amino acids in the postulated S2 pocket, corresponding to
9 residue 205 of the papain sequence (Barrett 1998). As reported previously
10 (Bruchhaus *et al.*, 2003), the EhCP-A and EhCP-B subfamilies differ in the
11 length of the pro regions as well as of the catalytic domains, and have distinct
12 sequence motifs in the N-terminal regions of the mature enzymes (DWR vs.
13 PCNC). Moreover, none of the EhCP-A subfamily but 10 of the 11 Eh CP-B
14 sequences contain hydrophobic stretches near or at the C-terminus, some of
15 which are predicted to constitute transmembrane helices (TMH) or GPI-
16 attachment moieties. This finding is consistent with previous reports on surface
17 localisation of *E. histolytica* cysteine proteinases but, so far, studies on the
18 cellular localisation of the various EhCP-B molecules have not been reported.

19

20 In contrast to the EhCP-A and EhCP-B subfamilies, primary structure
21 prediction indicates that EhCP-C members are not pre-pro enzymes, as they
22 lack hydrophobic signal sequences as well as identifiable pro regions. Instead,
23 they contain a hydrophobic region located 11 to 28 amino acids from the N-
24 terminus, which is predicted to form a TMH (Figure 4). Therefore, this new
25 group of molecules appears to be membrane associated via a signal anchor. All
26 EhCP-C enzymes have a conserved motif of the sequence H/I(X)₆L/ICP in the
27 C-terminal half but they differ substantially in their pI, with values ranging
28 from 4.6 to 8.8. As there is no example of a structurally related cysteine
29 endopeptidase corresponding to the EhCP-C subfamily in other organisms, the
30 specific functions of this group of molecules remain completely unknown.

31

1 In addition to the large number of C1 superfamily members, the *E. histolytica*
2 genome contains 2 genes encoding cysteine endopeptidases homologous to
3 family C2 or calpain-like cysteine proteases (EhCALP1 and EhCALP2).
4 Enzymes of this class contain several calcium-binding domains and have been
5 shown to participate in a variety of cellular processes including remodeling of
6 the cytoskeleton and membranes, signal transduction pathways and apoptosis.

7

8 Another 4 genes were identified coding for enzymes with homology to the
9 peptidase family C54 also termed autophagins (EhAUTO1-4). The process of
10 autophagy has been studied in human and yeast cells (Kirisako *et al.*, 2000;
11 Marino *et al.*, 2003). Autophagy is a mechanism for the degradation of
12 intracellular proteins and the removal of damaged organelles. During this
13 process the cellular components become enclosed in double membranes and are
14 subsequently degraded by lysosomal peptidases. Autophagins seem to be
15 important for cytoplasm-to-vacuole targeting.

16

17 Two other genes encoding putative cysteine endopeptidases of *E. histolytica*
18 show homology to the C19 and C65 families. These two groups of enzymes are
19 known to be involved in ubiquitin degradation. Family C19 are ubiquitinyl
20 hydrolases described as having ubiquitin-specific peptidase activity in humans.
21 C65 or otubains are a group of enzymes with isopeptidase activity, which
22 releases ubiquitin from polyubiquitin.

23

24 In summary, the *Entamoeba* genome contains a considerable number of
25 endopeptidase genes. Elucidation of the precise role of each of the various
26 enzymes will be a major challenge but may help us to understand the
27 mechanism(s) of virulence and other unique properties of this protistan parasite.

28

29 **3.3 Amoebapores and related proteins**

30 In the lysosome-like granular vesicles of *E. histolytica* is found a family of
31 small proteins, amoebapores, that are cytolytic towards human host cells,

1 display potent antibacterial activity, and cause ion channel formation in
2 artificial membranes (for a review see Leippe (1997)). Three amoebapore
3 isoforms have been isolated and biochemically characterised, and their primary
4 structure has been elucidated by molecular cloning of the genes encoding their
5 precursors (Leippe *et al.*, 1991; Leippe *et al.*, 1992; Leippe *et al.*, 1994b).
6 These membrane-permeabilising polypeptides are discharged by *E. histolytica*
7 into bacteria-containing phagosomes to combat growth of engulfed
8 microorganisms (Andrä *et al.*, 2003). Because of their potent cytolytic activity
9 against human cells in vitro (Berninghausen and Leippe, 1997; Leippe *et al.*,
10 1994a), amoebapores have been viewed as a crucial element of the machinery
11 use by the parasite to kill host cells. Trophozoites of *E. histolytica* lacking the
12 major isoform amoebapore A, whether through antisense inhibition of
13 translation (Bracha *et al.*, 1999) or epigenetic silencing of the gene (Bracha *et*
14 *al.*, 2003), became avirulent demonstrating that this protein plays a key role in
15 pathogenesis. Relatives of these protistan polypeptides are found in granules of
16 porcine and human cytotoxic lymphocytes where they are termed NK-lysin and
17 granulysin, respectively. All of these polypeptides are 70-80 amino acids in
18 length and are characterised by a compact alpha-helical, disulphide-bonded
19 structure known as the saposin-like fold. The structures of the amoebic and
20 mammalian polypeptides have been solved and compared (Anderson *et al.*,
21 2003; Hecht *et al.*, 2004; Leippe *et al.*, 2005; Liepinsh *et al.*, 1997). The
22 biological activities have also been measured in parallel (Bruhn *et al.*, 2003;
23 Gutschmann *et al.*, 2003) to evaluate the similarities and differences of these
24 effector molecules from organisms whose evolutionary paths diverged very
25 early. As they are active against both prokaryotic and eukaryotic target cells,
26 they may be viewed as broad-spectrum effector molecules.

27

28 In the genome of *E. histolytica*, 16 genes coding for putative saposin-like
29 proteins (SAPLIPs) were identified. All of these genes are transcribed by cells
30 growing in axenic culture (Winkelmann *et al.*, 2006). Like amoebapores, the
31 predicted proteins all contain one C-terminal SAPLIP domain and (with one

1 exception) a putative signal peptide (Table 6). As a transmembrane domain is
2 not apparent in these proteins, it may well be that they are secretory products
3 stored in the cytoplasmic vesicles and act synergistically with the amoebapores.
4 However, only four of them have a similar size to amoebapores, the others
5 being considerably larger (up to 1009 residues). At present, it is not clear
6 whether these larger gene products represent precursor molecules that are
7 processed further. None of the novel SAPLIPs contain the conserved unique
8 histidine residue at the C-terminus that is a key residue for the pore-forming
9 activity of amoebapores (Andrä and Leippe, 1994; Hecht *et al.*, 2004; Leippe *et*
10 *al.*, 2005). Indeed, it has recently been shown that recombinant SAPLIP3 has
11 no pore-forming or bactericidal activity, although it does cause membrane
12 fusion in vitro (Winkelmann *et al.*, 2006). This is in agreement with the
13 experimental evidence for only three pore-forming entities being present in
14 trophozoite extracts. Therefore, it is most likely that the three amoebapores are
15 the sole pore-forming molecules of the parasite. However, the lipid-interacting
16 activity present in all SAPLIP proteins (Munford *et al.*, 1995) and a function
17 that helps to kill bacterial prey may well characterise all members of the
18 amoebapore/SAPLIP superfamily of this voraciously phagocytic cell.

19

20 **3.4 Antioxidants**

21 *Entamoeba histolytica* trophozoites usually reside and multiply within the
22 human gut, which constitutes an anaerobic or microaerophilic environment.
23 However, during tissue invasion, the amoebae are exposed to an increased
24 oxygen pressure and have to eliminate toxic metabolites such as reactive
25 oxygen or nitrogen species (ROS/RNS) produced by activated phagocytes
26 during the respiratory burst. *E. histolytica* lacks a conventional respiratory
27 electron transport chain that terminates in the reduction of O₂ to H₂O. However,
28 *E. histolytica* does respire and tolerates up to 5% oxygen in the gas phase (Band
29 and Cirrito, 1979; Mehlotra, 1996; Weinbach and Diamond, 1974). Thus, *E.*
30 *histolytica* trophozoites must use different antioxidant enzymes for the removal
31 of ROS, RNS and oxygen (Figure 5).

1
2 Among the enzymes in the first line of oxidative defence are superoxide
3 dismutases (SODs), which are metalloproteins that use copper/zinc (Cu/Zn),
4 manganese (Mn) or iron (Fe) as metal cofactors. SODs catalyse the dismutation
5 of superoxide radical anions to form H₂O₂ and O₂ (Fridovich, 1995). Analysis
6 of the *E. histolytica* genome revealed only a single gene coding for a FeSOD
7 and no sequences encoding MnSOD or Cu/ZnSOD. This reflects the situation
8 found in most protistan parasites and is consistent with biochemical studies
9 previously performed on *E. histolytica* lysates (Tannich *et al.*, 1991a).

10

11 *Entamoeba histolytica* lacks the tripeptide glutathione (Fahey *et al.*, 1984),
12 which constitutes the major low molecular weight thiol found in almost all
13 aerobic cells (Sies, 1999). Instead, *E. histolytica* uses cysteine as its principal
14 low molecular weight thiol (Ariyanayagam and Fairlamb, 1999; Fahey *et al.*,
15 1984; Nozaki *et al.*, 1999). As expected, coding sequences for enzymes that use
16 glutathione as a cofactor, such as glutathione-S-transferase, glutathione-
17 dependent peroxidase, glutathione reductase or glutaredoxin, are all absent from
18 the *E. histolytica* genome. In addition, genes encoding catalases and
19 peroxidases are also missing, as previously suggested (Sykes and Band, 1977;
20 Weinbach and Diamond, 1974).

21

22 Other genes were identified that code for proteins involved in detoxification of
23 H₂O₂, including one with homology to rubrerythrin. Rubrerythrin is a non-
24 haeme iron protein thought to be able to reduce H₂O₂ as part of an oxidative
25 stress protection system (Weinberg *et al.*, 2004). So far, the nature of its redox
26 partner is unknown in *E. histolytica* and it remains to be determined whether
27 protection against oxidative stress is indeed its main function. Another group of
28 H₂O₂-detoxifying proteins identified in *E. histolytica* are peroxiredoxins.
29 Peroxiredoxins are known from a wide variety of organisms . They are able to
30 reduce H₂O₂ as well as peroxynitrite with the use of electrons provided by
31 thiols. In addition to involvement in the detoxification of reactive oxygen

1 species peroxiredoxins seem to play a role in other processes such as signalling
2 and differentiation (Hofmann *et al.*, 2002; Rhee *et al.*, 2005; Wood *et al.*,
3 2003a,b). All peroxiredoxins contain a conserved cysteine residue that
4 undergoes a cycle of peroxide-dependent oxidation and thiol-dependent
5 reduction during the reaction. The whole protein family can be divided into
6 three classes based on the number and position of active site Cys residues (2-
7 Cys, atypical 2-Cys, and 1-Cys peroxiredoxins; Wood *et al.*, 2003a,b). In *E.*
8 *histolytica* five different genes coding for peroxiredoxins were identified (Prx1-
9 5). They all belong to the 2-Cys peroxiredoxin family. Four of them (Prx1-4)
10 share 98% sequence identity and have an unusual N-terminal Cys-rich repeat
11 (KECCKKECQEKECQEKECCC) of unknown function. In contrast, the fifth
12 peroxiredoxin (Prx5) lacks the cysteine-rich N-terminal extension and shares
13 only 30% identity with Prx1-4. Biochemical studies have shown that *E.*
14 *histolytica* peroxiredoxins are able to detoxify H₂O₂ and cumene hydroperoxide
15 (Bruchhaus *et al.*, 1997; Poole *et al.*, 1997). Moreover, up-regulation of
16 peroxiredoxin and FeSOD was associated with metronidazole resistance in
17 cultured *E. histolytica* trophozoites (Samarawickrema *et al.*, 1997; Wassmann
18 *et al.*, 1999).

19

20 Reactions catalysed by peroxiredoxins are dependent on the presence of
21 physiological thiols like thioredoxin (Rhee *et al.*, 2005; Wood *et al.*, 2003b).
22 Thioredoxins are small proteins involved in thiol-redox processes (Holmgren,
23 2000). They contain two redox-active site cysteine residues of the motif CXXC
24 (Watson *et al.*, 2004). Five genes coding for classical cytoplasmic thioredoxins
25 were identified in the *E. histolytica* genome (Trx1-5). These thioredoxins have
26 a length of 103-114 amino acids and share 25 – 47 % sequence identity. Trx1-3
27 have identical active site motifs of the sequence WCGPC, whereas the active
28 sites of Trx4 and Trx5 have the sequences SCPSC and WCKDC, respectively.
29 In addition, another five thioredoxin-related proteins were identified (Trx6-10).
30 All have a signal sequence of 15 to 19 amino acid residues and the active site
31 motif WCGHC, which is also known from the active site of protein disulphide

1 isomerases. However, in contrast to the latter group of enzymes, the *E.*
2 *histolytica* thioredoxin-related molecules contain only one rather than two
3 active-site motifs and only two of the proteins have an ER membrane retention
4 signal (Freedman *et al.*, 2002). Thus it remains to be determined whether the
5 thioredoxin-related molecules of *E. histolytica* do constitute protein disulphide
6 isomerases or whether they undertake other functions within the cell.

7

8 Thioredoxins are kept in the reduced state by the enzyme thioredoxin reductase
9 which catalyses the reduction of oxidised thioredoxin by NADPH using FAD
10 and its redox-active disulphide (Nakamura, 2005). Two different genes with
11 homology to thioredoxin reductases have been previously described from *E.*
12 *histolytica* (thioredoxin reductase (TrxR) and NADPH:flavin oxidoreductase
13 (p34)). They share about 87% sequence identity and both contain the two
14 conserved sequence motifs forming the FAD and NAD(P)H binding domains.
15 p34 was shown to catalyse the NADPH-dependent reduction of oxygen to H₂O₂
16 as well as of disulphides like DTNB and cystine (Bruchhaus *et al.*, 1998; Lo
17 and Reeves, 1980). Therefore, in addition to disulphide reductase activity the
18 enzyme has H₂O₂-forming NADPH oxidase activity. It was also shown that p34
19 can transfer reducing equivalents to peroxiredoxin, converting the protein from
20 its non-active, oxidised form back into its active, reduced form (Bruchhaus *et*
21 *al.*, 1997). However, it is unlikely that peroxiredoxin is directly reduced by p34
22 *in vivo*. It is more likely that *E. histolytica* contains a classical thioredoxin
23 redox system consisting of thioredoxin reductase, thioredoxin and
24 peroxiredoxin (Poole *et al.*, 1997).

25

26 In addition to genes coding for proteins with homology to thioredoxin
27 reductase, four other gene families were identified that encode various
28 flavoproteins. One of these families includes 4 members that have between
29 53% and 61% sequence identity to A-type flavoproteins
30 (flavorubredoxin/flavodiiron). A-type flavoproteins belong to a large family of
31 enzymes that are widespread among anaerobic and facultatively anaerobic

1 prokaryotes. In addition to bacteria, homologous genes are also found in the
2 genomes of the pathogenic amitochondriate protistan parasites *Trichomonas*
3 *vaginalis* and *Giardia intestinalis* (Andersson *et al.*, 2003; Sarti *et al.*, 2004).
4 The A-type flavoproteins are made up of two independent structural modules.
5 The N-terminal region forms a metallo-beta-lactamase-like domain, containing
6 a non-haeme di-iron site, whereas the C-terminal region is a flavodoxin-like
7 domain, containing one FMN moiety. These enzymes have significant nitric
8 oxide reductase activity (Gomes *et al.*, 2002; Sarti *et al.*, 2004). For *Escherichia*
9 *coli* it is known that the nitric oxide reductase (FIRd) receives electrons from a
10 NADH:oxidoreductase (FIRd-red). Consistent with that situation, the *E.*
11 *histolytica* genome contains a gene encoding an NADH oxidase with 25%
12 sequence identity to several bacterial FIRd -reds.

13

14 The three other *E. histolytica* gene families with homology to iron-sulphur
15 flavoproteins (families B-D) are characterised by the presence of a flavodoxin-
16 like domain forming a typical FMN binding site. Family B and family C consist
17 of three members each, which share sequence identity of 42% and 46%,
18 respectively. Family D consists of two members, which share only 33%
19 sequence identity. At present, the function of the various flavodoxin-like
20 molecules remains to be determined and deserves to be investigated fully,
21 particularly as to whether they do indeed have antioxidant capacity.

22

23 **4. METABOLISM**

24 Biochemical analysis of *E. histolytica* metabolism has a long history (Reeves,
25 1984), dating back to shortly after the development of culture media that
26 allowed the generation of substantial numbers of axenic cells. The genome
27 sequence has confirmed most of the predicted metabolic pathways shown
28 biochemically to be present or absent in *E. histolytica* in the past. As with most
29 parasites, secondary loss of biosynthetic pathways is a recurring theme.
30 However, a few surprises have also been uncovered. Every single enzyme
31 involved in metabolism cannot realistically be discussed in this review. In this

1 section the only the major energy generating and biosynthetic aspects of
2 metabolism will be covered. Enzyme names, EC numbers and accession
3 numbers are given in the the supplementary table for this section.

4

5 **4.1 Energy Metabolism**

6 *4.1.1 Glycolysis*

7 *E. histolytica* lacks a functional tricarboxylic acid (TCA) cycle and oxidative
8 phosphorylation. It is not able to convert organic substrates such as glucose into
9 H₂O and CO₂, but has to rely on the energy generated by various types of
10 substrate level phosphorylation (Reeves, 1984). Glycolysis is the major
11 pathway of ATP generation, but in addition the genome project has identified a
12 number of genes that could result in more ATP generation through the
13 catabolism of amino acids. These enzymes will be described further below. As
14 *E. histolytica* lacks compartmentalised energy generation, it has been classified
15 as a type I amitochondriate protist (Martin and Müller , 1998) in contrast to the
16 type II amitochondriate protists containing hydrogenosomes, such as
17 *Trichomonas vaginalis*. Nevertheless, it does contain a mitochondrial remnant,
18 the mitosome (see section 8).

19

20 In *E. histolytica*, glycolysis appears to be localised in the cytosol. This is in
21 contrast to trypanosomes where a major part is carried out in the glycosomes
22 (Parsons, 2004) and the pathway is regarded as a potential target for
23 chemotherapy (Opperdoes and Michels, 2001). The kinetic properties of
24 recombinant *E. histolytica* glycolysis enzymes have recently been studied by
25 Saavedra *et al.* (2005). Their analysis suggested that fructose-1,6-bisphosphate
26 aldolase, phosphoglycerate mutase, glyceraldehyde-3-phosphate
27 dehydrogenase, and pyruvate phosphate dikinase might be regulating the
28 glycolytic flux.

29

30 4.1.1 (a) Hexokinases

1 Glucose taken up by *E. histolytica* is phosphorylated by two hexokinase (EC
2 2.7.1.1) isoenzymes (Hxk1 and Hxk2). The two *E. dispar* isoenzymes are
3 shifted towards a slightly more basic pI, which is the basis of the classical
4 biochemical method for distinguishing *E. histolytica* from *E. dispar* by starch
5 gel electrophoresis (Farri *et al.*, 1980). The pI differences among the two *E.*
6 *histolytica* isoforms (Ortner *et al.*, 1995) and between the two species (Ortner *et*
7 *al.*, 1997b) are the result of genetic differences that lead to different amino acid
8 sequences and charge differences. Hxk1 phosphorylates glucose and mannose,
9 while Hxk2 phosphorylates mainly glucose and is much less active with
10 mannose as a substrate (Kroschewski *et al.*, 2000).

11

12 4.1.1 (b) Glucose-6-phosphate isomerase

13 Glucose 6-phosphate is converted to fructose 6-phosphate by glucose-6-
14 phosphate isomerase (EC 5.3.1.9). The genome has 2 genes for this enzyme,
15 which code for proteins that differ only by a single insertion/deletion of 7
16 amino acid residues. Glucose-6-phosphate isomerase is another of the enzymes
17 for the classical differentiation of *Entamoeba zymodemes* by starch gel
18 electrophoresis (Sargeant, 1987).

19

20 4.1.1 (c) Phosphofructokinases

21 The main phosphofructokinase activity in *E. histolytica* is PPi-dependent (EC
22 2.7.1.90; Reeves *et al.*, 1976). There is a single gene (Deng *et al.*, 1998)
23 encoding this 60 kDa enzyme. The gene is a candidate for lateral transfer from
24 bacteria (Loftus *et al.*, 2005) (see section 10). The enzyme is expressed at a
25 tenfold higher level and displays about tenfold higher activity than a second
26 phosphofructokinase of 48 kDa (XP_653373) (Chi *et al.*, 2001). The substrate
27 specificity of the smaller enzyme is disputed. Whereas Bruchhaus *et al.* (1996)
28 reported that this minor enzyme also used PPi as phosphate donor, Chi *et al.*
29 (2001) found only an ATP-dependent activity. The 48 kDa and 60 kDa
30 enzymes are highly divergent with less than 20% sequence identity.
31 Interestingly, the specificity of the 60 kDa phosphofructokinase can be changed

1 from PPi to ATP by mutation of a single amino acid residue (Chi and Kemp,
2 2000). The authors concluded that ATP rather than PPi was the primordial high
3 energy compound. In the genome, there are two additional genes encoding
4 isoforms of the 48 kDa enzyme, which have not been studied at the protein
5 level.

6

7 4.1.1 (d) Fructose-1,6-bisphosphate aldolase

8 Fructose 1,6-bisphosphate is cleaved to glyceraldehyde 3-phosphate and
9 dihydroxyacetone 3-phosphate by fructose-1,6-bisphosphate aldolase (EC
10 4.1.2.13). The enzyme, a Class II aldolase (Marsh and Lebherz, 1992) has been
11 cloned (XP_650373) and exhibits strong sequence similarity to eubacterial
12 aldolases (Sanchez *et al.*, 2002). A second gene (XP_655966) encodes a protein
13 differing from the first by a single deletion of 28 amino acids flanked by short
14 divergent stretches. These bacterial-type aldolases are also found in
15 *Trichomonas vaginalis*, *Giardia intestinalis* and other protists (Sanchez *et al.*,
16 2002). *E. histolytica* has no gene coding for a Class I aldolase like those found
17 in animals, which might make aldolase an interesting target for chemotherapy.

18

19 4.1.1 (e) Triose-phosphate isomerase

20 Triose-phosphate isomerase (EC 5.3.1.1) converts dihydroxyacetone 3-
21 phosphate into glyceraldehyde 3-phosphate. The gene was previously cloned
22 (Landa *et al.*, 1997), and is highly similar to the annotated gene product. This
23 dimer-forming enzyme represents the first *E. histolytica* protein for which the
24 structure has been solved by X-ray crystallography (Rodriguez-Romero *et al.*,
25 2002).

26

27 4.1.1 (f) Glyceraldehyde-3-phosphate dehydrogenase

28 Glyceraldehyde-3-phosphate dehydrogenase (EC 1.2.1.12) oxidises and
29 phosphorylates glyceraldehyde 3-phosphate to 1,3-bisphosphoglycerate in two
30 coupled reactions using NAD⁺ as cofactor (Reeves, 1984). The genome project
31 revealed five putative genes, three of which encode the identical protein

1 sequence of 36.0 kDa and a predicted pI of 7.04. The fourth gene product,
2 XP_648981, differs from these three only by a 13 amino acid deletion, while
3 XP_650370 is a clearly distinct 34.8 kDa isoform with a lower predicted pI of
4 5.80. Interestingly, the isoforms XP_650356 and XP_650370 of different pI are
5 encoded within the same contig.

6

7 4.1.1 (g) Phosphoglycerate kinase

8 Phosphoglycerate kinase has an unusual substrate (Reeves and South, 1974),
9 transferring the high energy phosphate group from 3-phosphoglyceroyl
10 phosphate to GDP leading to the formation of GTP (EC 2.7.2.10). There is one
11 candidate gene encoding a 45 kDa protein.

12

13 4.1.1 (h) Phosphoglycerate mutase

14 Phosphoglycerate mutase (Reeves, 1984) isomerises 3-phosphoglycerate to 2-
15 phosphoglycerate (EC 5.4.2.1). Five divergent putative genes for this enzyme
16 are found in the genome. Two gene products of 62 kDa were classified as 2,3-
17 bisphosphoglycerate-independent phosphoglycerate mutases (XP_649031 and
18 XP_654182); they differ only at their C-termini and display significant
19 similarity to bacterial phosphoglycerate mutases. The three other genes are very
20 divergent. XP_651808 was identified as a candidate for lateral gene transfer
21 (Loftus *et al.*, 2005) (see section 10). The remaining two gene products
22 XP_649053 and XP_657284 are related to genes found in both prokaryotes and
23 eukaryotes.

24

25 4.1.1 (i) Enolase (2-phosphoglycerate dehydratase)

26 Enolase (EC 4.2.1.11) converts 2-phosphoglycerate to phosphoenolpyruvate.
27 The gene has been cloned (Beanan and Bailey, 1995) and the protein
28 characterised (Hidalgo *et al.*, 1997) previously. The 47 kDa gene product is a
29 typical eukaryotic enolase (XP_649161). A carboxy-terminally truncated
30 incomplete ORF is also found.

31

1 4.1.1 (j) Pyruvate, orthophosphate dikinase and pyruvate kinase

2 In *E. histolytica*, both activities forming ATP and pyruvate from
3 phosphoenolpyruvate have been found. The exergonic pyruvate kinase reaction
4 uses ADP (Saavedra *et al.*, 2004), and the pyruvate, orthophosphate dikinase
5 uses AMP and PPi in a slightly endergonic reaction (Varela-Gomez *et al.*,
6 2004). The dikinase activity is found in C4 plants where it is involved in
7 phosphoenolpyruvate generation for gluconeogenesis. In *E. histolytica* it was
8 discovered long before the pyruvate kinase (Reeves, 1968).

9

10 The cloning of pyruvate, orthophosphate dikinase (EC 2.7.9.1) was reported by
11 two groups. The published sequences (Bruchhaus and Tannich, 1993; Saavedra
12 Lira *et al.*, 1992) are highly similar or identical to XP_657332 and XP_654666.
13 In addition there are two shorter related open reading frames.

14

15 In the genome 3 putative pyruvate kinase genes (EC 2.7.1.40) have been
16 identified. The 3 are identical except for an amino-terminal deletion in
17 XP_648240 and an internal deletion in XP_653635.

18

19 4.1.1 (k) Pyruvate:ferredoxin oxidoreductase (PFOR) and ferredoxin

20 PFOR (EC 1.2.7.1) is an enzyme of major importance to *E. histolytica*, as the
21 parasite lacks NAD⁺-dependent pyruvate dehydrogenase and pyruvate
22 decarboxylase (Reeves, 1984). No evidence for the latter two genes was found
23 in the genome, confirming the biochemical results. PFOR oxidatively
24 decarboxylates pyruvate to acetyl-CoA. The electrons are transferred to
25 ferredoxin which, in its reduced form, can activate and reduce metronidazole,
26 the major anti-amoebic drug (Müller, 1986). The activated form of
27 metronidazole can potentially react with a number of biomolecules and is able
28 to cleave the parasite DNA. In human cells, metronidazole is not activated and
29 is much less toxic. In *T. vaginalis*, down-regulation of PFOR is one mechanism
30 of producing metronidazole resistance (Kulda, 1999); however PFOR
31 expression appears unaltered in partially resistant *E. histolytica*

1 (Samarawickrema *et al.*, 1997; Wassmann *et al.*, 1999). All eukaryotic PFOR
2 genes, including that of *E. histolytica*, appear to have been acquired during an
3 ancient lateral gene transfer event from bacteria (Horner *et al.*, 1999; Rotte *et*
4 *al.*, 2001). There are two putative PFORs in the *E. histolytica* genome
5 displaying minor sequence differences.

6

7 The genome contains 7 ferredoxin genes in total with 5 quite divergent
8 sequences. All are related to eubacterial and archaeal ferredoxins (Nixon *et al.*,
9 2002). The gene pairs XP_655183 / XP_655182 and XP_654311 / XP_652694
10 are identical. The other three gene products represent more divergent open
11 reading frames. The deduced proteins have similar molecular masses, between
12 6.1 kDa and 8.8 kDa, and different predicted isoelectric points between 4.2 and
13 8.6.

14

15 4.1.1 (l) Acetyl-CoA synthetase (acetate thiokinase)

16 The normal fate of acetyl-CoA in mitochondriate organisms is entry into the
17 tricarboxylic acid cycle. However this pathway is absent from *E. histolytica*.
18 Instead, the cleavage energy of the thioester bond of acetyl-CoA can be used to
19 generate one ATP molecule. One of the known acetyl-CoA synthetases
20 generates ATP from ADP and Pi (EC 6.2.1.13). Such an enzyme has been
21 characterised by Reeves *et al.* (1977) and cloned (Field *et al.*, 2000), and
22 reported to be a 77 kDa protein. The common acetyl-CoA synthetase activity
23 that produces ATP from AMP and P_{PPi} (EC 6.2.1.1) appears to be absent in *E.*
24 *histolytica*.

25

26 4.1.1 (m) Aldehyde and alcohol dehydrogenases

27 The *E. histolytica* genome encodes a complex system of alcohol and/or
28 aldehyde dehydrogenases. In total, there are 25 predicted genes, 3 of which are
29 on the list of lateral gene transfer (LGT) candidates.

30

1 Alcohol dehydrogenase ADH1 was the first alcohol dehydrogenase to be
2 characterised in *E. histolytica* (Reeves *et al.*, 1971) and is a NADPH-dependent
3 enzyme (EC 1.1.1.2). The gene was previously cloned (Kumar *et al.*, 1992); in
4 the genome three genes are almost identical to that sequence, while one
5 (XP_652772) has 67% identity.

6
7 Fermentation in *E. histolytica* uses the bifunctional alcohol dehydrogenase /
8 aldehyde dehydrogenase NADH-dependent enzyme ADH2, which belongs to
9 the ADHE family and has both alcohol dehydrogenase and aldehyde
10 dehydrogenase activities (Lo and Reeves, 1978). Under anaerobic conditions,
11 reduction of the acetyl-CoA generated by PFOR to ethanol is one way to
12 regenerate the NAD⁺ used by glyceraldehyde-3-phosphate dehydrogenase.
13 ADH2 first reduces acetyl-CoA to an enzyme-bound hemiacetal which is then
14 hydrolysed to acetaldehyde (EC 1.2.1.10) and further reduced to ethanol (EC
15 1.1.1.1). If the enzyme is also able to work in the reverse direction, *E.*
16 *histolytica* would be able to generate acetyl-CoA and energy from ethanol in
17 the presence of oxygen. This would explain older reports of ethanol stimulated
18 oxygen uptake in *E. histolytica* (Weinbach and Diamond, 1974). The enzyme is
19 closely related to AdhE from *E. coli* and other bacteria (Reid and Fewson,
20 1994) and there is strong support for its acquisition by LGT (Andersson *et al.*,
21 2006; Field *et al.*, 2000; Loftus *et al.*, 2005) (see section 10). Like its bacterial
22 homologue, ADH2 appears to form helical rods that sediment with membrane
23 fractions (Avila *et al.*, 2002). Two groups have previously cloned ADH2
24 (Bruchhaus and Tannich, 1994; Yang *et al.*, 1994), and in total the genome
25 contains 5 full-length ADH2 genes and one that is truncated. All share between
26 98% and 100% sequence identity.

27
28 In total, there are 11 alcohol dehydrogenase ADH3 genes in the genome, two of
29 which been reported previously (Kimura *et al.*, 1996; Rodriguez *et al.*, 1996).
30 The recombinant enzyme characterised by Rodriguez *et al.* (1996) was
31 NADPH-specific, like ADH1. There are five genes similar to these previously

1 reported sequences. The rest of the ADH3 sequences fall into two groups of 3
2 similar sequences. All 11 ADH3 sequences are between 44% and 100%
3 identical on the amino acid level. XP_649823 was originally on the list of LGT
4 candidates (Loftus *et al.*, 2005), and a similarity to ADH3 sequences of gram-
5 negative bacteria had been noted before (Nixon *et al.*, 2002). However a related
6 sequence is now known to exist in *T. vaginalis* also (see section 10).

7

8 The genome encodes three additional distinct alcohol dehydrogenases.
9 XP_656535 is a putative Zn-containing enzyme, and is on the list of LGT
10 candidates. XP_652753 has been annotated as a Fe-containing alcohol
11 dehydrogenase and XP_652262 simply as putative alcohol dehydrogenase.

12

13 One NADPH-dependent aldehyde dehydrogenase encoding gene (ALDH1) is
14 present and was reported previously (Zhang *et al.*, 1994).

15

16 4.1.2 Energy storage: the glycogen metabolism

17 *E. histolytica* uses glycogen as its major energy store. Glycogen is a polymer of
18 alpha-1,4-linked glucose chains with alpha-1,6 branch points, which in *E.*
19 *histolytica* has a compact structure as suggested by branch points every 5-6
20 glucose residues (Bakker-Grunwald *et al.*, 1995). The cytoplasm of
21 trophozoites contains numerous glycogen granules which were first observed
22 by electron microscopy (Rosenbaum and Wittner, 1970) and later characterised
23 biochemically (Takeuchi *et al.*, 1977). A glycogen phosphorylase activity (EC
24 2.4.1.1), associated with the glycogen granules, generates glucose 1-phosphate
25 from orthophosphate and the linear portion of various glucopolysaccharides
26 (Werries and Thurn, 1989). The genome contains at least 6 putative full-length
27 and truncated genes encoding glycogen phosphorylases, two of which were
28 cloned by Wu and Müller (2003). These authors noted a marked sequence
29 divergence in those regions of the enzymes involved in regulation by
30 phosphorylation and concluded that classical regulation by phosphorylation
31 may not occur.

1
2 Glycogen phosphorylase degrades the linear chains only down to the alpha-1,6
3 branch points. The remaining core molecule is called limit dextrin. Degradation
4 can proceed further with the help of a debranching enzyme that has been
5 purified (Werries *et al.*, 1990). It exhibits activities of both amylo-1,6-
6 glucosidase (EC 3.2.1.33) and 4-alpha-glucanotransferase (EC 2.4.1.25). The
7 genome contains two genes putatively encoding a full-length (XP_653608) and
8 a truncated glycogen debranching enzyme. The deduced molecular mass of the
9 large protein is 166 kDa which corresponds to the biochemical data (Werries *et*
10 *al.*, 1990).

11
12 Glucose 1-phosphate is isomerised to glucose 6-phosphate by
13 phosphoglucomutase (EC 5.4.2.2) before entering the glycolytic pathway. The
14 isoelectric points of the phosphoglucomutases from *E. histolytica* and *E. dispar*
15 differ which was exploited for differentiation of the two species by starch gel
16 electrophoresis (Sargeant *et al.*, 1978). The migration properties are
17 reproduced by recombinant enzymes and are the result of primary sequence
18 differences (Ortner *et al.*, 1997a). *E. histolytica* has one gene coding for this
19 important enzyme, and in addition there are two distantly related members of
20 the phosphoglucomutase / phosphomannomutase family.

21
22 Genes encoding the enzymes involved in glycogen biosynthesis in *E. histolytica*
23 have been identified: a glycogen synthase (EC 2.4.1.11) of 155 kDa and two
24 putative branching enzymes (EC 2.4.1.18). The glycogen precursor UDP-
25 glucose is generated from UTP and glucose 1-phosphate by UTP:glucose-1-
26 phosphate uridylyltransferase (EC 2.7.7.9). Two UTP-hexose-1-phosphate
27 uridylyltransferases have been characterised biochemically, a larger glucose 1-
28 phosphate-specific enzyme of 45 kDa and a less specific enzyme of 40 kDa
29 reported to use both galactose 1-phosphate and glucose 1-phosphate (Lobelle-
30 Rich and Reeves, 1983). The genome contains one larger open reading frame
31 encoding a putative UTP:glucose-1-phosphate uridylyltransferase of 54.7 kDa

1 and two smaller ones encoding enzymes of 46.3 kDa with high similarity
2 identified as UTP:N-acetyl-glucosamine-1-phosphate uridyltransferases. These
3 enzymes are interesting in that they could possibly be involved in the activation
4 of N-acetyl-glucosamine 1-phosphate as a precursor of the chitin cyst wall.

5

6 4.1.3 Catabolism of sugars other than glucose

7 4.1.3 (a) Activation of fructose and galactose for glycolysis

8 Neither Hxk1 nor Hxk2 can use fructose or galactose as a substrate, but there
9 are 2 genes encoding bacterial-type enzymes that may do so, a 33 kDa
10 fructokinase, which is one of the candidates for lateral gene transfer to the *E.*
11 *histolytica* lineage (see section 10), and a 43 kDa galactokinase. The
12 fructokinase groups with bacterial fructose 6-kinases (EC 2.7.1.4), and the
13 galactokinase groups with galactose 1-kinases (EC 2.7.1.6). This substrate
14 specificity has been noted before (Reeves, 1984). Fructose 6-phosphate enters
15 as an intermediate of the glycolytic pathway (see 4.1.1 (c)). As described above
16 (4.1.2), galactose 1-phosphate can be activated to UDP-galactose (Lobelle-Rich
17 and Reeves, 1983), and then epimerised to UDP-glucose by UDP-glucose 4-
18 epimerase (EC 5.1.3.2) (Reeves, 1984). In the genome, a single candidate 38
19 kDa ORF for the latter enzyme has been identified. The UDP-bound glucose
20 can then be used either for the synthesis of glycogen or fed into the glycolysis
21 pathway via glucose 1-phosphate and glucose 6-phosphate. This efficient
22 pathway allows *E. histolytica* to grow on galactose instead of glucose (Reeves,
23 1984).

24

25 4.1.3 (b) Anomerisation of aldoses

26 The 1-position in the pyranose form of aldoses has a hydroxyl group that can be
27 in either the α - or β -configuration. These forms can be interconverted by means
28 of an aldose 1-epimerase (EC 5.1.3.3) an enzyme that has recently been
29 characterised (Villalobo *et al.*, 2005). There is a single gene encoding this
30 product.

31

1 4.1.3 (c) Activation of pentoses

2 Two gene candidates encoding pentose-activating enzymes have been identified
3 in the *E. histolytica* genome: a 35 kDa ribokinase (EC 2.7.1.15) and a 56 kDa
4 xylulokinase (EC2.7.1.17). The latter is another bacterial-type sequence
5 putatively acquired by lateral gene transfer.

6

7 4.1.3 (d) Interconversion of hexoses and pentoses

8 The pathway of interconversion between hexoses and pentoses in *E. histolytica*
9 was described many years ago (Reeves, 1984; Susskind *et al.*, 1982). A
10 transketolase (EC 2.2.1.1) converts fructose 6-phosphate and glyceraldehyde 3-
11 phosphate into xylulose 5-phosphate and erythrose 4-phosphate. Erythrose 4-
12 phosphate and dihydroxyacetone phosphate are condensed by the glycolytic
13 enzyme fructose-1,6-bisphosphate aldolase to sedoheptulose 1,7-bisphosphate,
14 an extended substrate specificity of the aldolase. Phosphofructokinase then is
15 able to remove a phosphate group forming diphosphate and sedoheptulose 7-
16 phosphate. This molecule and glyceraldehyde 3-phosphate are then converted
17 by transketolase to the pentoses ribose 5-phosphate and xylulose 5-phosphate.
18 A transaldolase activity is absent (Reeves, 1984) consistent with there being no
19 such gene in the genome. In contrast, 7 gene products were identified as likely
20 transketolases: three highly similar proteins of 73 kDa and four truncated
21 versions.

22

23 **4.2 Amino acid catabolism**

24 *4.2.1 General features*

25 As discussed above, glycolysis under anaerobic conditions can use only part of
26 the energy contained in glucose for ATP generation. *E. histolytica* is capable
27 not only of taking up amino acids (Reeves, 1984), but also using them for the
28 generation of energy, as suggested by Zuo and Coombs (1995). The genome
29 has revealed a number of unusual genes, often with bacterial affinities, coding
30 for enzymes of amino acid catabolism (Anderson and Loftus, 2005).

31

1 In many cases, the degradation of amino acids starts with a transamination
2 reaction (EC 2.6.1. -) generating a 2-ketoacid. The *E. histolytica* genome has
3 five ORFs identified as aminotransferases. These ORFs are distinct from each
4 other with the exception of XP_655090 and XP_655099, which differ only by
5 one insertion and are LGT candidates. So far there is no enzymological data on
6 this group of enzymes, so their substrate specificities in *E. histolytica* are
7 unknown.

8

9 Both amino acid degradation and glycolysis have 2-ketoacids as intermediates.
10 Pyruvate is one common intermediate, as amino acid degradation can produce
11 either pyruvate or other 2-ketoacids. PFOR (see 4.1.1 (k)) is known to have a
12 relaxed specificity, and in addition to pyruvate it can oxidatively decarboxylate
13 2-ketobutanoate, oxaloacetate, and 2-ketoglutarate (Samarawickrema *et al.*,
14 1997). The reaction generates CoA-thioesters with the potential of producing
15 one ATP per molecule.

16

17 The amino acids asparagine, aspartate, serine, alanine, tryptophan, cysteine,
18 threonine, methionine, glutamine, and glutamate can all be transformed into one
19 of these 2-ketoacids in one or very few steps. This underlines the major
20 importance of the PFOR in the energy metabolism of *E. histolytica*. The
21 enzyme is indispensable, and as it always generates reduced ferredoxin it will
22 always activate metronidazole. Consequently, it would be very difficult for *E.*
23 *histolytica* to become resistant to metronidazole.

24

25 4.2.1 Aspartate and asparagine

26 *E. histolytica* takes up asparagine and aspartate in the presence or absence of
27 glucose (Zuo and Coombs, 1995). Four putative asparaginases (EC 3.5.1.1) are
28 found in the genome. Three are identical and share only 48% amino acid
29 identity with the fourth (XP_656586). Asparaginase mediates the formation of
30 aspartate from asparagine by releasing ammonia. The predicted sequences
31 appear to possess a signal sequence, as suggested by TargetP program

1 (www.cbs.dtu.dk/services/TargetP/), which is reminiscent of a periplasmic
2 isotype (EcA, type II) (Swain *et al.*, 1993) that is up-regulated under anaerobic
3 and carbon-restricted conditions (Cedar and Schwartz, 1967).

4

5 Aspartate can be converted to fumarate and ammonia by aspartate ammonia-
6 lyase (aspartase, EC 4.3.1.1). Addition of a water molecule by fumarase (EC
7 4.2.1.2) produces malate. The genome encodes a putative fumarase that is
8 related to bacterial Class I fumarases. The aspartase is a member of the bacterial
9 Class II fumarase / aspartase protein family (Woods *et al.*, 1988), and also on
10 the list of LGT candidates.

11

12 Aspartate is also decomposed into oxaloacetate and ammonia by aspartate
13 aminotransferase, with the concomitant production of glutamate from 2-
14 oxoglutarate. Oxaloacetate is then converted into malate via malate
15 dehydrogenase (EC 1.1.1.37) and, since *E. histolytica* lacks both a functional
16 TCA cycle and phosphoenolpyruvate carboxykinase, the malate generated can
17 be oxidatively decarboxylated to pyruvate by malic enzyme (EC 1.1.1.39). Both
18 of these enzymes are present in *E. histolytica*. Two very similar genes have
19 been identified as encoding malic enzyme and are LGT candidates.

20

21 4.2.2 Serine, threonine

22 Serine and threonine are also taken up by *E. histolytica* in the presence and
23 absence of glucose (Zuo and Coombs, 1995). Serine can be deaminated by the
24 pyridoxal phosphate-dependent serine dehydratase (L-serine ammonia-lyase,
25 EC 4.3.1.17) to pyruvate and ammonia. The enzyme was characterised by
26 Takeuchi *et al.* (1979) who showed that addition of serine to the culture
27 medium stimulated oxygen consumption. In an analogous reaction, threonine
28 dehydratase (threonine ammonia-lyase, EC 4.3.1.19) breaks down threonine to
29 2-oxobutanoate. Both ketoacids can then be oxidised by PFOR to acetyl-CoA
30 or propionyl-CoA. Both catabolic reactions can be carried out by the same
31 enzyme, as has been shown in yeast for example (Ramos and Wiame, 1982). In

1 the *E. histolytica* genome annotation, four gene products have been annotated
2 as threonine dehydratases, but none as serine dehydratase. XP_650405 and
3 XP_652480 are identical while XP_655614 and XP_657171 share 95% and
4 37% identity with the others, respectively. The exact substrate specificities of
5 these 4 putative serine / threonine dehydratases have not been reported.

6
7 Degradation of serine via the non-phosphorylated serine pathway, by the
8 sequential reactions of L-serine: pyruvate aminotransferase (EC 2.6.1.51), D-
9 glycerate dehydrogenase (EC 1.1.1.29), and D-glycerate kinase (EC 2.7.1.31)
10 (Snell, 1986) results in the glycolytic intermediate 3-phosphoglycerate. The
11 genome encodes several putative aminotransferases (see section 4.2.1), but it is
12 not yet known if serine is among their substrates. An unusual bacterial-type
13 NADPH-dependent D-glycerate dehydrogenase was characterised by Ali *et al.*
14 (2003) and there are 2 genes encoding D-glycerate dehydrogenases, one of
15 which (XP_648124) is among the weaker LGT candidates (see section 10). The
16 genome also contains 2 genes encoding identical glycerate kinases. The enzyme
17 has recently been characterised by Ali and Nozaki (unpublished).

18
19 *4.2.3 Methionine, homocysteine and cysteine*

20 Methionine γ -lyase (EC 4.4.1.11) decomposes methionine to methanethiol
21 (mercaptomethane), ammonia, and 2-oxobutanoate. In *E. histolytica*, two
22 methionine γ -lyases, EhMGL1 and EhMGL2, of similar molecular weights
23 have been characterised (Tokoro *et al.*, 2003). These two isoenzymes show
24 marked differences in substrate specificity, isoelectric point, enzymological and
25 biochemical parameters (Tokoro *et al.*, 2003). Both enzymes can also act on
26 other amino acids. In addition to degrading methionine, both EhMGL1 (pI
27 6.01) and EhMGL2 (pI 6.63) can convert homocysteine to hydrogen sulphide,
28 ammonia and 2-oxobutanoate. EhMGL2 also decomposes cysteine to hydrogen
29 sulphide, ammonia, and pyruvate, whereas EhMGL1 is only weakly active
30 against cysteine. Decomposition of homocysteine by methionine γ -lyase is
31 essential since this parasite lacks the other known enzymes capable of

1 destroying this toxic amino acid. In the genome, three open reading frames
2 correspond to EhMGL1 and one to EhMGL2. So far, the only eukaryotes
3 known to possess methionine γ -lyases are *E. histolytica* and *T. vaginalis*
4 (Lockwood and Coombs, 1991). As the enzymes are absent from the human
5 host and important for the generation of metabolic energy, they could be targets
6 for chemotherapy (Coombs and Mottram, 2001; Tokoro *et al.*, 2003).

7

8 In addition to serving as a source of metabolic energy, another important role of
9 methionine is as a donor of methyl groups via S-adenosylmethionine synthetase
10 (synonymous with methionine adenosyltransferase, EC 2.5.1.6). Seven gene
11 candidates were identified, four full-length and three truncated. The S-
12 adenosylhomoserine left after the transfer of the activated methyl group can be
13 hydrolysed by S-adenosylhomocysteine hydrolase (EC 3.3.1.1) giving
14 adenosine and homocysteine. Two candidate genes with identical sequences
15 and one truncated form are present.

16

17 However, *E. histolytica* lacks the remaining enzymes for the reverse
18 transsulphuration pathway (forming cysteine from methionine) (Nozaki *et al.*,
19 2005), i.e. cystathionine β -synthase and cystathionine γ -lyase. In addition, *E.*
20 *histolytica* lacks all enzymes involved in the forward transsulphuration
21 (forming methionine from cysteine) including cobalamin-dependent methionine
22 synthase (EC 2.1.1.13) or cobalamin-independent methionine synthase (EC
23 2.1.1.14), which suggests that *E. histolytica* is capable of neither converting
24 homocysteine to cystathionine nor recycling homocysteine to methionine.

25

26 *E. histolytica* lacks the methylthioadenosine cycle enzymes except for two, 5'-
27 methylthioadenosine/S-adenosyl homocysteine nucleosidase (EC 3.2.2.9) and
28 aspartate aminotransferase (AT, EC 2.6.1.1). The significance of these two
29 enzymes in *E. histolytica* is unknown.

30

31 *4.2.4 Arginine*

1 In *G. intestinalis* and *T. vaginalis* the arginine deiminase (EC 3.5.3.6) pathway
2 is important for energy generation (Knodler *et al.*, 1994; Linstead and
3 Cranshaw, 1983; Schofield and Edwards, 1994), generating one ATP molecule
4 from the breakdown of arginine to ornithine. In contrast, no arginine deiminase
5 gene or dihydrolase pathway was detected in the *E. histolytica* genome.

6
7 In *E. histolytica*, arginine can either be degraded by arginase (EC 3.5.3.1) via
8 ornithine or arginine decarboxylase (EC 4.1.1.19) via agmatine. The arginine
9 decarboxylase reaction uses up protons and may be involved in the acid
10 resistance needed for the passage of cysts through the human stomach
11 (Anderson and Loftus, 2005). Another function suggested for arginine
12 degradation was that it depletes arginine as a substrate for human macrophages,
13 preventing NO synthesis and amoebicidal activity (Elnekave *et al.*, 2003). Both
14 enzymes could also be important for the generation of the polyamine putrescine
15 (see 4.3 below). The genome contains a single gene encoding a 96 kDa
16 polypeptide annotated as ornithine/arginine/lysine decarboxylase, the substrate
17 specificity of which has not yet been examined on the recombinant protein
18 level. There is a single gene encoding a putative 33 kDa arginase.

19

20 4.2.5 Glutamate, glutamine

21 In aerobic organisms, the 2-oxoglutarate generated from glutamate in a
22 transaminase reaction enters the citric acid cycle for further catabolism. In *E.*
23 *histolytica*, which also contains transaminases, 2-oxoglutarate can be oxidised
24 by PFOR to give succinyl-CoA from which one molecule of ATP can be
25 generated.

26

27 Several other gene products of *E. histolytica* could act on glutamine and
28 glutamate. The genome lacks a glutaminase (EC 3.5.1.2) to carry out the simple
29 hydrolysis of glutamine. Instead there is a putative glucosamine-fructose-6-
30 phosphate aminotransferase (EC 2.6.1.16), which uses the energy in the amide

1 group of glutamine to generate glucosamine 6-phosphate from fructose 6-
2 phosphate. This product may be used for cyst wall biosynthesis.

3

4 *4.2.6 Tryptophan*

5 Tryptophan can be degraded to indole, pyruvate, and ammonia by the PLP-
6 dependent enzyme tryptophanase (EC 4.1.99.1), for which one candidate gene
7 exists. To date, tryptophanase has only been found in bacteria, never in
8 eukaryotes, and it is also on the list of LGT candidates.

9

10 *4.2.7 Alanine: a possible special case*

11 Alanine could potentially be transformed into pyruvate by alanine
12 aminotransferase (synonymous with alanine:pyruvate transaminase, EC
13 2.6.1.2). However, *E. histolytica* is reported to excrete alanine (Zuo and
14 Coombs, 1995) suggesting that this enzyme is not used under the culture
15 conditions tested. Conceivably, the purpose of the excretion process may be to
16 carry excess nitrogen out of the cell in the absence of a functional urea cycle.

17

18 *4.2.8 Catabolism of other amino acids*

19 Most of the enzymes for branched-chain amino acid metabolism are missing in
20 *E. histolytica* but leucine, isoleucine, and valine could be transformed to 2-
21 oxoisocaproate, 2-oxo-3-methylvalerate, and 2-oxovalerate, respectively, by a
22 putative branched-chain amino acid aminotransferase (EC 2.6.1.42), one of the
23 aminotransferases mentioned above (section 4.2). This could produce ammonia
24 or transfer the amino group to 2-oxoglutarate to form glutamate. Subsequent
25 oxidative decarboxylation to give the respective CoA-derivatives could be
26 envisaged but so far no gene candidates for the necessary dehydrogenases have
27 been identified.

28

29 One gene encodes a putative histidine ammonia-lyase (EC 4.3.1.3), which is
30 responsible for the decomposition of histidine into urocanate and ammonia.

31 Other than the formation of ammonia, the significance of this enzyme is not

1 clear since the downstream enzymes involved in histidine catabolism from
2 urocanate to glutamate were not found.

3

4 Currently, there is little information regarding the fate of the amino acids
5 glycine, proline, phenylalanine, tyrosine, and lysine in *E. histolytica*. No genes
6 for the catabolic enzymes necessary were detected except for a LGT candidate
7 bacterial-type 96 kDa broad-specificity ornithine/arginine/lysine decarboxylase
8 that may be acting on lysine.

9

10 **4.3 Polyamine Metabolism**

11 The absence of *S*-adenosyl-L-methionine decarboxylase (EC 4.1.1.50), which
12 converts *S*-adenosyl methionine into decarboxylated *S*-adenosyl methionine,
13 spermidine synthase (EC 2.5.1.16), and spermine synthase (EC 2.5.1.22)
14 suggests a complete lack of polyamine metabolism in this parasite (Anderson
15 and Loftus, 2005). However, as mentioned above, *E. histolytica* possesses
16 genes encoding arginase and arginine decarboxylase. Both could be involved
17 in the production of putrescine via agmatine and agmatinase (EC 3.5.3.11) or
18 via ornithine and ornithine decarboxylase (EC 4.1.1.17). The high putrescine
19 concentration in trophozoites demonstrated by NMR spectroscopy (9.5 mM)
20 (Bakker-Grunwald *et al.*, 1995) reinforces the physiological significance of
21 putrescine. However, the fate of putrescine is unknown as neither spermine nor
22 spermidine has been demonstrated in *E. histolytica*.

23

24 There is controversy regarding the presence or absence of trypanothione, a
25 spermidine-containing thiol, in *E. histolytica*. Trypanothione is a major thiol in
26 trypanosomes and leishmania (Fairlamb and Cerami, 1992) and contains two
27 molecules of glutathione joined by a spermidine linker. The first reports
28 detected the presence of trypanothione in *E. histolytica* (Ondarza *et al.*, 1997)
29 but were contradicted soon after (Ariyanayagam and Fairlamb, 1999). More
30 recently another study reaffirmed its presence (Ondarza *et al.*, 2005). However,
31 the gene encoding trypanothione reductase reported from *E. histolytica* strain

1 HK-9 (AF503571) has no homologue in the genome of HM-1:IMSS. Although
2 this matter has not been resolved, there is general agreement that the major thiol
3 in *E. histolytica* is cysteine (Fahey *et al.*, 1984).

4

5 The *E. histolytica* genome encodes a 46 kDa ornithine decarboxylase with
6 similarity to both plant and vertebrate enzymes and there is also the 96 kDa
7 ornithine/arginine/lysine decarboxylase (see section 4.2.4). Only the former
8 enzyme has been characterised at the biochemical level (Arteaga-Nieto *et al.*,
9 2002) and has been shown to be insensitive to difluoromethylornithine
10 (DFMO), as is *E. histolytica* (Gillin *et al.*, 1984).

11

12 The conversion of arginine into putrescine via agmatine, in a reaction initiated
13 by arginine decarboxylase, is generally present in bacteria and plants. Although
14 arginine decarboxylase is present in *E. histolytica*, agmatinase (EC 3.5.3.11),
15 which further catalyses conversion of agmatine to putrescine and urea, appears
16 absent. However, one gene identified as a 33 kDa arginase also shares 21%
17 sequence identity with human mitochondrial agmatinase and therefore its
18 substrates need to be examined on the biochemical level to see whether the
19 enzyme can act on arginine, agmatine, or both. At present, the role of arginine
20 decarboxylase in *E. histolytica* is not clear, although as mentioned above this
21 enzyme may also be involved in acid resistance in *E. histolytica*.

22

23 **4.4 Biosynthesis of Amino Acids**

24 *4.4.1 Cysteine and serine*

25 One of the areas in which reduction of metabolism is most evident is in amino
26 acid biosynthesis. Biosynthetic pathways for most amino acids other than serine
27 and cysteine (Ali *et al.*, 2003; Ali *et al.*, 2004a; Nozaki *et al.*, 1998a; Nozaki *et*
28 *al.*, 1999) have been lost in *E. histolytica*. Similarly, *P. falciparum*, which
29 predominantly acquires amino acids from host haemoglobins, lacks
30 biosynthesis of most amino acids (Gardner *et al.*, 2002). Intracellular
31 concentrations of some amino acids (glutamate, leucine, valine, and proline, in

1 descending order of abundance) are very high in *E. histolytica* ranging from 6-
2 21 mM (Bakker-Grunwald *et al.*, 1995). In particular, the glutamate and proline
3 concentrations are much higher in the cells than in the growth medium (21 and
4 7.3 mM vs. 5.9 and 1.8 mM, respectively). Glutamate accounts for over one
5 third of the total amino acid pool (Bakker-Grunwald *et al.*, 1995), and is likely
6 to play a central role in homeostasis not only of amino acids but of energy
7 metabolism in general. Thus, it is likely that these amino acids are actively
8 taken up by as-yet unidentified amino acid transporters.

9

10 Retention of the serine and cysteine biosynthetic pathways when the others
11 have been lost is likely related to the physiological importance of cysteine,
12 which is the major intracellular thiol of this parasite. The cysteine biosynthetic
13 pathway consists of two major steps, catalysed by serine acetyltransferase (EC
14 2.3.1.30), which produces *O*-acetylserine from serine and acetyl-coenzyme A,
15 and cysteine synthase (EC 2.5.1.47), which subsequently transfers an alanyl
16 moiety from *O*-acetylserine to sulphide to produce cysteine. *E. histolytica*
17 possesses three genes each for cysteine synthase and serine acetyltransferase.
18 Cysteine synthase 1 and 2 were considered to be allelic isotypes (Nozaki *et al.*,
19 1998b), while cysteine synthase 3 appears to be distinct, with only 83% identity
20 to cysteine synthase 1 and 2. In contrast, all three serine acetyltransferase genes
21 seem to be distinct, showing only 48-73% identity (Ali and Nozaki,
22 unpublished). It was previously shown that cysteine synthase 1/2 and serine
23 acetyltransferase 1 are unique in that (a) they do not form a heterocomplex, in
24 contrast to other organisms (Bogdanova and Hell, 1997; Droux *et al.*, 1998) and
25 (b) serine acetyltransferase 1 is sensitive to allosteric inhibition by both L-
26 cysteine and L-cystine (Nozaki *et al.*, 1999). Since all variants of these two
27 enzymes lack organelle-targeting sequences, the significance of the multiple
28 isotypes is unknown. It is important to determine subcellular distribution and
29 specific functions of these isotypes to understand the significance of the
30 redundancy. As this pathway is absent in humans, it is a rational target for
31 development of new chemotherapeutic drugs against amoebiasis.

1
2 Serine is synthesised de novo utilising the glycolytic intermediate 3-
3 phosphoglycerate, in a pathway that includes three sequential reactions
4 catalysed by D-phosphoglycerate dehydrogenase (EC 1.1.1.95), phospho-L-
5 serine aminotransferase (EC 2.6.1.52), and O-phospho L-serine phosphatase
6 (EC 3.1.3.3). Although the final enzyme has not yet been enzymologically and
7 functionally analysed, the first two enzymes have been characterised (Ali and
8 Nozaki, 2006; Ali *et al.*, 2004a).

9

10 4.4.2 Interconversion of glutamate-glutamine and aspartate-asparagine

11 The single step interconversions of glutamate and glutamine, catalysed by
12 glutamate synthase (EC 1.4.1.13) and glutamine synthetase (EC 6.3.1.2), and of
13 aspartate and asparagine by asparagine synthase (EC 6.3.5.4) are found in *E.*
14 *histolytica*. There are two isotypes of glutamine synthetase with 47% amino
15 acid identity and 5 candidate genes. NADPH-dependent glutamate synthase
16 (EC 1.4.1.13) catalyses the formation of two glutamates from glutamine and 2-
17 oxo-glutarate in bacteria, yeast, and plants, and together with glutamine
18 synthetase is involved in ammonia fixation under ammonia-restricted
19 conditions. NADPH-dependent glutamate synthase is normally composed of
20 two large and two small subunits (Petoukhov *et al.*, 2003). Although three
21 genes encoding the small subunit are present, the large subunit appears to be
22 absent in *E. histolytica*. These putative NADPH-dependent glutamate synthase
23 small subunits share 80% amino acid identity and show 44% amino acid
24 identity to homologues from the Archaea. The similarity to archaeal-type
25 glutamate synthase (Nesbo *et al.*, 2001) suggests that the *E. histolytica* small
26 subunits may function as a glutamate synthase without the large subunit, as
27 shown for *gltA* from the archaean *Pyrococcus* (Jongsareejit *et al.*, 1997).

28

29 The two enzymes that catalyse interconversion between aspartate and
30 asparagine, aspartate ammonia ligase (EC 6.3.1.1) and asparaginase (EC
31 3.5.1.1; see 4.2.1), are present in *E. histolytica*. Two types of aspartate

1 ammonia ligases, AsnA and AsnB, are known from other organisms: the former
2 utilises only ammonia, while the latter uses both ammonia and glutamine as
3 amide donors in a reverse reaction. Mammals possess only AsnA, whereas
4 prokaryotes have both AsnA and AsnB (Boehlein *et al.*, 1996; Nakamura *et al.*,
5 1981). Interestingly, *E. histolytica* possesses only the AsnB homologue. Thus,
6 the amoebic enzyme is likely involved in the formation of glutamate from
7 glutamine, in addition to asparagine formation from aspartate.

8

9 4.4.3 Synthesis of glutamate and aspartate

10 Glutamate can be formed from 2-oxo-glutarate and ammonia in a reversible
11 reaction catalysed by glutamate dehydrogenase (EC 1.4.1.2), which is present
12 in *E. histolytica*. It is known that this enzyme plays a dominant role in ammonia
13 fixation under ammonia-non-restricted conditions as this reaction consumes no
14 ATP. In addition, glutamate dehydrogenase is also involved in gluconeogenesis
15 from glutamate.

16

17 Aspartate ammonia lyase (synonymous with aspartase, EC 4.3.1.1), which
18 decomposes aspartate into fumarate and ammonia in a reversible reaction, is
19 also present in *E. histolytica* (see 4.2.1 above).

20

21 4.5 Lipid Metabolism

22 For *E. histolytica*, the lack of oxidative phosphorylation means that the high
23 energy content of lipids such as fatty acids cannot be exploited. Therefore lipids
24 such as phospholipids and cholesterol are primarily membrane components in
25 *E. histolytica* (Das *et al.*, 2002; Sawyer *et al.*, 1967). Although these
26 components are acquired mainly from their food or from the human host, *E.*
27 *histolytica* does have some capability for biosynthesis, as well as extending and
28 remodeling lipids, and for attaching lipids to proteins.

29

30 4.5.1 Lipid biosynthetic capabilities

31 4.5.1 (a) Polyisoprene biosynthesis and protein prenylation

1 Cholesterol is an important membrane constituent generated from C₅ isoprene
2 precursors. *E. histolytica* trophozoites in axenic culture need cholesterol in their
3 growth medium (Reeves, 1984), and it is likely that they acquire it from their
4 human host. Reeves (1984) even cites several studies which show that
5 hypercholesteremia in the host increases the damage inflicted by amoebic
6 infection. *E. histolytica* lacks several enzymes for the classical sterol
7 biosynthesis pathway (Schroepfer, 1981). The first stage of sterol biosynthesis
8 is the formation of isopentenyl- or dimethylallyl diphosphate. In the *E.*
9 *histolytica* genome no candidate genes for the generation of these intermediates
10 were found, neither for the mevalonate pathway nor for the mevalonate-
11 independent methylerythritol 4-phosphate (MEP) pathway that operates in
12 bacteria and plants (Hunter *et al.*, 2003; Rohmer *et al.*, 1993). In a later step
13 towards cholesterol synthesis, two molecules of C₁₅ farnesyl diphosphate are
14 dimerised to give C₃₀ presqualene diphosphate (EC 2.5.1.21). This enzyme
15 activity and those catalysing the subsequent steps also appear to be absent. The
16 genome data thus support the long standing conclusion that cholesterol
17 biosynthesis is absent from *E. histolytica*.

18

19 Unexpectedly, the *E. histolytica* genome appears to encode enzymes involved
20 in the intermediate stages of cholesterol biosynthesis from C₅ isopentenyl
21 diphosphate to C₁₅ farnesyl diphosphate. The latter compound, and the larger
22 C₂₀ compound geranylgeranyl diphosphate, may serve as precursors for the
23 hydrophobic modification of GTP-binding proteins allowing them to bind to
24 membranes (Grunler *et al.*, 1994). Protein prenylation is a ubiquitous process. It
25 is important in human cell biology, health and disease (McTaggart, 2006), but it
26 is also essential for parasites such that protein farnesylation has been proposed
27 as a potential novel target for anti-parasitic chemotherapy (Maurer-Stroh *et al.*,
28 2003) including anti-*E. histolytica* chemotherapy (Ghosh *et al.*, 2004).

29

30 The first enzyme in this pathway is the isopentenyl-diphosphate delta-isomerase
31 which catalyses the conversion of isopentenyl diphosphate to dimethylallyl

1 diphosphate (EC 5.3.3.2). There is a single gene encoding this enzyme that is of
2 presumed bacterial origin and is on the list of LGT candidates. The two
3 isomeric C₅ isoprenyl diphosphates undergo condensation to C₁₀ geranyl
4 diphosphate, catalysed by geranyl-diphosphate synthase (EC 2.5.1.1). Farnesyl-
5 diphosphate synthase (EC 2.5.1.10) then adds another C₅ unit to give C₁₅
6 farnesyl diphosphate. Finally geranylgeranyl-diphosphate synthase (EC
7 2.5.1.29) adds another C₅ prenyl unit to give C₂₀ geranylgeranyl diphosphate.
8 The genome contains five putative prenyl transferase genes, which all have
9 been annotated as geranylgeranyl-diphosphate synthases. Their sequences are
10 highly similar, with the exception that the open reading frames are disrupted in
11 two of them (XP_650479 and XP_655958). These prenyl transferases appear to
12 be of bacterial origin as well, and XP_650913 is on the list of LGT candidates.
13 When searching for geranyl-diphosphate synthase or farnesyl-diphosphate
14 synthase in the *E. histolytica* genome, the closest matches are for the same
15 genes, so that the substrate specificity of these enzymes is unclear and needs to
16 be examined biochemically.

17

18 The *E. histolytica* genome contains one sequence each for the alpha and beta
19 chains of protein farnesyltransferase (EC 2.5.1.58), which were previously
20 cloned and characterised as recombinant proteins (Kumagai *et al.*, 2004).

21

22 In addition to the protein farnesyltransferase, a protein
23 geranylgeranyltransferase I (EC 2.5.1.59) beta chain has recently been cloned
24 and expressed together with the protein farnesyltransferase alpha chain
25 (Makioka *et al.*, 2006). The heterodimeric molecule had protein
26 geranylgeranyltransferase activity of unusually broad substrate specificity. The
27 alpha and beta chains of the protein (Rab-) geranylgeranyltransferase II (EC
28 2.5.1.60) have also been cloned, as cDNAs (Kumagai *et al.* unpublished
29 results).

30

1 The *E. histolytica* genome encodes candidate enzymes for the modification of
2 prenylated proteins. There are two highly divergent proteins both identified as
3 CAAX prenyl proteases (EC 3.4.24.84). CAAX is the carboxy-terminus of the
4 substrate protein, in which C is the prenylated cysteine residue, A is an aliphatic
5 amino acid and X is the terminal residue. The proteases cleave after the
6 modified cysteine. After the processing step, a prenylcysteine carboxyl
7 methyltransferase (EC 2.1.1.100) methylates the carboxy-terminal residue;
8 there are two divergent candidate genes for this enzyme.

9
10 Taken together, the *E. histolytica* genome contains all the necessary genes to
11 encode the pathway from isopentenyl diphosphate to a processed farnesylated
12 or geranylgeranylated protein. The source of the starting material, isopentenyl
13 diphosphate, remains unknown at this time, but there may be a previously
14 unknown pathway for its synthesis or *E. histolytica* may be able to acquire it
15 from its environment.

16

17 4.5.1 (b) Fatty acid biosynthesis

18 *E. histolytica* encodes an unusual 138 kDa acetyl-CoA carboxylase with two
19 bacterial-type carboxylase domains, an acetyl-CoA carboxylase and a pyruvate
20 carboxylase. Since no biotin carboxylase domain is found in the *E. histolytica*
21 genome, it was proposed that the enzyme removes a carboxyl group from
22 oxaloacetate and transfers it to acetyl-CoA, forming malonyl-CoA and pyruvate
23 (Jordan *et al.*, 2003; Loftus *et al.*, 2005). This fusion protein has not been
24 identified in any organisms other than *Giardia* and *Entamoeba*.

25

26 In the classical pathway of fatty acid biosynthesis, starting from acetyl-CoA
27 sequential two-carbon units are added from malonyl-CoA. In each round of
28 extension, the beta-keto group is reduced in three steps before a new two-
29 carbon unit is added. The whole pathway is carried out in a large fatty acid
30 synthase complex, where the growing chain is linked to an acyl carrier protein.
31 *E. histolytica* lacks this classical pathway. There are, however, plant

1 homologues of fatty acid chain elongases such as *Arabidopsis thaliana* KCS1
2 (Todd *et al.*, 1999). There are eight putative fatty acid elongases in the *E.*
3 *histolytica* genome, and all are very similar to each other. These enzymes could
4 be involved in elongation of fatty acids taken up from the host or food sources,
5 but their function and substrate specificity are unknown at this time.

6

7 4.5.2 Phospholipid metabolism

8 Phospholipids amount to 60-70% of the total lipids in *E. histolytica* (Sawyer *et*
9 *al.*, 1967). So far little information is available at the biochemical level on how
10 phospholipids are synthesised, acquired or remodelled. The genome project has
11 revealed a number of genes indicating that the phospholipid metabolism could
12 be more complex than expected.

13

14 4.5.2 (a) Phospholipid biosynthesis

15 In order to produce phospholipids one has to generate the important
16 intermediate phosphadidate (1,2-diacylglycerol 3-phosphate) by
17 phosphorylation and acylation of glycerol. *E. histolytica* contains one gene for a
18 glycerol kinase (EC 2.7.1.30). The second step would be the transfer of the acyl
19 group to glycerol-3-phosphate by glycerol-3-phosphate O-acyltransferase (EC
20 2.3.1.15), but no candidate gene for this enzyme has been found in the genome.
21 There are, however, two potential 1-acylglycerol-3-phosphate O-
22 acyltransferases (EC 2.3.1.51) that could attach the second acyl group. After the
23 attachment of the acyl groups, and in preparation for the attachment of the
24 activated aminoalcohols, the phosphate is removed by phosphadidate
25 phosphatase (EC 3.1.3.4), for which there is one gene, resulting in a
26 diacylglycerol.

27

28 The activation of ethanolamine (EC 2.7.1.82) or choline (EC 2.7.1.32) for
29 attachment to the phosphadidate starts with phosphorylation. There are two
30 genes identified as choline/ethanolamine kinases that share 37% amino acid
31 identity. Next, ethanolamine phosphate and choline phosphate are converted to

1 CDP-ethanolamine (EC 2.7.7.14) and CDP-choline (EC 2.7.7.15), respectively.
2 The genome encodes two enzymes sharing 57% sequence identity that are
3 identified as ethanolamine-phosphate cytidylyltransferases. The substrate
4 specificity of these enzymes needs to be examined on the biochemical level.
5 Finally the activated ethanolamine or choline is attached to diacylglycerol by
6 the enzymes ethanolaminephosphotransferase (EC 2.7.8.1) or diacylglycerol
7 cholinephosphotransferase (EC 2.7.8.2) producing phosphatidylethanolamine or
8 phosphatidylcholine, respectively. For these activities a total of 8 possible
9 genes are found that share varying degrees of sequence similarity.
10
11 In *E. histolytica*, an alternative pathway of phospholipid biosynthesis could
12 involve the biosynthesis of phosphatidylserine. In this pathway, the
13 phosphatidate itself is activated by CTP in a reaction catalysed by
14 phosphatidate cytidylyltransferase (EC 2.7.7.41) resulting in CDP-
15 diacylglycerol. Three genes have been identified. Phosphatidylserine synthase
16 then catalyses the reaction of CDP-diacylglycerol with serine to give
17 phosphatidylserine (EC 2.7.8.8); one gene has been found.
18
19 Some organisms can form phosphatidylethanolamine from phosphatidylserine
20 using a decarboxylase, but such an enzyme appears to be absent from the *E.*
21 *histolytica* genome. There are, however, several candidate methyltransferases of
22 yet unknown substrate specificity, which might be able to generate
23 phosphatidylcholine from phosphatidylethanolamine.
24
25 Taken together, large portions of the pathways needed to generate the most
26 important phospholipids can be assembled from genes tentatively identified to
27 date in the *E. histolytica* genome. The first acylation of glycerol 3-phosphate to
28 lysophosphatidate remains an important gap. As *E. histolytica* could potentially
29 acquire all the necessary phospholipids from the host, the functional relevance of
30 the described biosynthetic pathways may not be high.
31

1 Finally, two additional interesting enzymes present in *E. histolytica* should be
2 mentioned. The first was previously characterised using cDNA sequences and
3 recombinant proteins as L-myo-inositol 1-phosphate synthase (EC 5.5.1.4;
4 Lohia *et al.*, 1999). This enzyme catalyses the complicated isomerisation of
5 glucose 6-phosphate to L-myo-inositol 1-phosphate. Inositol is found in
6 phosphatidylinositol and in glycosylphosphatidylinositol- (GPI-) anchors of
7 some membrane proteins, as well as playing a major role in signal transduction
8 via the secondary messenger 1,4,5-inositol trisphosphate. There are three myo-
9 inositol 1-phosphate synthase genes, all highly similar to each other and to the
10 previously sequenced cDNA.

11

12 The second is phospholipid-cholesterol acyltransferase (EC 2.3.1.43), which
13 transfers an acyl group from phospholipids such as phosphatidylcholine to
14 cholesterol giving a cholesterol ester. The genome contains 7 genes for this
15 enzyme. So far nothing is known about the importance of cholesterol esters for
16 *E. histolytica*.

17

18 4.5.2 (b) Phospholipid degradation

19 Phospholipids are degraded by phospholipases. Whereas phospholipases A1
20 (EC 3.1.1.32) and A2 (EC 3.1.1.4) cleave the acyl residues in the 1 or 2 position
21 of the glycerol core, phospholipases C (EC 3.1.4.3) and D (EC 3.1.4.4) cleave
22 at the phosphate, phospholipase C on the glycerol side, and phospholipase D on
23 the aminoalcohol side. In *E. histolytica* phospholipase A activity has been
24 implicated in virulence (Ravdin *et al.*, 1985) as it liberates toxic fatty acids and
25 lysophospholipids (Said-Fernandez and Lopez-Revilla, 1988). Phospholipases
26 A have been found in two forms, a membrane-bound Ca-dependent form active
27 at alkaline pH and a soluble Ca-independent form active at acid pH (Long-Krug
28 *et al.*, 1985; Vargas-Villarreal *et al.*, 1998). The genome encodes 11 potential
29 phospholipases A with predicted pI values between 4.8 and 8.8 and various
30 degrees of sequence similarity. In addition the *E. histolytica* genome encodes 3
31 potential phospholipases D.

1
2 Finally, there are 2 highly similar genes for phospholipases C, but these are
3 homologous to phosphatidylinositol-specific phospholipases C (EC 3.1.4.11)
4 and most likely do not cleave phosphatidylinositol or phosphatidylcholine but
5 GPI-anchors instead. So far there are no studies using individual recombinant
6 phospholipases, and it is not yet known how much these enzymes may
7 contribute to the virulence of *E. histolytica*.

8

9 **4.6 Coenzyme A Biosynthesis and Pantothenate Metabolism**

10 Analysis of the genome revealed a complete lack of known folate-dependent
11 enzymes and folate transporters, suggesting this cofactor is not utilised by *E.*
12 *histolytica*. This is at odds with a study on the nutritional requirements of *E.*
13 *histolytica* in which folate was found to be essential for growth (Diamond and
14 Cunnick, 1991). More experimental research will be needed to resolve this
15 discrepancy. Most organisms require folate as a cofactor for several reactions
16 of amino acid metabolism and for synthesis of thymidylate, a component of
17 DNA. The microsporidian *Encephalitozoon cuniculi*, which possesses the
18 smallest known eukaryotic genome, still contains a folate transporter and
19 several folate-dependent enzymes (Katinka *et al.*, 2001). In eukaryotes
20 possessing mitochondria or chloroplasts, folate is required for the formylation
21 of methionine on the initiator tRNA used for organelle protein synthesis.
22 Although *E. histolytica* possesses a mitochondrion-derived organelle, the
23 mitosome, there is no organellar genome (Leon-Avila and Tovar, 2004) and so
24 no need for organellar protein synthesis. The most important metabolic
25 consequences of the loss of folate metabolism for *E. histolytica* are therefore
26 the absence of thymidylate synthesis and methionine recycling, although it
27 remains possible that *E. histolytica* possesses folate-independent enzymes
28 carrying out these steps.

29

30 Phosphopantothenoyl-cysteine decarboxylase (EC 4.1.1.36) and
31 phosphopantothenoyl-cysteine synthetase (EC 6.3.2.5, synonymous with

1 phosphopantothenate-cysteine ligase) exist as a fusion protein in *E. histolytica*,
2 as in Bacteria and Archaea. The amino- and carboxyl-terminal domain
3 possesses decarboxylase and synthetase activity, respectively (Kupke, 2002;
4 Kupke, 2004; Kupke *et al.*, 2000; Strauss *et al.*, 2001). The role of this enzyme
5 in coenzyme A biosynthesis is not well understood in *E. histolytica* as the other
6 necessary enzymes are absent.

7

8 **4.7 Nucleic Acid Metabolism**

9 Like many protistan parasites, *E. histolytica* lacks *de novo* purine synthesis
10 (Reeves, 1984). The genome reveals that nucleic acid metabolism of *E.*
11 *histolytica* is similar to that of the other luminal parasites *G. intestinalis* and *T.*
12 *vaginalis* in lacking pyrimidine synthesis and thymidylate synthase (Aldritt *et*
13 *al.*, 1985; Wang and Cheng, 1984). In addition *E. histolytica* appears to lack
14 ribonucleotide reductase, a characteristic shared with *G. intestinalis* (Baum *et*
15 *al.*, 1989). Ribonucleotide reductase was found, however, in genomic
16 sequences of the species *E. invadens* and *E. moshkovskii*, indicating that the
17 enzyme was lost or replaced relatively recently. Among eukaryotes, the loss of
18 these areas of nucleic acid metabolism is otherwise rare. The enzymes were
19 likely lost during adaptation to living in an organic nutrient rich environment.

20

21 **4.8 Missing Pieces**

22 Several important enzymes and pathways could not be found within the genome
23 and their presumed sequence divergence from known enzymes and pathways
24 labels them as possible drug targets once they are identified. Phosphopyruvate
25 carboxylase, which reversibly converts phosphoenolpyruvate to oxaloacetate, is
26 a central enzyme of carbon metabolism in *E. histolytica* (Reeves, 1970), but
27 could not be identified. Isoprenyl-PP synthesis and aminoethylphosphonate
28 synthesis are also likely to be present but no candidate genes could be
29 identified.

30

31 **4.9 Transporters**

1 A total of 174 transporters were identified within the genome, a number
2 intermediate between the 62 transporters of *P. falciparum* and the 286
3 transporters of *S. cerevisiae* (membranetransport.org). *E. histolytica* has a
4 number of ion transporters similar to those of yeast, but fewer identifiable
5 nutrient and organellar transporters. *Plasmodium* and *Entamoeba* both have
6 reduced metabolisms and take up many complex nutrients. The higher number
7 of transporters in *Entamoeba* suggests that they may be more substrate specific
8 than the *Plasmodium* transporters or that they may have a higher level of
9 redundancy.

10

11 Since glucose transport activity has been experimentally characterised in *E.*
12 *histolytica* and glucose is thought to be the major energy source, it was
13 surprising to find no homologues of known hexose transporters in the genome.
14 Most hexose transporters belong to the sugar porter subfamily of the major
15 facilitator superfamily (TC 2.A.1.1), members of which are found in
16 prokaryotes, animals, fungi, plants, and other protists, including *D. discoideum*,
17 but no proteins of this family were found in the *E. histolytica* genome. A group
18 of candidate monosaccharide transporters found within the genome is related to
19 the glucose/ribose porter family from prokaryotes (TC 2.A.7.5). These
20 transporters consist of two related domains, and the *Entamoeba* proteins appear
21 to have the N-terminal and C-terminal domains switched relative to the
22 bacterial proteins. Functional characterisation of transporter-encoding genes
23 will be necessary for a more complete picture.

24

25 **5. THE CYTOSKELETON**

26 The eukaryotic cytoskeleton is composed of three main elements: actin
27 microfilaments, tubulin-based microtubules and intermediate filaments. Despite
28 the fact that *E. histolytica* is very motile and performs phagocytosis very
29 efficiently, its cytoskeletal components are rather simple. No genes encoding
30 homologues of intermediate filament network proteins, including keratins,
31 desmin and vimentin, have been identified in *E. histolytica*, providing further

1 evidence that these particular cytoskeletal components are rather poorly
2 conserved in evolution. In contrast, microfilament and microtubule components
3 have been readily identified.

4

5 **5.1 Actin and Microfilaments**

6 Genome information suggests that *E. histolytica* has a greater dependence than
7 other protists on an actin-rich cytoskeletal network. Microfilament proteins are
8 represented by actin and several actin-binding proteins, although there are
9 notable differences with respect to analogous proteins in other eukaryotes.
10 There are eight actin genes in the *E. histolytica* genome, in addition to six
11 others that encode divergent actins. Three divergent actins surprisingly contain
12 an extra N-terminal domain with as yet unknown functional characteristics.
13 Examples of hybrid actins are rather scarce and have been found as ubiquitin
14 fusions (Archibald *et al.*, 2003). The functional significance of these *E.*
15 *histolytica* hybrid actins is as yet unknown.

16

17 Under physiological salt concentrations, monomeric actin assembles into
18 polymers of F-actin, thus building microfilaments. Actin assembles and
19 disassembles in an extremely dynamic and highly controlled process which is
20 dependent on many different actin-binding proteins (Winder and Ayscough,
21 2005). The *E. histolytica* genome encodes homologues of actin-binding proteins
22 involved in the severing, bundling, cross-linking and capping of filamentous
23 actin. The number and variety of actin-binding proteins support the view that
24 the actin-rich cytoskeleton is very dynamic in *E. histolytica*.

25

26 Since the spontaneous polymerisation of actin monomers is inhibited by the
27 action of sequestering proteins such as thymosin β 4 and profilin, efficient actin
28 polymerisation requires the intervention of an actin polymerisation-promoting
29 factor. The best described promoting factors are the Arp2/3 complex and the
30 formin protein family.

31

1 The Arp2/3 complex is composed of two actin-related proteins (Arp2 and Arp3,
2 which act as a template for new actin filaments) and works in conjunction with
3 five additional subunits: ARPC1 -to 5 (Vartiainen and Machesky, 2004). All
4 subunits have been clearly identified in the *E. histolytica* genome, and among
5 these the Arp2 and Arp 3 subunits are the best conserved. The Arp2/3
6 complex's ability to nucleate new actin filaments is stimulated by its interaction
7 with nucleation promoting factors such as the Wiskott-Aldrich Syndrome
8 protein (WASP) or the suppressor of cAMP-receptor (SCAR) factor.
9 Surprisingly, no proteins with homology to WASP/SCAR components were
10 found in the genome, suggesting that actin nucleation depends on the activity of
11 other, as yet unidentified proteins.

12

13 In contrast, *E. histolytica* possesses six genes coding for formins, which have
14 emerged as potent regulators of actin dynamics in eukaryotic cells through their
15 ability to increase actin filament assembly (Higgs and Peterson, 2005). Formins
16 control rearrangements of the actin cytoskeleton, especially in the context of
17 cytokinesis and cell polarisation. Members of this family have been found to
18 interact with Rho-GTPases, profilin and other actin-associated proteins. The
19 precise nature of this polymerisation-accelerating activity differs from one
20 formin to another: some nucleate filaments *de novo*, some require profilin for
21 effective nucleation, while yet others seem to use filament severing as their
22 basic mechanism. However, the Formin Homology 2 Domain (FH2, comprising
23 roughly 400 amino acids) is central to formin activity (Otomo *et al.*, 2005; Xu
24 *et al.*, 2004). Actin nucleation by formins is thought to occur by stabilisation of
25 an unfavourable nucleation intermediate, possibly through FH2 domains
26 binding to monomers in the same manner that they bind to barbed ends (an
27 activity influenced by profilin). The formin homologues from *E. histolytica* all
28 contain an FH2 domain, suggesting that they are potential actin nucleation
29 factors.

30

1 Once nucleated, actin filaments are able to grow rapidly by addition of
2 monomers at their barbed ends. Filaments are regulated by several mechanisms
3 (Winder and Ayscough, 2005). Filament length is controlled by capping
4 proteins: barbed end cappers (such as capping protein and gelsolin) block
5 addition of new monomers and thus act to decrease the overall length of the
6 filament. In addition, gelsolin severs actin filaments, thereby rapidly increasing
7 actin dynamics. Actin filaments appear to be significantly shorter in *E.*
8 *histolytica* when compared with those from fibroblasts and stress fibres are not
9 formed in this amoeba. Although *E. histolytica* actin has been shown not to
10 bind DNase I (Meza *et al.*, 1983), the inferred amino acid sequence indicates
11 conservation of all the residues likely to participate in this binding event -
12 suggesting that post-translational modifications of actin monomers may prevent
13 DNase I-actin binding. It remains to be determined whether such modifications
14 of actin participate in the regulation of actin polymerisation. The genome
15 encodes multiple genes associated with filament capping and severing, as well
16 as candidates for proteins that cross-link actin filaments and thus organise them
17 into a supramolecular network. The organisation of actin into networks and
18 higher-order structures is crucial for both cell shape and function. These
19 structures can be responsible for overall cell shape and related processes, such
20 as bundle formation through α -actinin activity, for example. The arrangement
21 of actin filaments into cross-linked arrays is also mediated by proteins with
22 multiple actin-binding domains, which allows a more perpendicular
23 arrangement of actin filaments. Examples of this type of protein are the large,
24 flexible filamin dimer (Vargas *et al.*, 1996) and the spectrin tetramer. Genome
25 analysis has now identified many candidate genes for actin-binding proteins in
26 *E. histolytica*, and additional protein partners of this versatile family
27 responsible for cytoskeleton regulation are likely to emerge from curation of the
28 sequence and cellular studies of cell motility and phagocytosis in this parasite.
29

1 **5.2 Tubulins and microtubules**

2 *E. histolytica* has a lower dependence on a tubulin-based cytoskeleton than
3 most other eukaryotic cells. Protein homologues of the basic (α , β and γ)
4 tubulins are present, although other tubulins more characteristic of organisms
5 with basal bodies and flagella (e.g.: ϵ - and δ -tubulins) are absent from *E.*
6 *histolytica* (Dutcher, 2001). Nine different tubulins (grouped into multigene
7 families) exist in most eukaryotic cells. Microtubules (MTs) composed of α -
8 and β - tubulin are intranuclear in *E. histolytica* (Vayssie *et al.*, 2004), and this
9 raises the question of how such structures are modulated within the nucleus,
10 given that MT dynamics require MT nucleation-based renewal at the minus end
11 and MT capping at the plus end. Proteins involved in MT nucleation act in
12 concert with γ -tubulin (which is also intra-nuclear in *E. histolytica*), and this
13 parasite possesses at least one homologue to the Spc98 factor, a component of
14 the MT-nucleating Tub4p- γ tubulin complex. In contrast, no homologues of
15 EB1, CLIP-170, APC (all involved in MT capping) or centrins (which operate
16 at the MT organising centre) have yet been identified, suggesting that other
17 factors (or mechanical constraints within the MT) may be required in blocking
18 MT growth. *E. histolytica* does encode candidate proteins involved in MT
19 severing or chromosome segregation. All these proteins are good candidates for
20 experimental analysis of the mechanisms of intranuclear MT localisation and
21 turnover, as well as of the trafficking of tubulins between the cytoplasm and
22 nucleus.

23

24 There is little information available on the precise organisation of microtubules
25 and F-actin cytoskeleton during *E. histolytica* motility. In many eukaryotic
26 cells, F-actin-microtubule interactions can be observed in lamellipodia at all
27 stages. Interestingly, microtubules preferentially grow along actin bundles in
28 filopodia, suggesting that a physical link between the structures exists (Leung *et*
29 *al.*, 2002). Multifunctional MT-associated proteins (MAPs, like MAP1B,
30 MAP2 and plakins) are promising candidates for acting as such links, either via
31 dimerisation of MAPs with single microtubule and actin binding sites or by

1 direct bridging of the two cytoskeletons (for example via plakins, which contain
2 binding sites for both microtubules and actin within a single molecule). Plakin
3 homologues have not been identified in the *E. histolytica* genome but a MAP is
4 present. Furthermore, proteins with domains that can bind to actin (and
5 potentially to MT) have been described in *E. histolytica* - the ABP-120 gelation
6 factor, for example (Vargas *et al.*, 1996).

7

8 **5.3 Molecular motors**

9 The distribution of intracellular factors and vesicles is performed using three
10 sets of molecular transporters: myosin along microfilaments and kinesin and
11 dynein along MTs. Although *E. histolytica* is a highly motile cell, stress fibres
12 and cytoplasmic MTs have never been observed. The fluidity of the parasite's
13 cytoplasm may be related to features of its molecular motors some of which are
14 very surprising. The myosin family of actin filament-based molecular motors
15 consists of at least 20 structurally and functionally distinct classes. The human
16 genome contains nearly 40 myosin genes, representing 12 of these classes.
17 Remarkably, *E. histolytica* is the first reported instance of a eukaryote with only
18 one unconventional myosin. This myosin heavy chain (myosin IB) belongs to
19 the type I myosin family, of which 12 are present in the *Dictyostelium* genome
20 (Eichinger *et al.*, 2005).

21

22 All members of the myosin family share a common structure composed of three
23 modules: the head, neck and tail domains. The N-terminal region harbours the
24 motor unit, which uses ATP to power movement along the actin filaments. By
25 interacting with specific proteins and 'cargoes', the tail is responsible for the
26 myosin's specific function and location. In particular, the presence of an SH3
27 domain in the tail region is important for linking these myosin I molecules with
28 the endocytic machinery and the Arp2/3 complex. Protistan class I myosins are
29 able to recruit the Arp2/3 complex towards the CARMIL adapter protein and
30 Acan125. These homologous adapters consist of multiple, leucine-rich repeat
31 sequences and bear two carboxyl-terminal polyproline motifs that are ligands

1 for the myosin I SH3 domains. CARMIL has been shown to bind the Arp2/3
2 complex via an acidic motif similar to those found in WASP. In view of the fact
3 that *E. histolytica* does not have WASP homologues, the discovery of a
4 CARMIL homologue through proteomic analysis of *E. histolytica* phagosomes
5 (Marion et al, 2005) provides an important clue for understanding actin
6 nucleation in *E. histolytica*. Interestingly, myosin IB in *E. histolytica* plays a
7 structural role in the actin network, due to its ability to cross-link filaments
8 (Marion *et al.*, 2004). The cytoskeletal structuring activity of myosin IB
9 regulates the gelation state of cell cytoplasm and the dynamics of cortical F-
10 actin during phagocytosis.

11
12 The most studied myosin has been the conventional or class II myosin. This
13 double-headed molecule is composed of two heavy chains and two pairs of
14 essential and regulatory light chains. The heavy chain tail consists of an α -
15 helical, coiled coil protein able to form a parallel dimer that in turn can self-
16 associate into bipolar, thick filaments. This enables myosin II to operate in huge
17 filament arrays, which drive high speed motility. In addition to myosin IB, *E.*
18 *histolytica* also has a conventional myosin II heavy chain (very closely related
19 to its homologue in *Dictyostelium*) which has been reported to be involved in
20 crucial phases of parasite motility, surface receptor capping and phagocytosis
21 (Arhets *et al.*, 1998). *E. histolytica*'s sole isoform shapes the actin network and
22 maintains cytoskeletal integrity. Candidate genes for the regulatory and
23 essential light chain activities were also found, and these possess the EF hand
24 domains necessary for Ca^{2+} binding.

25
26 Directional transport along the MTs depends on dynein and kinesin, both MT-
27 associated motor proteins which convert the chemical energy from ATP
28 hydrolysis into movement. These motors are unidirectional and move towards
29 either the MT plus- or minus- ends (Mallik and Gross, 2004). Kinesins and
30 dyneins have been implicated in a wide range of functions - principally
31 intracellular organelle transport during interphase and spindle function during

1 mitosis and meiosis. Members of the dynein family are minus-end directed,
2 although this remains to be confirmed for a few uncharacterised, vertebrate,
3 cytoplasmic dynein heavy chains. It has not yet been reliably established that
4 the *E. histolytica* genome contains a dynein heavy chain gene, although a
5 dynein light chain gene is present: improvements in gene assembly should
6 provide us with more information on this high molecular mass protein.

7

8 Kinesins are microtubule-dependent molecular motors that play important roles
9 in intracellular transport and cell division. Even though the motor domain is
10 found within the N-terminus in most kinesins (N-type), it is located within the
11 middle or C-terminal domains in some members of the family (M-type and C-
12 type kinesins, respectively) (Asbury, 2005). The position of the motor domain
13 dictates the polarity of the movement of kinesin along the MT: whereas N- and
14 M-type kinesins are plus-end directed, the C-type kinesins are minus-end
15 directed. Humans possess 31 different kinesins and trypanosomes have more
16 than 40. The *E. histolytica* genome sequence predicts only six kinesin-encoding
17 genes (four N-type, two C-type and no M-type homologues have been found).
18 One of the N-kinesins also contains a domain homologous to the HOOK
19 protein required for the correct positioning of microtubular structures within the
20 cell (Walenta *et al.*, 2001). Bearing in mind that *E. histolytica* MTs are
21 intranuclear, the study of kinesin function and trafficking should help us
22 elucidate what is likely to be a very interesting MT functional mechanism.

23

24 **6. VESICULAR TRAFFIC**

25 The requirement for nutritional uptake from the extracellular milieu in the host
26 intestine imposes a heavy reliance on endocytic and phagocytic activities in
27 *Entamoeba* (Espinosa-Cantellano and Martínez-Palomo, 2000). Proliferating
28 trophozoites secrete a number of peptides and proteins including cysteine
29 proteases (Que and Reed, 2000) and amoebapores (Leippe, 1999) required for
30 bacterial cell killing and degradation as well as being implicated in virulence
31 (Petri, 2002). During encystation, the cells also secrete substrates used for the

1 formation of the cyst wall (Eichinger, 1997). Electron micrographic studies
2 have revealed a complex membrane organisation. The trophozoites contain
3 numerous vesicles and vacuoles varying in size and shape (Clark *et al.*, 2000;
4 Mazzuco *et al.*, 1997). Intracellular transport of both endocytosed and
5 synthesised molecules between compartments is regulated by the elaborate
6 orchestration of vesicle formation, transport, docking and fusion to the target
7 compartment (Bonifacino and Glick, 2004; Kirchhausen, 2000).

8

9 **6.1 Complexity of Vesicle Trafficking**

10 Among a number of molecules and structures involved in vesicular trafficking,
11 three types of coated vesicles, named coatamer protein (COP) I, COPII, and
12 clathrin-coated vesicles are the best characterised (Bonifacino and Glick, 2004;
13 Kirchhausen, 2000). COPI vesicles primarily mediate transport from the Golgi
14 to the endoplasmic reticulum (ER) and between the Golgi cisternae, while
15 COPII vesicles are involved in the transport from the ER to the *cis*-Golgi. The
16 clathrin-dependent pathway has a few independent routes: from the plasma
17 membrane to endosomes, from the Golgi to endosomes, and from endosomes to
18 the Golgi. It has been well established that certain subfamilies of Ras-like small
19 GTPases, widely conserved among eukaryotes, regulate both the formation of
20 transport vesicles and their docking and fusion to the target organelles. The
21 ADP-ribosylation factor (Arf) and secretion-associated Ras-related protein
22 (Sar) families of GTPases regulate the formation of COPI and COPII vesicles
23 (Memon, 2004), respectively. In contrast, the Rab family of GTPases (Novick
24 and Zerial, 1997) is involved in the targeting and fusion of vesicles to the
25 acceptor organelles together with the tethering machinery SNARE (a soluble *N*-
26 ethylmaleimide-sensitive factor attachment protein receptor) (Chen and
27 Scheller, 2001). Since individual coat proteins, small GTPases, SNAREs, and
28 their associated proteins show distinct intracellular distributions in both
29 unicellular and multicellular organisms, they are believed to play a critical role
30 in the determination of membrane trafficking specificity (Chen and Scheller,
31 2001; Munro, 2004; Novick and Zerial, 1997). It is generally believed that the

1 total number of proteins involved in the membrane traffic reflects the
2 complexity and multiplicity of its organism. The total number of the putative
3 amoebic genes encoding Arf/Sar, Rab, SNARE, and coat proteins together with
4 those from *S. cerevisiae*, *C. elegans*, *D. melanogaster*, *H. sapiens*, and *A.*
5 *thaliana*, is shown in Table 7. *E. histolytica* reveals complexity similar to yeast,
6 fly, and worm in case of Sar/Arf and SNAREs, while the number of genes
7 encoding three coat proteins (COPI, COPII, and Adapter Proteins (APs)) was
8 higher in *E. histolytica* than these organisms and comparable to that in
9 mammals and plants. In contrast, the number of Rab proteins in *E. histolytica* is
10 exceptionally high, exceeding that in mammals and plants.

11

12 **6.2 Proteins Involved in Vesicle Formation**

13 *6.2.1 COPII-coated vesicles and Sar1 GTPase*

14 COPII components were originally discovered in yeast using genetic and
15 biochemical approaches (reviewed in Bonifacino and Glick (2004)). COPII
16 vesicles mediate the transport from the ER to the Golgi and consists of three
17 major cytosolic components and a total of five essential proteins: the Sec23p-
18 Sec24p complex, the Sec13p-Sec31p complex, and the small GTPase Sar1p
19 (Barlowe *et al.*, 1994). Sar1p and Sec23p-Sec24p complex are involved in the
20 formation of the membrane-proximal layer of the coat, while Sec13p-Sec31p
21 complex mediates the formation of the second membrane-distal layer (Shaywitz
22 *et al.*, 1997). These proteins are well conserved among various organisms
23 (Table 7). *E. histolytica* encodes one each of Sar1, Sec13 and Sec31, two of
24 Sec23, and five proteins corresponding to Sec24 (Table 7). The yeast and
25 human genomes also encode multiple Sec24 isotypes (3 and 4, respectively).
26 Although Sec24 isotypes have been shown to be responsible for the selection of
27 transmembrane cargo proteins in yeast (Peng *et al.*, 2000; Roberg *et al.*, 1999),
28 the significance of the Sec24 redundancy in *E. histolytica* is not clear.
29 Additional regulatory proteins participate in COPII assembly in yeast, including
30 Sec16p, a putative scaffold protein (Espenshade *et al.*, 1995), and Sec12p, a

1 guanine nucleotide exchange factor (GEF) for Sar1p (Barlowe and Schekman,
2 1993). Homologues of Sec12p and Sec16p appear to be absent in *E. histolytica*.
3 The p24 protein is a non-essential component of vesicle formation (Springer *et*
4 *al.*, 2000) and in yeast it functions as a cargo adaptor through binding to Sec23p
5 (Kaiser, 2000; Schimmoller *et al.*, 1995). *E. histolytica* encodes four p24
6 proteins, fewer than in yeast and humans which have eight. GTPase-activating
7 protein (GAP) Sec23p is also present in *E. histolytica*; this activates the
8 intrinsic GTPase activity of Sar1p after the formation of COPII vesicle, and
9 inactivates the function of Sar1p (Yoshihisa *et al.*, 1993), resulting in the
10 uncoating of COPII vesicles.

11

12 6.2.2 COPI-coated vesicles and Arf GTPases

13 COPI-coated vesicles, which mediate transport from the Golgi to the ER and
14 between the Golgi cisternae (Kirchhausen, 2000), consist of seven proteins (α ,
15 β , β' , γ , δ , ϵ , and ζ -COP) (Hara-Kuge *et al.*, 1994). The number of proteins
16 making up the COPI coat, and thus the complexity of COPI components, varies
17 among organisms (Table 7). While human possesses two isotypes of γ -COP and
18 ζ -COP, yeast has a single gene for each. In humans, the two isotypes of γ -COP
19 and ζ -COP form three different COPI complexes ($\gamma 1/\zeta 1$, $\gamma 1/\zeta 2$, and $\gamma 2/\zeta 1$),
20 which have different intracellular distributions (Wegmann *et al.*, 2004). This
21 implies that COPI-coated vesicles are also involved in functions other than
22 Golgi-to-ER transport (Whitney *et al.*, 1995). In *E. histolytica*, the COPI
23 complex appears more heterogeneous: *E. histolytica* encodes two isotypes each
24 of γ -COP, δ -COP, and α -COP and three isotypes of β -COP. In contrast, *E.*
25 *histolytica* lacks ϵ -COP, which is known to stabilise α -COP (Duden *et al.*,
26 1998). It has been shown in yeast that all genes encoding components of COPI
27 coat except for Sec28p, the yeast ϵ -COP homologue, are essential for growth
28 (Duden *et al.*, 1998).

29

1 Recruitment of COPI to the Golgi membrane requires the association of a GTP-
2 bound GTPase called Arf (Donaldson *et al.*, 1992; Kahn *et al.*, 2006). Arf was
3 initially identified due to its ability to stimulate the ADP-ribosyltransferase
4 activity of cholera toxin A (Kahn and Gilman, 1984). To recruit the COPI coat,
5 Arfs are activated by a Sec7 domain-containing protein, Arf-GEF, which is a
6 target of a fungal metabolite brefeldin A (Helms and Rothman, 1992; Sata *et*
7 *al.*, 1998). Among Arf family proteins, Arf1 is involved in the formation of
8 COPI-coated vesicles in the retrograde transport from the Golgi to ER, and is
9 also involved in the assembly of clathrin-AP1 (see next section) on the *trans*-
10 Golgi network (TGN) (Stamnes and Rothman, 1993), clathrin-AP3 on
11 endosomes (Ooi *et al.*, 1998), and the recruitment of AP-4 to the TGN (Boehm
12 *et al.*, 2001). The specific roles of Arfs3-5 are less clear, although Arf4 and
13 Arf5 show *in vitro* activities similar to Arf1. Functional cooperativity of Arfs in
14 the vesicular formation has also been demonstrated recently. At least two of
15 four human Arf isotypes (Arf1, Arf3-5) are essential for a retrograde pathway
16 from the Golgi to the ER, in the secretory pathway from the Golgi to the TGN,
17 and in the recycling from endosomes to the plasma membrane (Volpicelli-
18 Daley *et al.*, 2005). In contrast to these Arfs, Arf6 regulates the assembly of
19 actin filaments and is involved in endocytosis on the plasma membrane
20 (Radhakrishna and Donaldson, 1997).

21

22 GTPases that share significant similarity to Arf, but do not either activate
23 cholera toxin A or rescue *S. cerevisiae* Arf mutants are known as Arls (Arf-like
24 GTPases) (Lee *et al.*, 1997a). Arl1 is involved in endosome -to-Golgi trafficking
25 (Lu *et al.*, 2001; Lu *et al.*, 2004). Other Arls (Arls 2-11) and Arf-related
26 proteins (Arp or ArfRP 1-2) have been localised to the cytosol, nucleus,
27 cytoskeleton and mitochondria (Burd *et al.*, 2004; Pasqualato *et al.*, 2002). The
28 number of Arf, Arl, and Arf-related proteins varies among organisms (Table 7).
29 Among 27 members identified in humans, only about a half dozen Arf/Arl/Arp
30 proteins, including Arf1-6 and Arl1 (Wennerberg *et al.*, 2005), have been

1 shown to function in membrane traffic (Lu *et al.*, 2001). The localisation and
2 function of the remaining Arf/Arl/Arp remained unclear.

3

4 *E. histolytica* encodes ten Arf/Arl proteins (Table 7). Only two *E. histolytica*
5 Arfs (A1 and A2) have a high percentage identity to human Arfs 1, 3, 5, and 6
6 and yeast Arfs 1-3 (57-76% identity), while the remaining eight Arf/Arl fall
7 into three groups (A4-6, B1-3, and C) and are equally divergent from one
8 another and from other organisms. Both the intracellular distributions and the
9 specific steps in vesicular trafficking mediated by these *Entamoeba* Arf/Arl
10 proteins are unknown. It is worth noting that five of these Arfs lack a conserved
11 glycine residue at the second amino acid position of the amino terminus; this
12 glycine is known to be myristylated and essential for membrane association in
13 other organisms (Randazzo *et al.*, 1995). *EhArfA4* also lacks one of the
14 conserved GTP-binding consensus regions (Box2). Similar deletion of GTP-
15 binding domains has also been observed in proteins belonging to the Rab
16 family (see section 6.3.1).

17

18 6.2.3 Clathrin-coated vesicle and its adaptor proteins

19 Clathrin-coated vesicles and pits, as demonstrated by electron microscopy, are
20 often indicative of clathrin-mediated endocytosis. However, there is no clear
21 ultrastructural evidence for their occurrence in *Entamoeba* (Chavez-Munguia *et al.*,
22 2000). Interestingly, heavy- but not light-chain clathrin is encoded in the
23 genome. Since a majority of proteins, including adaptor proteins (APs,
24 Adaptins), known to be involved in the assembly of clathrin-coated vesicles are
25 encoded in *E. histolytica*, the fundamental mechanisms and components of
26 clathrin-mediated endocytosis are probably present in this organism, but are
27 likely to be divergent from other eukaryotes. AP is a cytosolic heterotetramer
28 that mainly mediates the integration of membrane proteins into clathrin-coated
29 vesicles in the secretory and endocytic pathways (Boehm and Bonifacino, 2001;
30 Kirchhausen, 2000). AP is composed of two large, one medium, and one small

1 subunit (Keen, 1987). Four major types of AP complexes (AP1-4) have been
 2 identified (Boehm and Bonifacino, 2001; Nakatsu and Ohno, 2003). AP-2
 3 (consisting of α , β 2, σ 2, and μ 2) mediates endocytosis from the plasma
 4 membrane (Conner and Schmid, 2003; Motley *et al.*, 2003), while AP-1
 5 (γ , β 1, σ 1, and μ 1A) (Meyer *et al.*, 2000), AP-3 (δ , β 3A, σ 3, and μ 3A) (Le
 6 Borgne *et al.*, 2001; Vowels and Payne, 1998), and AP-4 (ϵ , β 4, σ 4, and μ 4)
 7 (Aguilar *et al.*, 2001), play a role in the Golgi-endosome, endosomal-lysosomal,
 8 or the Golgi/lysosome sorting pathway, respectively. AP-4, which is present
 9 only in mammals and plants (Boehm and Bonifacino, 2001), was also identified
 10 in non-clathrin-coated vesicles mediating the transport from TGN to the plasma
 11 membrane or endosomes (Hirst *et al.*, 1999). A few isotypes of AP-1 and AP-3,
 12 e.g., AP-1B (γ , β 1, σ 1, and μ 1B) and AP-3B (δ , β 3B, σ 3, and μ 3B), showed
 13 tissue specific expression (Faundez *et al.*, 1998; Folsch *et al.*, 1999). *E.*
 14 *histolytica* encodes ten large subunits (α , β , γ , δ , and ϵ), four medium
 15 subunits (one each of μ 1 and μ 2, and two μ 3), and four small subunits (ϵ 1– ϵ 4).
 16 This suggests that *E. histolytica* produces four types of AP complex, as in
 17 humans and plants.

18

19 **6.3 Proteins Involved in Vesicle Fusion**

20 *6.3.1 Rab GTPases*

21 The docking and fusion of transport vesicles to a specific target compartment
 22 requires the appropriate Rab protein. Specific interaction of a Rab with its
 23 effector molecules in conjunction with the interaction between SNAREs plays a
 24 central role in vesicle fusion (Zerial and McBride, 2001). In general, the
 25 complexity of the Rab gene family correlates with the degree of
 26 multicellularity. For example, *S. pombe*, *S. cerevisiae*, *C. elegans*, *D.*
 27 *melanogaster*, and *H. sapiens* consist of one, one, ca. 10^3 , 10^9 , and 10^{13} cells,
 28 and have 7, 11, 29, 29, and 60 Rab genes, respectively (Pereira-Leal and
 29 Seabra, 2001). It has been also shown that in multicellular organisms, Rab
 30 proteins are expressed in a highly coordinated (i.e. tissue-, organ-, or

1 developmental stage-specific) fashion (Seabra *et al.*, 2002; Zerial and McBride,
2 2001). *E. histolytica* possesses an extremely high number of Rab genes - 91
3 (Figure 6). Among its 91 Rabs only 22, including *EhRab1*, *EhRab2*, *EhRab5*,
4 *EhRab7*, *EhRab8*, *EhRab11*, *EhRab21*, and their isotypes showed >40%
5 identity to Rabs from other organisms. The 69 remaining *E. histolytica* Rab
6 proteins showed only moderate similarity (<40% identity) and represent unique,
7 presumably *Entamoeba*-specific, Rab proteins. Approximately one third of Rab
8 proteins form 15 subfamilies, including Rab1, Rab2, Rab7, Rab8, Rab11, and
9 RabC-P, each of which contains up to 9 isoforms. Interestingly, approximately
10 70% of *E. histolytica* Rab genes contain one or more introns (Saito-Nakano *et*
11 *al.*, 2005). SNARE genes are also intron-rich whereas the Sar/Arf GTPase and
12 the three coat protein genes have a low frequency of introns. The high
13 frequency of introns in the Rab and SNARE gene families may indicate the
14 presence of post-transcriptional regulation of these genes.

15

16 Although Rab proteins generally possess a CXC or CC at the carboxyl
17 terminus, twenty-five *E. histolytica* Rabs have an atypical carboxyl terminus,
18 such as CXXX, XCXX, XXCX, XXXC, or no cysteine at all. The enzyme(s)
19 involved in the lipid modification of these unusual Rab proteins remain poorly
20 understood (see 4.5.1 (a)). It is also worth noting that >20 *E. histolytica* Rab
21 lack or contain only a degenerate form of the consensus sequence for structural
22 elements such as the GTP-binding regions and the Switch I and II regions,
23 implicated in the binding to GEF, GAP, effectors, or guanine nucleotides
24 (Saito-Nakano *et al.*, 2005). These non-conventional *EhRabs* are not
25 pseudogenes since at least some of the genes are known to be expressed as
26 mRNA (Saito-Nakano *et al.*, 2001). It has been shown that neither *EhRab5* nor
27 *EhRab7A* rescued the corresponding yeast mutant (Saito-Nakano *et al.*, 2004).
28 Therefore, many, if not all, *E. histolytica* Rabs may have lost functional
29 interchangeability with their homologues in other organisms despite the
30 relatively high percentage of sequence identities. Classification and annotation

1 of the *E. histolytica* Rab proteins has been previously described (Saito-Nakano
2 *et al.*, 2005).

3

4 One of the peculiarities of *E. histolytica* Rab proteins was demonstrated by the
5 unprecedented function of *EhRab7A*, which plays an important role in the
6 transport of cysteine proteases via interaction with the retromer complex. The
7 *E. histolytica* retromer complex consists of three components, Vps26, Vps29
8 and Vps35, rather than the 4-5 found in yeast and mammals (Nakada-Tsukui *et*
9 *al.*, 2005). Homologues of Vps5, Vps17, and sorting nexins are not encoded in
10 the genome. It has been suggested that the *EhRab7A*-retromer interaction,
11 mediated by direct binding of *EhRab7A* to a unique carboxyl-terminal region of
12 Vps26, regulates intracellular trafficking of cysteine proteases, and possibly
13 other hydrolases as well, by modulating the recycling of a putative cysteine
14 protease receptor from lysosomes and phagosomes to the Golgi or post-Golgi
15 compartment (Nakada-Tsukui *et al.*, 2005).

16

17 6.3.2 SNARE and their accessory proteins

18 The final step in membrane trafficking is the fusion of a transport vesicle with
19 its target membrane, which is mediated by the SNARE family of proteins.
20 SNAREs are integral membrane proteins that are present on both donor and
21 acceptor membranes and form a stable complex to tether the two membranes. It
22 is believed that the formation of a SNARE complex pulls the vesicle and target
23 membrane together and provides the energy to drive fusion of the lipid bilayers
24 (Chen and Scheller, 2001; Chen *et al.*, 1999). In a prototypical model, a
25 SNARE complex, which consists of four helices, is formed at each fusion site
26 (Hanson *et al.*, 1997; Poirier *et al.*, 1998). For instance, the fusion of synaptic
27 vesicles with the presynaptic nerve terminus is mediated by the formation of a
28 complex comprising one helix each from syntaxin 1A (Qa-SNARE, also termed
29 target-SNARE (t-SNARE)) and VAMP2 (R-SNARE, vesicular SNARE (v-
30 SNARE)) and two helices from SNAP-25 (Qb- and Qc-SNARE).

1
2 The complexity of SNAREs has remained largely unchanged in yeast, fly, and
3 worm, but has increased remarkably in mammals and plants (Table 7)
4 indicating that although expansion of SNARE repertoires occurs, a set of core
5 SNAREs is sufficient to mediate vesicular fusion of most pathways in
6 multicellular organisms. *E. histolytica* encodes 28 putative SNAREs, 18 Q-
7 SNAREs and 10 R-SNAREs, which is comparable to the complexity to humans
8 and plants. A notable peculiarity of SNAREs in *E. histolytica* is the lack of a
9 group of proteins possessing two helices (Qb and Qc SNAREs) such as SNAP-
10 25. Thus, the prototype model of membrane tethering by a combination of four
11 helices (from Qa, R, and Qb/Qc) does not appear to be possible in this
12 organism.

13

14 A group of proteins that interact directly with the syntaxin subfamily, including
15 the prototypical member yeast Sec1p and mammalian Munc-18, are essential
16 cytosolic proteins peripherally associated with membranes (Toonen and
17 Verhage, 2003). They are presumed to be chaperones, putting syntaxins into the
18 conformations required for interaction with other SNAREs (Dulubova *et al.*,
19 1999; Yang *et al.*, 2000). Sec1/Munc-18 proteins are also conserved in *E.*
20 *histolytica* (there are 5 Sec1 genes). Two additional important components
21 involved in the recycling of fusion machinery, *N*-ethylmaleimide sensitive
22 factor (NSF) (Beckers *et al.*, 1989) and soluble NSF attachment protein (SNAP)
23 (Clary *et al.*, 1990; Mayer *et al.*, 1996) are also found in *E. histolytica*.

24

25 Other proteins involved in vesicle fusion are the saposin-like proteins
26 mentioned earlier (Section 3.3). The membrane-fusogenic activity of the *E.*
27 *histolytica* SAPLIPs may play a role in vesicle fusion (Winkelmann *et al.*,
28 2006) but how they interface with the Rab/SNARE processes remains to be
29 determined.

30

1 **6.4 Comparisons and Implications**

2 While the fundamental machinery of vesicular trafficking is conserved in *E.*
3 *histolytica*, the high activity of the endocytic and biosynthetic transport
4 pathway in this organism appears to have resulted in the dramatic expansion of
5 the Rab gene repertoire. The diversity and complexity of Rab proteins present
6 in *E. histolytica* likely reflect the vigorous dynamism of membrane transport
7 and the reliance on Rab proteins for the specificity of vesicular trafficking. The
8 high degree of Rab complexity observed in *E. histolytica* (91) has no precedent
9 in other organisms, although the incomplete genome of *Trichomonas vaginalis*
10 appears to encode 65 Rabs (Lal *et al.*, 2005) while *Dictyostelium* encodes 50
11 (Eichinger *et al.*, 2005). Rab proteins have been extensively studied in
12 *Trypanosoma brucei* and the recent completion of *T. brucei*, *T. cruzi* and
13 *Leishmania major* genomes led to identification of all Rab genes in these
14 haemoflagellates (Ackers *et al.*, 2005; Berriman *et al.*, 2005; Quevillon *et al.*,
15 2003). Among the 16 Rab present in *T. brucei*, there are only three Rab proteins
16 (RabX1-X3) that appear to be unique to kinetoplastids. *T. brucei* possesses 11
17 Rab proteins homologous to those in humans, suggesting significant
18 conservation of the Rab-dependent core endomembrane systems in
19 kinetoplastids. *Plasmodium falciparum* possesses only 11 Rab genes all of
20 which are considered orthologues of yeast and mammalian Rabs, although
21 Rab5a, 5b, and 6 revealed unique features (Quevillon *et al.*, 2003).
22 Interestingly, some of these Rabs are expressed in a stage-dependent manner
23 (Quevillon *et al.*, 2003). The comparatively small number of Rabs in these
24 protists reinforces the tremendous diversity and complexity of Rabs seen in *E.*
25 *histolytica* (Table 7).

26

27 In marked contrast to the complexity of Rab proteins in *E. histolytica*, the
28 number of SNARE proteins, the other major components of vesicular fusion, is
29 comparable to that in yeast. The apparent disparity in the number of Rab and
30 SNARE proteins suggests one of three possibilities: 1) *EhRab* proteins share a
31 single SNARE complex as an interacting partner (Huber *et al.*, 1993; Rowe *et*

1 *al.*, 2001; Torii *et al.*, 2004), 2) a majority of *EhRabs* do not require SNARE
 2 proteins for membrane fusion (Demarque *et al.*, 2002), 3) some *EhRabs* are
 3 primarily involved in cellular functions other than membrane fusion, like Arl
 4 GTPases (Burd *et al.*, 2004; Pasqualato *et al.*, 2002). Genome-wide surveys of
 5 SNAREs in other protists are not available. The three major types of coatomer
 6 protein, which are conserved in *E. histolytica*, are also conserved in
 7 kinetoplastids (Berriman *et al.*, 2005). However, in contrast to *E. histolytica*, *T.*
 8 *brucei* does not possess multiple isotypes of COPI and II components except
 9 for Sec24, which has two isotypes. *T. cruzi* encodes all four AP complexes
 10 while *L. major* and *T. brucei* lack AP-4 or AP-2, respectively, which suggests
 11 that the repertoire of AP complexes in kinetoplastids is variable and species-
 12 specific. Although low similarity of the *E. histolytica* components to either
 13 yeast or mammalian orthologues make unequivocal assignment of *Entamoeba*
 14 AP complexes challenging, tentative assignments have been made. It is likely
 15 that *E. histolytica* encodes four kinds of AP complex corresponding to APs 1-4.

16

17 **6.5 Glycosylation and Protein Folding.**

18 *6.5.1 Asparagine-linked glycan precursors.*

19 Mammals, plants, *Dictyostelium*, and most fungi synthesise asparagine-linked
 20 glycans (N-glycans) by means of a common 14-sugar precursor dolichol-PP-
 21 $\text{Glc}_3\text{Man}_9\text{GlcNAc}_2$ (Figures 7 and 8) (Helenius and Aebi, 2004). This lipid-
 22 linked precursor is made by at least fourteen glycosyltransferases, which are
 23 present in the cytosolic aspect or lumen of the ER. The reducing end of the
 24 glycan contains two N-acetylglucosamines, while nine mannoses are present on
 25 three distinct arms. Three glucoses are added to the left arm, which is the same
 26 arm that is involved in the quality control (QC) of protein folding (see next
 27 section) (Trombetta and Parodi, 2003).

28

29 *Entamoeba* is missing luminal glucosylating and mannosylating enzymes and
 30 so makes the truncated, 7-sugar N-glycan precursor dolichol-PP- $\text{Man}_5\text{GlcNAc}_2$
 31 (Figures 7 and 8) (Samuelson *et al.*, 2005). Five mannoses on this N-glycan

1 include the left arm, which is involved in the quality control of protein folding.
2 In contrast, *Entamoeba* is missing the middle and the right arms, which are
3 involved in N-glycan associated QC of protein degradation (see next section).
4 Because *Dictyostelium*, which is phylogenetically related to *Entamoeba*, makes
5 a complete 14-sugar N-glycan precursor, it is likely that *Entamoeba* has lost
6 sets of glycosyltransferases in the ER lumen (Samuelson *et al.*, 2005).
7 Similarly, secondary loss of glycosyltransferases best explains the diversity of
8 N-glycan precursors in fungi, which contain 0-14 sugars, and apicomplexa,
9 which contain 2-10 sugars (Samuelson *et al.*, 2005).

10

11 The 14-sugar N-glycan precursor of mammals, plants, *Dictyostelium*, and most
12 fungi is transferred to the nascent peptide by an oligosaccharyltransferase
13 (OST), which is composed of a catalytic peptide and 6-7 non-catalytic peptides
14 (Kelleher and Gilmore, 2006). In contrast, the *Entamoeba* OST contains a
15 catalytic peptide and just three non-catalytic peptides, while other protists (e.g.
16 *Giardia* and *Trypanosoma*) have an OST with a single catalytic peptide. This
17 reduced complexity does not likely affect the site of N-glycan addition to the
18 nascent peptides, which is NxS or NxT (the so-called sequon) (Kornfeld and
19 Kornfeld, 1985).

20

21 6.5.2 N-glycans and quality control of protein folding.

22 Protein folding in the lumen of the ER is a complex process that involves N-
23 glycan-dependent and N-glycan-independent QC systems (Helenius and Aebi,
24 2004; Trombetta and Parodi, 2003). *Entamoeba* has four of five systems
25 present in higher eukaryotes for protein folding (Figure 9).

26 1) *Entamoeba* has the minimum component parts for N-glycan-dependent QC
27 of protein folding (Helenius and Aebi, 2004; Trombetta and Parodi, 2003;
28 Banerjee, Robbins, and Samuelson, unpublished data). These include a UDP-
29 glucose-dependent glycosyltransferase (UGGT), which adds a single glucose to
30 the left arm of the N-glycans of misfolded proteins and so form
31 GlcMan₅GlcNAc₂ (Figure 7). The glucosylated N-glycan is then bound and

1 refolded by the lectin calreticulin (CRT), which is a chaperone that works with
2 a protein disulfide isomerase (PDI) to make and break disulfide bonds. A
3 glucosidase (Gls2) removes glucose from the well-folded protein, which is
4 transferred to the Golgi by a mannose-binding lectin (ERGIC-53). The
5 *Entamoeba* system is similar to that of mammals and fungi, which add glucose
6 to the $\text{Man}_9\text{GlcNAc}_2$ precursor to make $\text{GlcMan}_9\text{GlcNAc}_2$ (Figure 7).
7 Mammals have a second glucosidase to remove glucose from the
8 $\text{Glc}_3\text{Man}_9\text{GlcNAc}_2$ precursor (Figure 7).

9 2) *Entamoeba* has N-glycan-independent QC of protein folding within the
10 lumen of the ER, which includes the chaperones Hsp70 and Hsp90 (also
11 known as BiP and Grp94, respectively) (Figure 9) (Helenius and Aebi, 2004;
12 Trombetta and Parodi, 2003; Banerjee, Cui, Robbins, and Samuelson,
13 unpublished data). Also involved in this QC system are PDIs; DnaJ proteins
14 that increase the ATPase activity of Hsp70 and Hsp90; and peptidyl-prolyl cis-
15 trans isomerases (PPIases). This N-glycan-independent QC system for protein
16 folding is present in all eukaryotes (Banerjee, Cui, Robbins, and Samuelson,
17 unpublished data).

18 3) *Entamoeba* and all other eukaryotes have an N-glycan-independent system
19 for ER-associated degradation (ERAD) of misfolded proteins (Figure 9) (Hirsch
20 *et al.*, 2004; Banerjee, Cui, Robbins, and Samuelson, unpublished data). This
21 system is composed of proteins (Sec61 and Der1) that dislocate misfolded
22 proteins from the ER lumen to the cytosol. There a complex of proteins
23 (Cdc48, Npl4, and Ufd1) ubiquinate misfolded proteins, which are then
24 degraded in the proteasome. In contrast, *Entamoeba* and the vast majority of
25 eukaryotes are missing an N-glycan-dependent system of ERAD of misfolded
26 proteins (Helenius and Aebi, 2004; Trombetta and Parodi, 2003; Banerjee, Cui,
27 Robbins, and Samuelson, unpublished data). In this system, the middle arm of
28 $\text{Man}_9\text{GlcNAc}_2$ is trimmed to $\text{Man}_8\text{GlcNAc}_2$, which is recognised by a unique
29 mannose-binding lectin (EDEM) before dislocation into the cytosol for
30 degradation (Figure 9).

1 4) *Entamoeba* has a transmembrane kinase (Ire1), which recognises misfolded
2 proteins in the lumen of the ER and triggers the unfolded protein response
3 (Figure 9) (Patil and Walter, 2001; Banerjee, Cui, Robbins, and Samuelson,
4 unpublished data; and see section 7.2.2). The amoebic unfolded protein
5 response is likely to be different from those of mammals and fungi, because
6 *Entamoeba* is missing an important downstream target, which is a transcription
7 factor called Hac1.

8

9 6.5.3 Unique N-glycans.

10 Mammals make complex N-glycans in the Golgi by trimming back the
11 precursor to $\text{Man}_3\text{GlcNAc}_2$ and then adding N-acetyl glucosamine, galactose,
12 sialic acid, and fucose (Figure 8) (Hubbard and Ivatt, 1981). In each case, the
13 activated sugars (UDP-GlcNAc, UDP-Gal, CMP-sialic acid, and GDP-fucose)
14 are transferred from the cytosol to the lumen of the Golgi by a specific
15 nucleotide-sugar transporter (NST) (Hirschberg *et al.*, 1998). In turn, each
16 activated sugar is added to the N-glycans by a specific glycosyltransferase.
17 *Entamoeba* N-glycans are remarkable for two properties. First, the most
18 abundant N-glycan is unprocessed $\text{Man}_5\text{GlcNAc}_2$ (Figure 7) (Magnelli, Ratner,
19 Robbins, and Samuelson, unpublished data). This N-glycan is recognised by
20 the mannose-binding lectin Concanavalin A, which caps glycoproteins on the
21 *Entamoeba* surface (Silva *et al.*, 1975). Unprocessed $\text{Man}_5\text{GlcNAc}_2$ is also
22 recognised by the anti-retroviral lectin cyanovirin, which binds $\text{Man}_9\text{GlcNAc}_2$
23 on the surface of gp120 (Adams *et al.*, 2004; Magnelli, Ratner, Robbins, and
24 Samuelson, unpublished data). This result suggests the possibility that the anti-
25 retroviral lectin may be active against numerous protists.
26 Second, complex N-glycans of *Entamoeba*, which are built upon the same
27 $\text{Man}_3\text{GlcNAc}_2$ core as higher eukaryotes, contain just two additional sugars
28 (galactose and glucose) (Figure 9, D and H) (Magnelli, Ratner, Robbins, and
29 Samuelson, unpublished data). Galactose is added first to both arms of
30 $\text{Man}_3\text{GlcNAc}_2$, and then glucose is added to galactose. To make these complex
31 N-glycans, *Entamoeba* has NSTs for glucose (UDP-Glc) and galactose (UDP-

1 Gal) (Bredeson *et al.*, 2005). Glucose is also transferred to N-glycans during
2 the QC of protein folding in the ER, while both galactose and glucose are
3 transferred to proteophosphoglycans (PPGs) (see next section) (Moody-Haupt
4 *et al.*, 2000). Because the complex N-glycans of *Entamoeba* are unique, it is
5 possible that they may be targets of anti-amoebic antibodies.

6

7 6.5.4 O-glycans and GPI anchors.

8 The surface of *E. histolytica* trophozoites is rich in glycoconjugates as shown
9 by the ability of many lectins and carbohydrate specific antibodies to recognise
10 the cell surface (Srivastava *et al.*, 1995; Zhang *et al.*, 2002).

11 Proteophosphoglycans (PPG) constitute the major glycoconjugate of the *E.*
12 *histolytica* cell surface. PPG is anchored to the cell surface through a GPI
13 moiety (Bhattacharya *et al.*, 1992). The structure of the PPG GPI has been
14 tentatively determined (Moody-Haupt *et al.*, 2000). In most eukaryotes,
15 phosphatidylinositol (PI) is glycosidically linked to the reducing end of de-
16 acetylated glucosamine followed by three mannoses which are in turn attached
17 to the ethanolamine that links the protein to the GPI. However, the GPI anchor
18 of *E. histolytica* PPG was found to have a unique backbone that is not observed
19 in other eukaryotes, namely Gal-Man-Man-GlcN-myoinositol. The
20 intermediate and light subunits of the *E. histolytica* Gal/GalNAc lectin, among
21 other cell surface molecules, are anchored to the cell surface through GPI
22 anchors. Though the structure of the GPI anchors is not known, they are
23 thought to be functionally important (Ramakrishnan *et al.*, 2000). In humans,
24 23 genes are known to participate in the biosynthesis of GPI anchors. However,
25 only 15 of these were identified in *E. histolytica* (Vats *et al.*, 2005).

26 Interestingly, all the catalytic subunits were identified in *E. histolytica*, the
27 missing genes encoding the accessory subunits suggesting that the biosynthetic
28 pathway may not be significantly different from that in other eukaryotes. The
29 presence of the pathway was also confirmed by detecting the biochemical
30 activities of the first two enzymes - N-acetyl glucosamine transferase and
31 deacetylase. In addition, antisense inhibition of the deacetylase blocked GPI

1 anchor biosynthesis and reduced virulence of the parasite (Vats *et al.*, 2005). A
2 novel GIPL (glycosylated inositol phospholipid) was also identified in *E.*
3 *histolytica* (Vishwakarma *et al.*, 2006). Structural studies indicate that a
4 galactose residue is attached to glucosamine as the terminal sugar instead of
5 mannose. This suggests that *E. histolytica* is capable of synthesising unusual
6 GPI-containing glycoconjugates not observed in other organisms.
7 In PPG, glycans are attached to a peptide backbone by an *O*-Phosphodiester-
8 linkage (O-P glycans). The *E. histolytica* O-P-glycans have galactose at the
9 reducing end followed by a chain of glucoses. *Entamoeba invadens* also has O-
10 P-glycans on its cyst wall proteins but the reducing sugar is a deoxysugar rather
11 than galactose (Van Dellen *et al.*, 2006b). While *Dictyostelium* also has O-P-
12 glycans on glycoproteins in its spore wall, glycoproteins with O-P-glycans are
13 absent from the vast majority of animals and plants (West, 2003).

14

15 6.5.5 Significance.

16 The unique glycans of *Entamoeba* lead to three important evolutionary
17 inferences. First, much of the diversity of eukaryotic N-glycans is due to
18 secondary loss of enzymes that make the 14-sugar lipid-linked precursor, which
19 was present in the common ancestor to extant eukaryotes. Despite the truncated
20 N-glycan precursor, *Entamoeba* has conserved the relatively complex N-
21 glycan-dependent QC system for protein folding. Third, the unique N-glycans
22 and O-P-linked glycans are based upon a novel set of glycosyltransferases,
23 which are present in *Entamoeba* and remain to be characterised molecularly.

24

25 7. PROTEINS INVOLVED IN SIGNALLING

26 7.1 Phosphatases

27 The combined actions of protein kinases and phosphatases regulate many
28 cellular activities through reversible phosphorylation of proteins. These
29 activities include such basic functions as growth, motility, and metabolism.
30 Although it was once assumed that kinases played the major regulatory role, it
31 is now clear that phosphatases can also be critical participants in some cellular

1 events (Li and Dixon, 2000). There are few publications on the role of
2 phosphatases in *E. histolytica*, however, several investigators have established a
3 role for phosphatases in proliferation, and growth. Chaudhuri *et al.* (1999)
4 observed that there was an increase in phospho-tyrosine levels in serum starved,
5 growth inhibited, *E. histolytica* cultures. Upon the additional serum and
6 subsequent growth simulation, an increase in tyrosine phosphatase activity
7 occurred. These investigators also demonstrated that genistein, a tyrosine
8 kinase inhibitor, had no effect on growth, while the addition of sodium
9 orthovanadate, a phosphatase inhibitor, produced a major decrease in cell
10 proliferation. Membrane-bound and secreted acid phosphatase activities have
11 been detected in *E. histolytica* (Aguirre-Garcia *et al.*, 1997; Anaya-Ruiz *et al.*,
12 1997). The secreted acid phosphatase activity is absent from *E. dispar*
13 (Talamas-Rohana *et al.*, 1999). This secreted acid phosphatase was found to
14 have phosphotyrosine hydrolase activity, and caused cell rounding and
15 detachment of HeLa cells (Anaya-Ruiz *et al.*, 2003), suggesting that
16 phosphatase activity contributes to the virulence of the organism.

17
18 There are four families of phosphatases (Stark, 1996). Members of the PPP
19 (protein phosphatase P) family are serine/threonine phosphatases, and include
20 PP1, PP2A, and PP2B (calcineurin-like) classes. The PPM (protein
21 phosphatase M) family phosphatases also dephosphorylate serine/threonine
22 residues but are unrelated to the PPP family proteins. A third family consists of
23 protein tyrosine phosphatases (PTP) and dual phosphatases. Low molecular
24 weight phosphatases make up the fourth family. In eukaryotic cells, greater than
25 99% of protein phosphorylation is on serine or threonine residues (Chinkers,
26 2001). Human cells have about 500 serine/threonine phosphatases and 100
27 tyrosine phosphatases (Hoof van Huijsduijnen, 1998; Hunter, 1995).
28 *Saccharomyces cerevisiae* has 31 identified or putative protein phosphatases
29 (Stark, 1996). *E. histolytica* has over 100 putative protein phosphatases. Only a
30 few of these phosphatases have potential transmembrane domains. Some *E.*
31 *histolytica* phosphatases have varying numbers of leucine-rich-repeats (LRR).

1 The LRR domain is thought to be a site for protein:protein interactions (Hsiung
2 *et al.*, 2001; Kobe and Deisenhofer, 1994). LRR domains have been found in a
3 few kinases, but had not been identified in any phosphatases until recently (Gao
4 *et al.*, 2005).

5

6 7.1.1 Serine/Threonine Protein Phosphatases

7 Members of the PPP family of protein phosphatases are closely related
8 metalloenzymes, and complex with regulatory subunits. In contrast, PPM
9 family members are generally monomeric, ranging 42-61 kDa in size. By Blast
10 analysis, the serine/threonine protein phosphatases of *E. histolytica* are most
11 closely related to PPP phosphatases PP2A, PP2B, and PPM phosphatase PP2C.

12

13 7.1.1 (a) PP2A and PP2B (Calcineurin-like) serine/threonine phosphatases

14 PP2A phosphatases are trimeric enzymes consisting of catalytic, regulatory, and
15 variable subunits (Wera and Hemmings, 1995). Calcineurin is a calcium-
16 dependent protein serine/threonine phosphatase (Rusnak and Mertz, 2000).

17 Orthologues of calcineurin are widespread, from yeast to mammalian cells.

18 Calcineurin is a heterodimeric complex with catalytic (CaNA) and regulatory
19 (CaNB) subunits. CaNA ranges in size from 58-64 kDa. Its conserved domain
20 structure includes a catalytic domain, a CaNB-binding domain, a calmodulin
21 binding domain, and an autoinhibitory (AI) domain. The binding of CaNB and
22 calmodulin activates CaNA. CaNB subunit is 19 kDa, contains 4 EF hand
23 calcium binding motifs and has similarity to calmodulin. The binding of
24 calmodulin releases the autoinhibitory domain and results in activation of the
25 phosphatase. Deletion of the AI domain results in a constitutively active
26 protein. Calcineurin is specifically inhibited by cyclosporin A and FK506.

27 Cyclosporin A and FK506 first bind to specific proteins, cyclophilin A and
28 FK506BP, respectively, then bind to CaNA at the CaNB binding site.

29 Cyclophilin A has been identified in *E. histolytica* and treatment with
30 cyclosporin A decreases growth and viability (Carrero *et al.*, 2000; Carrero *et*
31 *al.*, 2004; Ostoa-Saloma *et al.*, 2000).

1
2 The *E. histolytica* genome has 51 PP2A and calcineurin-like protein
3 phosphatases. The Pfam motif that classifies proteins as PPP phosphatases is
4 Metallophos (PF00149, calcineurin-like phosphoesterase). This motif is also
5 found in a large number of proteins involved in phosphorylation, including
6 DNA polymerase, exonucleases and other phosphatases. The genome
7 annotation identifies three loci as CaNA orthologues. However, due to the
8 similarity among this family of phosphatases, it is difficult to tell by sequence
9 analyses alone those that are calcium-dependent. Identification of CaNA will
10 have to be confirmed experimentally.

11
12 Two of the PPM phosphatases contain a TPR domain (PF00515). TPR is
13 thought to be involved in protein:protein interactions (Das *et al.*, 1998).
14 Activities that have been ascribed to TPR include regulatory roles, lipid binding
15 and auto-inhibition.

16

17 7.1.1 (b) PP2C phosphatases

18 PP2C phosphatases are also widespread and are often involved in
19 terminating/attenuating phosphorylation during the cell cycle or in response to
20 environmental stresses such as osmotic and heat shock (Kennelly, 2001).
21 Thirty-five genes were identified as PP2C phosphatases. These proteins can be
22 divided into three broad categories: 1) PP2C domain only- small (235-381
23 amino acids), 2) PP2C domain only- large (608-959 amino acids), and 3) PP2C
24 with LRR domains.

25

26 7.1.2 Tyrosine phosphatases (PTP)

27 Tyrosine phosphorylation-dephosphorylation is a key regulatory mechanism for
28 many aspects of cell biology, and development (Li and Dixon, 2000). PTPs are
29 a large class of enzymes that have catalytic domains of ~300 amino acids.
30 Forty of these residues are highly conserved (Hooft van Huijsduijnen, 1998).
31 PTPs can be divided into membrane (receptor) and non-membrane (soluble)

1 PTPs (Li and Dixon, 2000). The soluble PTP group includes those that contain
2 conserved SH2, PEST, Ezrin, PDZ, or CH2 domains. Two other classes of
3 PTPs are the low molecular weight and dual phosphatases. *Saccharomyces*
4 *cerevisiae* lacks classic PTPs but does contain dual phosphatases, such as the
5 MAP kinase kinases.

6
7 *E. histolytica* has only four potential PTPs none of which are receptor PTPs,
8 (*i.e.* PTPs with recognisable transmembrane spanning regions). Two of the
9 PTPs (XM_650778, XM_645883) are 350 and 342 amino acids in length and
10 share 48% identity. Neither of these phosphatases has any other recognisable
11 conserved domain. Non-receptor type 1 PTPs are the closest match to these
12 proteins (Li and Dixon, 2000). Membrane and secreted forms of a PTP that
13 cross-react with anti human PTP1B have been reported in *E. histolytica*
14 (Aguirre-García *et al.*, 2003; Talamas-Rohana *et al.*, 1999). Both forms have an
15 apparent molecular weight of 55 kDa and disrupt host actin stress fibers.
16 However, since none of the putative PTPs identified by the genome project
17 appear to encode secreted or membrane forms it is unlikely that these loci
18 represent these previously reported PTP1B cross-reacting proteins.

19
20 A third PTP contains a protein tyrosine phosphatase like protein (PTPLA)
21 domain (PF04387). The PTPLA domain is related to the catalytic domains of
22 tyrosine kinases, but it has an arginine for proline substitution at the active site
23 (Uwanogho *et al.*, 1999). It is not yet clear whether this family of proteins
24 actually has phosphatase activity or serves some other regulatory role.

25
26 An orthologue of a low molecular weight PTP has also been identified. Low
27 molecular weight protein tyrosine phosphatases have been found in bacteria,
28 yeast, and mammalian cells (Ramponi and Stefani, 1997). They are not similar
29 to other PTPs except in the conserved catalytic domain.

1

2 *7.1.3 Dual-specificity protein phosphatases*

3 Dual specificity PTPs (DSP) can hydrolyse both tyrosine and serine/threonine
4 residues, though they hydrolyse phosphorylated tyrosine substrates 40-500 fold
5 faster (Zhang and Van Etten, 1991). In other organisms, DSPs are found
6 mostly in the nucleus and have roles in cell cycle control, nuclear
7 dephosphorylation and inactivation of MAP kinase.

8

9 The *E. histolytica* genome has 23 sequences related to DSPs. They fall into
10 three main subclasses: those with the DSP domain only, those with DSP plus a
11 variable number (1-5) of LRRs, and those with the Rhodanese homology
12 domain (RHOD; IPR001763). Rhodanese is a sulphurtransferase involved in
13 cyanide detoxification. Its active site, RHOD, is also found in the catalytic site
14 of the dual specificity phosphatase CDC25 (Bordo and Bork, 2002).

15

16 *7.1.4 Leucine Rich Repeats (LRRs)*

17 LRRs are tandem arrays of 20-29 amino acid, leucine-rich motifs. LRRs have
18 been found in a number of proteins with varied functions including enzyme
19 inhibition, regulation of gene expression, morphology and cytoskeleton
20 formation (Kobe and Deisenhofer, 1994). LRRs are thought to provide versatile
21 sites for protein:protein interaction and have been found linked to a variety of
22 secondary domains. Most LRRs form curved horseshoe-shaped structures with
23 “a parallel beta sheet on the concave side and mostly helical elements on the
24 convex side” (IPR001611).

25

26 The LRR_1 Pfam is the second most abundant Pfam domain found in the *E.*
27 *histolytica* genome (Table 3). The LRR motifs in *E. histolytica* most closely
28 resemble the LRR found in BspA (section 2.7; Davis *et al.*, 2006). Several *E.*
29 *histolytica* proteins that contain LRRs are associated with other recognised
30 domains. These include the protein phosphatases PP2C and DSP, as well as
31 protein kinase (PK), F-box (PF00646), gelsolin/villin headpiece (IPR007122),

1 DNA J (IPR001623), Band 41 (B41;IPR000299), WD-40 (IPR001680), and
2 Zinc binding (IPR000967) domains. The association of LRRs with
3 phosphatases is unusual. One published example is the phosphatase that
4 dephosphorylates the kinase Akt (Gao *et al.*, 2005). Fungal adenylate cyclases
5 have both LRR and PP2C-like domains but this is not a wide spread feature of
6 adenylate cyclases in other species (Mallet *et al.*, 2000; Yamawaki-Kataoka *et*
7 *al.*, 1989). The LRR may be a site for interaction with phosphorylated residues
8 in *E. histolytica*. This speculation is supported by the example of the Grr1
9 protein of yeast, which contains an F-box and a LRR (Hsiung *et al.*, 2001).
10 Grr1 is involved in ubiquitin-dependent proteolysis. The LRR domain of Grr1
11 binds to phosphorylated targets in the proteasome complex. Another example
12 is the fission yeast phosphatase regulatory subunit, Sds22, which also has LRRs
13 (MacKelvie *et al.*, 1995). The LRR containing phosphatases of *E. histolytica*
14 may represent fusions of regulatory and catalytic subunits.

15

16 **7.2 Kinases**

17 *7.2.1. Cytosolic kinases.*

18 Eukaryotic protein kinases are a superfamily of enzymes, which are important
19 for signal transduction and cell-cycle regulation. Six families of
20 Serine/Threonine kinases (STKs), which include AGC, Ste, CK1, CaMK,
21 CMGC, and TKL (tyrosine kinase-like), have conserved aspartic acid and
22 lysine amino acids in their active sites and phosphorylate serine or threonine on
23 target proteins (Hanks and Hunter, 1995). Tyrosine kinases (TK), which lack
24 active site lysine, phosphorylate tyrosine on target proteins. Phosphorylated
25 tyrosine is in turn recognised by Src-homology 2 (SH2) domains that are
26 present on some kinases and other proteins. All seven families of protein
27 kinases are present in metazoa and in *D. discoideum*, while plants lack TK, and
28 *S. cerevisiae* lacks both TK and TKL.

29

30 Over 150 predicted *E. histolytica* cytosolic kinases, those that lack signal
31 peptides and trans-membrane helices, can be identified, including

1 representatives of each of the seven groups of kinases (AGC, CAMK, CK1,
2 CMGC, STE, TKL, and TK) (Loftus *et al.*, 2005; Cui and Samuelson,
3 unpublished data). Two predicted *E. histolytica* TKs, which group with human
4 TKs in phylogenetic trees, contain an AAR peptide in the active site and a
5 Kelch domain at the C-terminus (Gu and Gu, 2003). Four cytosolic protein
6 kinases contain C-terminal SH2 domains, which bind phosphorylated tyrosine
7 residues. Phosphotyrosine has been identified in *E. histolytica* using specific
8 antibodies (Hernandez-Ramirez *et al.*, 2000). The thirty-five predicted
9 cytosolic *E. histolytica* TKLs include some that contain Leu-rich repeats (LRR)
10 and ankyrin repeats at their N-termini. In contrast, the vast majority of
11 *Entamoeba* cytosolic kinases lack accessory domains.

12

13 7.2.2. Receptor-kinases.

14 Five distinct families of eukaryotic proteins have an N-terminal ectoplasmic
15 domain, a single transmembrane helix, and a C-terminal cytoplasmic kinase
16 domain (Blume-Jensen and Hunter, 2001). Ire-1 transmembrane kinases, which
17 are present in *S. cerevisiae*, plants, and metazoa, detect unfolded proteins in the
18 lumen of the ER and help splice a transcription factor mRNA by means of a
19 unique C-terminal ribonuclease (Patil and Walter, 2001). Receptor tyrosine
20 kinases (RTKs), which include growth hormone and epidermal growth factor
21 (EGF) receptors, are restricted to metazoa and have a diverse set of N-terminal
22 ectoplasmic domains and a conserved C-terminal cytosolic TK (Schlessinger,
23 2000). Receptor serine/threonine kinases (RSK) of metazoa and receptor-like
24 kinases (RLKs) of plants each contain a C-terminal TKL domain (Massague *et*
25 *al.*, 2000; McCarty and Chory, 2000; Shiu and Bleecker, 2001). Phylogenetic
26 analyses suggest that plant RLKs, animal RSKs, and animal RTKs each form
27 monophyletic groups, and that plant RLKs closely resemble cytosolic TKLs of
28 animals called Pelle or IRAK (Shiu and Bleecker, 2001).

29

30 *E. histolytica* contains >80 novel receptor RSKs, each of which has a N-
31 terminal signal sequence, a conserved ectoplasmic domain, a single

1 transmembrane helix (TMH), and a cytosolic kinase domain (Beck *et al.*, 2005).
2 The largest group of *E. histolytica* RSKs has a CXXC-rich ectoplasmic domain
3 with 6 to 31 internal repeats that each contains 4 to 6 cysteine residues (Figure
4 10). Very similar CXXC-rich domains are present in the ectoplasmic domain
5 intermediate subunit of the Gal/GalNAc lectin (section 3.1.3). CXXC-rich
6 domains are also present in hypothetical secreted proteins of *E. histolytica*,
7 while cysteine-rich domains are also present in the heavy subunit of the
8 Gal/GalNAc lectin and at the cytosolic aspect of some cysteine proteases
9 (Figure 10).

10

11 Ectoplasmic domains of other large families of *Entamoeba* RSKs have one or
12 two 6-Cys domains at the N-terminus and four 6-Cys domains proximal to the
13 plasma membrane. There are no plasma membrane proteins or secreted
14 proteins with similar domains. A minority of RSKs do not contain Cys-rich
15 ectoplasmic domains. Numerous *Entamoeba* RSKs are expressed at the same
16 time, but the specific ligands for the *Entamoeba* RSKs have not been identified
17 (Beck *et al.*, 2005).

18

19 As discussed in the section on protein folding (6.5.2), *Entamoeba* has an Ire1
20 transmembrane kinase, which recognises misfolded proteins in the lumen of the
21 ER and triggers the unfolded protein response (Figure 8).

22

23 7.2.3 Significance

24 While most protists lack TK, TKL, receptor-kinases, and Ire1 *E. histolytica* has
25 all four. It is very likely that the *E. histolytica* receptor-kinases, which are
26 extensively duplicated, will have important roles in pathogenesis (Beck *et al.*,
27 2005; Okada *et al.*, 2005). Similarly, trimeric G-proteins and the associated
28 adenylyl-cyclases likely have important roles in cyst formation and virulence
29 (Coppi *et al.*, 2002; Frederick and Eichinger, 2004).

30

31 7.3 Calcium Binding Proteins

1 Ca²⁺ signaling plays a crucial role in the pathogenesis of many protozoan
2 parasites, including *E. histolytica* (Ravdin *et al.*, 1985). Many of the calcium-
3 mediated processes are carried out with the help of calcium binding proteins
4 (CaBPs). CaBPs have been identified and characterised in almost all eukaryotic
5 systems. Some of these, such as calmodulin (CaM) and troponin C, have been
6 studied extensively. A number of CaBPs have also been identified in *E.*
7 *histolytica*. Among these are two related EF-hand containing proteins, granin
8 1 and granin 2, which are likely to be localised in intracellular granules (Nickel
9 *et al.*, 2000). Another protein, URE3-BP, was shown to have a transcription
10 regulatory function (Gilchrist *et al.*, 2001). The CaM-dependent secretion of
11 collagenases from electron dense granules has been demonstrated using *E.*
12 *histolytica* lysate. However, there is as yet no direct molecular evidence for the
13 presence of CaM in *E. histolytica* (Muñoz *et al.*, 1991). The CaM-like protein
14 EhCaBP1 has four canonical EF-hand Ca²⁺ binding domains but no functional
15 similarity to CaM (Yadava *et al.*, 1997). Inducible expression of EhCaBP1
16 antisense RNA demonstrated this protein's role in actin-mediated processes
17 (Sahoo *et al.*, 2004).

18

19 Analysis of the whole genome revealed presence of 27 CaBPs with multiple
20 EF-hand calcium binding domains (Bhattacharya *et al.*, 2006). Many of these
21 proteins are architecturally very similar but functionally distinct from CaM.
22 Moreover, functional diversity was also observed among closely related CaBPs,
23 such as EhCaBP1 and EhCaBP2 (79% identical at the amino acid level;
24 Chakrabarty *et al.*, 2004). Analysis of partial EST and proteomic databases
25 combined with Northern blots and RT-PCR shows that at least one third of
26 these genes are expressed in trophozoites, suggesting that many if not all of the
27 27 are functional genes (Bhattacharya *et al.*, 2006).

28

29 What are the roles of these proteins in the context of *E. histolytica* biology? At
30 present the function of only two EhCaBPs are known, EhCaBP1 and URE3-BP.
31 The rest of the proteins are likely to be Ca²⁺ sensors involved in a number of

1 different signal transduction pathways. After binding Ca^{2+} these may undergo
2 conformational changes and the bound form then activates downstream target
3 proteins. It is not clear why *E. histolytica* would need so many Ca^{2+} sensors
4 when many other organisms do not. It is likely that with Ca^{2+} being involved in
5 many functions, some of which are localised in different cellular locations, the
6 various CaBPs may participate in different functions that are spatially and
7 temporally separated.

8

9 **8. THE MITOSOME**

10 One of the expectations for the *E. histolytica* genome project was that it would
11 identify the function of the mitochondrial remnant known as the mitosome
12 (Tovar *et al.*, 1999) or crypton (Mai *et al.*, 1999). Under the microscope
13 mitosomes are ovoid structures smaller than 0.5 μm in diameter (Leon-Avila
14 and Tovar, 2004). While it is now clear that no mitochondrial genome still
15 persists, from both genome sequencing and cellular localisation data (Leon-
16 Avila and Tovar, 2004), the protein complement of the organelle is still
17 somewhat obscure. The number of identifiable mitosomal proteins remains very
18 small and does not provide great insight into the organelle's function. Genes
19 encoding mitochondrial-type chaperonins (cpn60, hsp10 and mt-hsp70) have
20 been identified and appear to be synthesised with amino-terminal signal
21 sequences. The importation machinery has been shown to be conserved with
22 that in true mitochondria (Mai *et al.*, 1999; Tovar *et al.*, 1999) but none of the
23 proteins involved in mitosomal protein import have been identified with
24 certainty.

25

26 Other genes encoding putative mitosomal proteins include pyridine nucleotide
27 transhydrogenase (which moves reducing equivalents between NAD and
28 NADP, and acts as a proton pump (Clark and Roger, 1995); only an incomplete
29 gene is present in the assembly), and ADP/ATP transporter (Chan *et al.*, 2005),
30 a P-glycoprotein-like protein (Pgp6), and a mitochondrial type thioredoxin,
31 although the latter two are identified based largely on their amino terminal

1 extensions. The only enzymatic pathway that is normally mitochondrial in
2 location is iron-sulphur cluster synthesis. Genes encoding homologues of both
3 IscS/NifS and IscU/NifU proteins are present, but uniquely among eukaryotes
4 the *E. histolytica* homologues are not of mitochondrial origin, having been
5 acquired by distinct lateral gene transfer from an ϵ -proteobacterium (Ali *et al.*,
6 2004b; van der Giezen *et al.*, 2004). The location of these proteins appears to be
7 cytoplasmic as determined by immunofluorescence, using antibodies against
8 both the native proteins as well as detection of epitope-tagged proteins in
9 transformed *E. histolytica* (Ali and Nozaki, unpublished). The same pathway
10 has been localised to mitosomes in *Giardia* and is also retained in all other
11 organisms with remnant mitochondria. Given the apparently unique non-
12 compartmentalised nature of iron-sulphur cluster synthesis in *E. histolytica* the
13 location of the proteins needs to be confirmed by immuno-electron-microscopy;
14 such experiments are currently underway (Ali and Nozaki, unpublished). The
15 function of the *E. histolytica* mitosome therefore remains an enigma.

16

17 **9. ENCYSTATION**

18 The infectious stage of *Entamoeba histolytica*, and also that most often used for
19 diagnosis, is the quadrinucleate cyst. Because it is not possible to encyst *E.*
20 *histolytica* in axenic culture, *Entamoeba invadens*, which is a reptilian parasite,
21 has been used as a model organism for encystation (Eichinger, 2001; Wang *et*
22 *al.*, 2003). The *E. invadens* cyst wall is composed of three parts: deacetylated
23 chitin (also known as chitosan), lectins that bind chitin (e.g. Jacob and Jessie)
24 or cyst wall glycoproteins (e.g. plasma membrane Gal/GalNAc lectin), and
25 enzymes that modify chitin or cyst wall proteins (e.g. chitin deacetylase,
26 chitinase, and cysteine proteases) (Figure 11).

27

28 **9.1 Chitin synthases**

29 Chitin fibrils, which are homopolymers of β -1,4-linked N-acetyl glucosamine
30 (GlcNAc), are synthesised by chitin synthases. Chitin synthases share common
31 ancestry with cellulose synthases and hyaluronan synthase. They are

1 transmembrane proteins with a catalytic domain in the cytosol (Bulawa, 1993),
2 where UDP-GlcNAc is made into a homopolymer and is threaded through the
3 transmembrane domains into the extracellular space. In *Saccharomyces*
4 *cerevisiae*, four accessory peptides, encoded by the Chs4-7 genes, are necessary
5 for the function of its chitin synthases (Trilla *et al.*, 1999). Remarkably, the *E.*
6 *histolytica* chitin synthase 2 (EhChs2) complements a *S. cerevisiae* chs1/chs3
7 mutant and the function of EhChs2 is independent of the four accessory
8 peptides (Van Dellen *et al.*, 2006a). This result suggests the possibility that
9 chimaeras of *E. histolytica* and *S. cerevisiae* chitin synthases may be used to
10 map domains in the *S. cerevisiae* chitin synthase that interact with the accessory
11 peptides.

12

13 **9.2 Chitin Deacetylases**

14 Chitin fibrils in the cyst wall are modified by deacetylases and chitinases (see
15 section 9.3). There are two *E. invadens* chitin deacetylases, which convert
16 chitin to chitosan (Das *et al.*, 2006). Chitosan is a mixture of N-acetyl
17 glucosamine and glucosamine and so has a positive charge. It is also present in
18 spore walls of *S. cerevisiae* and in lateral walls of *Mucor* (Kafetzopoulos *et al.*,
19 1993; Mishra *et al.*, 1997). It is likely that the positive charge of chitosan
20 fibrils contributes to the binding of cyst wall proteins, all of which are acidic
21 (de la Vega *et al.*, 1997; Frisardi *et al.*, 2000; Van Dellen *et al.*, 2002b).
22 Monosaccharide analyses of the *E. invadens* cyst walls following treatment
23 with SDS to remove proteins strongly suggest that chitosan is the only sugar
24 homopolymer present (Das *et al.*, 2006).

25

26 **9.3 Chitinases**

27 *Entamoeba* species encode numerous chitinases with a conserved type 18
28 glycohydrolase domain (de la Vega *et al.*, 1997). Recombinant *Entamoeba*
29 chitinases have both endo- and exo-chitinase activities. Two other domains are
30 important in *Entamoeba* chitinases: 1) At the N-terminus is a unique 8-Cys
31 chitin-binding domain (CBD), which is also present as a single domain in *E.*

1 *histolytica* Jessie lectins (Figure 11) (Van Dellen *et al.*, 2002b). Chitinase and
2 Jessie-3 lectin bind to the *E. invadens* cyst wall by means of this 8-Cys CBD
3 (Van Dellen *et al.*, submitted). This *E. histolytica* chitinase CBD has the same
4 function as CBDs in chitinases of fungi, nematodes, insects, and bacteria, but
5 has no sequence similarity (i.e. it has arisen by convergent evolution) (Shen and
6 Jacobs-Lorena, 1999). 2) Between the CBD and chitinase domains of
7 *Entamoeba* species are low complexity sequences that contain heptapeptide
8 repeats (Ghosh *et al.*, 2000). These polymorphic repeats may be used to
9 distinguish isolates of *E. histolytica* within the same population and may be
10 able to discriminate among isolates from New and Old World (Haghighi *et al.*,
11 2003). These polymorphic repeats, which are rich in serine and resemble
12 mucin-like domains in other glycoproteins, may also be the sites for addition of
13 *O*-phosphodiester linked sugars (see section 6.5.4).

14

15 **9.4 Jacob lectins**

16 Chitin fibrils in the cyst wall of *E. invadens* are cross-linked by Jacob lectins,
17 which contain 3 to 5 unique 6-Cys CBDs (Frisardi *et al.*, 2000). *E. invadens*
18 has at least nine genes encoding Jacob lectins, and the mRNA levels from each
19 gene increase during encystation (Van Dellen *et al.*, submitted). In addition, at
20 least six Jacob lectin proteins are present in *E. invadens* cyst walls (Van Dellen
21 *et al.*, submitted). Between the CBDs, Jacob lectins have low complexity
22 sequences that are rich in serine as in the case of chitinase [5]. Jacob lectins are
23 post-translationally modified in two ways. First, they are cleaved by cysteine
24 proteinases at conserved sites in the serine- and threonine-rich spacers between
25 CBDs. Second, they have *O*-phosphodiester-linked sugars added to serine and
26 threonine residues. *O*-phosphodiester-linked glycans are also present in
27 proteophosphoglycans (PPGs) on the surface of *E. histolytica* trophozoites
28 (Moody-Haupt *et al.*, 2000).

29

30 **9.5 Gal/GalNAc lectins**

1 The Gal/GalNAc lectins present on the surface of *E. histolytica* trophozoites
2 have been described above (section 3.1) and in the literature (Mann *et al.*, 1991;
3 Petri *et al.*, 2002). Their possible role in encystation is suggested by two
4 independent experiments. First, the signal for encystation likely depends in part
5 on aggregation of *E. invadens*, which is inhibited by exogenous galactose
6 (Coppi and Eichinger, 1999). Aggregated *E. invadens* secrete catecholamines,
7 which in an autocrine manner stimulate amoebae to encyst (Coppi *et al.*, 2002).
8 Second, in the presence of excess galactose, *E. invadens* forms wall-less cysts
9 that contain four nuclei and makes Jacob lectins and chitinase (Frisardi *et al.*,
10 2000). Because *E. invadens* trophozoites have a Gal/GalNAc lectin on their
11 surface that is capable of binding sugars on Jacob lectin, and because Jacob
12 lectins have no carboxy-terminal transmembrane helix or GPI-anchor, it is
13 likely that the cyst wall is bound to the plasma membrane by the Gal/GalNAc
14 lectin.

15

16 **9.6 Summary and Comparisons**

17 Similar to the cyst wall of *Giardia*, the cyst wall of *E. invadens* is a single
18 homogeneous layer and contains a single homopolymer, chitosan (Figure 11)
19 (Frisardi *et al.*, 2000; Gerwig *et al.*, 2002; Shen and Jacobs-Lorena, 1999). In
20 contrast, *S. cerevisiae* spore walls have multiple layers and contain β -1,3-
21 glucans in addition to chitin, while *Dictyostelium* walls have multiple layers
22 and contain N-acetyl galactosamine polymers in addition to cellulose (West,
23 2003; Yin *et al.*, 2005).

24

25 Similar to *Dictyostelium* and in contrast to fungi, the vast majority of
26 *Entamoeba* cyst wall glycoproteins are released by SDS (Van Dellen *et al.*,
27 submitted; Frisardi *et al.*, 2000; West, 2003; Yin *et al.*, 2005). While some
28 *Dictyostelium* cyst wall proteins have been shown to be cellulose-binding
29 lectins, all of the proteins bound to the cyst wall of *E. invadens* have 6-Cys
30 CBDs (Jacob lectins) or 8-Cys CBDs (Jessie 3 lectin and chitinase) (Frisardi *et*
31 *al.*, 2000; Van Dellen *et al.*, 2002b; Van Dellen *et al.*, submitted). In the same

1 way that *Giardia* cyst wall protein 2 is cleaved by a cysteine proteinase, Jacob
2 lectins are cleaved by an endogenous cysteine proteinase at sites between
3 chitin-binding domains (Touz *et al.*, 2002).

4
5 Like *Dictyostelium* spore coat proteins and insect peritrophins, cysteine-rich
6 lectin domains of *E. invadens* cyst wall proteins are separated by serine- and
7 threonine-rich domains that are heavily glycosylated (Frisardi *et al.*, 2000;
8 West, 2003; Yin *et al.*, 2005; Van Dellen *et al.*, submitted). *S. cerevisiae* cyst
9 wall proteins also have extensive serine- and threonine-rich domains that are
10 heavily glycosylated (Yin *et al.*, 2005). These glycans likely protect proteins in
11 cyst walls or fungal walls from exogenous proteases. While glycoproteins of
12 the *E. invadens* cyst wall and *Dictyostelium* spore coat contain *O*-
13 phosphodiester-linked glycans, *S. cerevisiae* wall glycoproteins contain *O*-
14 glycans (Gemmill and Trimble, 1999; West *et al.*, 2005).

15
16 Like *S. cerevisiae*, *E. invadens* has enzymes in its wall that modify chitin (Yin
17 *et al.*, 2005). Similar to chitinases of *S. cerevisiae* and bacteria, *E. invadens*
18 chitinase has a CBD in addition to the catalytic domain (Kuranda and Robbins,
19 1991). It is likely that the CBD is present to localise chitinase to the cyst wall
20 (*E. invadens*) or cell wall (*S. cerevisiae*). Finally, while *E. invadens* uses
21 catecholamines as autocrines for encystation, *Dictyostelium* uses cAMP as an
22 autocrine for sporulation (Coppi *et al.*, 2002; Kriebel and Parent, 2004). An
23 important goal of future research will be to translate what is known about the *E.*
24 *invadens* cyst wall to that of *E. histolytica*.

25

26 **10. EVIDENCE OF LATERAL GENE TRANSFER IN THE *E.*** 27 ***HISTOLYTICA* GENOME**

28 Lateral (or horizontal) gene transfer (LGT) plays a significant role in
29 prokaryotic genome evolution, contributing up to ~20% of the content of a
30 given genome (Doolittle *et al.*, 2003). LGT has therefore been an important
31 means of acquiring new phenotypes, such as resistance to antibiotics and new

1 physiological and metabolic capabilities, that may permit or facilitate
2 adaptation to new ecological niches (Koonin *et al.*, 2001; Lawrence, 2005b;
3 Ochman *et al.*, 2000). More recently, data from microbial eukaryote genomes
4 suggest that LGT has also played a role in eukaryotic genome evolution,
5 particularly among protists that eat bacteria (Andersson, 2005; Doolittle, 1998;
6 Doolittle *et al.*, 2003; Lawrence, 2005a; Richards *et al.*, 2003). *Entamoeba*
7 *histolytica* lives in the human gut, an environment that is rich in
8 microorganisms and where LGT is thought to be common between bacteria
9 (Shoemaker *et al.*, 2001). The *E. histolytica* genome thus provides a nice
10 model for investigating prokaryote to eukaryote LGT. In the original genome
11 description (Loftus *et al.*, 2005) 96 putative cases of LGT were identified using
12 phylogenetic analyses of the *E. histolytica* proteome. These have now been
13 reanalysed in the light of more recently published (August 2005) eukaryotic and
14 prokaryotic genomes. This has allowed evaluation of how previous inferences
15 were influenced by the sparse sampling of eukaryotic and prokaryotic genes
16 and species available at the time of the original analysis. Sparse gene and
17 species sampling is, and is likely to remain, a very serious problem for
18 reconstructing global trees and inferring LGT (Andersson *et al.*, 2001; Richards
19 *et al.*, 2003; Salzberg *et al.*, 2001). Thus, although ecologists differ in their
20 claims for the extent of the unsampled microbial world, they all agree that those
21 species in culture, and the even smaller subset for which genome data exist,
22 represent the smallest tip of a very large iceberg.

23

24 **10.1 How Do The 96 LGT Cases Stand Up?**

25 As before (Loftus *et al.*, 2005), Bayesian and maximum likelihood distance
26 bootstrap phylogenetic analyses were used to identify putative LGT using the
27 following *ad hoc* conservative criteria: Putative LGT was inferred where either
28 no other eukaryote possessed the gene, or where the *E. histolytica* sequence was
29 grouped with bacteria and separated from other eukaryotes by at least two
30 strongly supported nodes (bootstrap support >70%, posterior probabilities
31 >0.95). In cases where tree topologies were more weakly supported but still

1 suggested a possible LGT, bootstrap partition tables were examined for
2 partitions where the *E. histolytica* sequence clustered with another eukaryote.
3 If no such partitions were found that gene was considered to be a putative LGT.
4 Table 8 lists the results of the new analyses and also gives BlastP statistics for
5 each sequence.

6
7 A total of 41 LGT remain as strongly supported as before based upon the
8 original criteria. For the remaining 55 tree topologies, support for recent LGT
9 into the *Entamoeba* lineage is not as strong as before. For 27 of these 55 trees,
10 two strongly supported nodes separating *E. histolytica* from other eukaryotes
11 has been reduced to only one well-supported node. However, close scrutiny of
12 the bootstrap partition tables for these trees revealed that, as before, there are no
13 trees in which *E. histolytica* is found together with another eukaryote. Thus,
14 LGT still remains the strongest hypothesis to explain 68 (70%) of the original
15 96 tree topologies. In a further 14 cases, the position of *E. histolytica* among
16 prokaryotes and eukaryotes was not well supported. The taxonomic sampling of
17 eukaryotes in these trees is very patchy and the trees do not depict consensus
18 eukaryotic relationships. Thus, although the trees do not fulfill the conservative
19 criteria for LGT they also do not provide strong support for the alternative
20 hypothesis, that the *E. histolytica* genes were vertically inherited from a
21 common ancestor shared with all other eukaryotes.

22
23 In nine trees *E. histolytica* either clustered with a single newly published
24 eukaryotic sequence, or such a relationship could not be ruled out. In six of
25 these nine trees *E. histolytica* and *Trichomonas vaginalis* grouped together, and
26 two trees grouped *E. histolytica* with the diatom *Thalassiosira* (for example see
27 Figure 12). Such trees are also not easy to explain within the current consensus
28 for eukaryotic relationships (Baldauf, 2003). Similar topologies have been
29 previously reported for other eukaryotes (Andersson, 2005). The explanations
30 advanced to explain the absence of the gene in other eukaryotes include
31 massive gene loss from multiple eukaryotic lineages, or LGT between the

1 eukaryotic lineages concerned. *Entamoeba* species can ingest both eukaryotes
2 and prokaryotes and it has been suggested that LGT between eukaryotes,
3 subsequent to one lineage acquiring the gene from a prokaryote, could explain
4 such peculiar tree topologies and sparse distribution (Andersson, 2005). The
5 fact that six of the nine cases recover a relationship between *Entamoeba* and
6 *Trichomonas*, whose relatives often share the same niche, is consistent with this
7 idea. In prokaryotes, recent large-scale analyses support the hypothesis that
8 species from the same environment may share a set of niche specific genes
9 (Beiko *et al.*, 2005; Mira *et al.*, 2004).

10

11 For five trees, the *E. histolytica* gene now appears to be present in eukaryotes
12 from a different taxonomic group and the analysis cannot exclude a common
13 origin for all eukaryotic sequences. Thus, for about 5% of the original 96 cases
14 the simplest explanation is no longer LGT, but vertical inheritance from a
15 common ancestor shared with other eukaryotes.

16

17 **10.2 Where Do The Genes Come From?**

18 As before, certain prokaryotic groups are favoured as the potential donors of
19 LGT genes in the *E. histolytica* genome (Loftus *et al.*, 2005). In 15 well-
20 resolved trees *E. histolytica* is recovered next to a member of the
21 Bacteroidetes/Chlorobii group. Bacteroidetes/Chlorobii are abundant members
22 of the intestinal microflora (Shoemaker *et al.*, 2001) providing plenty of
23 opportunity for LGT to occur. Members of the Bacteroidetes/Chlorobii and
24 Fusobacterium (one tree) groups are all obligate anaerobes. This bias is
25 consistent with the idea that prokaryotic and eukaryotic cohabitants of the same
26 anaerobic niche are sharing genes (Andersson *et al.*, 2001; Beiko *et al.*, 2005;
27 Lawrence, 2005a). For example, Figure 13 shows an intriguing example where
28 the *T. vaginalis* gene clusters with members of the Bacteroidetes/Chlorobii and
29 *E. histolytica* clusters with *Fusobacterium*.

30

31 **10.3 What Kinds of Gene Are Being Transferred?**

1 Most of the 68 laterally transferred genes that can be assigned to a functional
2 category encode enzymes involved in metabolism (Figure 14). This is
3 consistent with the complexity hypothesis, which posits that LGT of genes
4 involved in processing a single substrate are more likely to be transferred than
5 those genes encoding proteins that interact with many other cellular
6 components, such as ribosomal proteins for example (Jain *et al.*, 1999).
7 Mapping the LGT enzymes on the *E. histolytica* metabolic pathway (Loftus *et*
8 *al.*, 2005) indicates that LGT has affected some important pathways, including
9 iron-sulphur cluster biosynthesis, amino acid metabolism, and nucleotide
10 metabolism. Since only eight of the 68 LGT have obvious homologues in the
11 human genome, the proteins are potentially specific to the parasite and may
12 thus be worth exploring as potential drug targets. The rest of the LGT cases
13 involve hypothetical or unclassified proteins.

14

15 **11. MICROARRAY ANALYSIS**

16 Microarray-based analyses can be utilised in conjunction with genome
17 sequencing to assign functional roles to annotated genes and to clarify genomic
18 architecture. A number of groups have utilised DNA microarrays in *E.*
19 *histolytica* (made from random genomic DNA fragments or long or short
20 oligonucleotides based on annotated genes) to successfully study transcriptional
21 differences between virulent and avirulent *E. histolytica* as well transcriptional
22 responses to heat shock, collagen and calcium exposure, tissue invasion, and
23 cyst development (Debnath *et al.*, 2004; Gilchrist *et al.*, 2006; MacFarlane and
24 Singh, 2006; Weber *et al.*, 2006; Davis *et al.*, 2007; Ehrenkauffer *et al.*, 2007).
25 Additionally, using a genomic DNA microarray, comparative genomic
26 hybridisations (CGH) between strains and species of *Entamoeba* have been
27 performed (Shah *et al.*, 2005).

28

29 Some interesting aspects of amoebic biology have been uncovered using DNA
30 microarray based expression profiling. To investigate the hypothesis that
31 virulence determinants will be more highly expressed in virulent strains, the

1 transcriptomes of virulent and avirulent *Entamoeba* species and strains have
2 been studied. It has been confirmed that a number of known virulence
3 determinants have decreased expression in avirulent *Entamoeba* (MacFarlane
4 and Singh, 2006; Davis *et al.*, 2007). A genomic DNA microarray composed of
5 2,110 genes identified 29 genes with decreased expression in both an attenuated
6 *E. histolytica* strain (Rahman) and the avirulent *E. dispar* (strain SAW760)
7 (MacFarlane and Singh, 2006), while an oligonucleotide microarray composed
8 of 6,242 genes identified 152 genes with a higher level of expression in the
9 virulent *E. histolytica* HM-1:IMSS than in the attenuated Rahman strain (Davis
10 *et al.*, 2007). A majority of these genes are annotated as hypothetical and
11 whether these genes encode novel virulence factors will require genetic analysis
12 of their functions. A peroxiredoxin gene identified as having decreased
13 expression in *E. histolytica* Rahman has been shown to be a virulence factor
14 (Davis *et al.*, 2006), indicating that these comparisons between virulent and
15 avirulent strains are likely to be a fruitful avenue of investigation.

16

17 In other microarray based studies, the large family of transmembrane receptor
18 kinases identified in *E. histolytica* has been found to be differentially expressed
19 under *in vitro* trophozoite culture conditions (Beck *et al.*, 2005). One can easily
20 envision that these kinases may have roles in signaling, allowing the parasite to
21 adapt to its ever changing environmental milieu. A substantial transcriptional
22 response to heat shock has been demonstrated (Weber *et al.*, 2006), and
23 interestingly lectin gene family members were identified as being differentially
24 regulated under heat shock conditions.

25

26 The most comprehensive microarray data to date used a whole genome short
27 oligonucleotide microarray (based on the Affymetrix platform) to profile the
28 transcriptional changes that occur as the parasite colonises and invades the host
29 colon (Gilchrist *et al.*, 2006). Using a mouse model of colitis, in which the
30 microscopic features replicate human disease and substantial pathology can be
31 seen, the transcriptional response of parasites was assayed soon after

1 colonisation (1 day after injection into the caecum) and in a long-term (29 days)
2 disease state. Overall, 326 genes were modulated at day 1 after infection, 109
3 at 29 days after infection, and 88 at both time points. A number of the well-
4 characterised “virulence determinants” in *E. histolytica* were highly expressed
5 under all conditions tested and not transcriptionally modulated, although some
6 members of the cysteine proteinase gene family were highly regulated during
7 tissue invasion. A summary of the genes and gene families that have been
8 identified as being transcriptionally active under the conditions mentioned
9 above are listed in Table 9.

10

11 The life cycle of *E. histolytica* involves transition between the trophozoite
12 stage, responsible for colonisation as well as invasive disease and the cyst,
13 responsible for infection transmission. Despite its central role, little is known
14 about cyst development in *E. histolytica*, largely due to our inability to generate
15 *E. histolytica* cysts in axenic culture. Using a whole genome microarray and
16 axenic cultures of recently isolated *E. histolytica* strains that contained
17 spontaneously produced cysts, a cyst transcriptome was developed that
18 identified 1,439 developmentally regulated genes (672 cyst-specific and 767
19 trophozoite-specific genes; Ehrenkaufner *et al.*, 2007). This first large-scale
20 insight into encystation indicates that ca. 15% of *E. histolytica* genes are
21 transcriptionally controlled in this developmental pathway. Among the genes
22 identified were a number of stage-specific cysteine proteinases, transmembrane
23 kinases, transcriptional regulators, and other potential initiators of the
24 developmental cascade. Future characterisation of these genes and pathways
25 will provide important insights into developmental processes in this parasite.

26

27 The above microarray studies used expression data to identify interesting genes
28 and pathways potentially involved in amoebic pathogenesis or development. In
29 another application of microarrays, comparative genomic hybridisations (CGH)
30 identified a number of interesting genomic characteristics of *Entamoeba* (Shah
31 *et al.*, 2005). The *E. histolytica* genome project revealed that a large number of

1 genes are multi-copy or members of highly similar gene families. Due to the
2 repetitive nature of the genome there has been difficulty with genome assembly
3 and thus the large number of gene duplications could have represented an
4 assembly artifact. The data from CGH confirmed the high copy number of a
5 significant portion (ca. 14%) of the genome and validated the genome
6 assembly. Additionally, genome-wide genetic diversity was demonstrated
7 among strains of *E. histolytica* (Shah *et al.*, 2005) including the observation that
8 the attenuated *E. histolytica* strain Rahman had a unique genetic pattern
9 suggesting the possibility that a genomic signature may correlate with invasive
10 potential. Since genome sequencing for different *E. histolytica* strains,
11 including clinical isolates, is unlikely the promise of CGH to study genetic
12 diversity and identify genotype-phenotype associations is substantial.

13

14 *E. dispar*, the closely related but avirulent species, had been identified early on
15 as having some genetic divergence from the virulent *E. histolytica*. CGH
16 analysis of *E. histolytica* and *E. dispar* revealed a significant amount of
17 difference between the two species. Whether the genetic drift in these genes is
18 responsible for the non-invasive phenotype of *E. dispar* is not known, but the
19 work has highlighted a number of genes for further functional analyses.

20

21 Taken together the DNA microarray analyses of *Entamoeba* have been useful to
22 begin to dissect the genome of this parasite and provide functional context to
23 the genes identified in the genome sequencing effort. Future directions will
24 include analysis of the parasite transcriptome in invasive hepatic disease as well
25 as further characterisation of the developmental conversion to the cyst form.
26 Those data may be useful in the development of novel diagnostic and
27 therapeutic options. Additionally, genetic approaches can now be applied to
28 definitively assign a role for these genes in amoebic biology and pathogenesis.

29

30 **12. FUTURE PROSPECTS FOR THE *E. HISTOLYTICA* GENOME**

1 Although the genome of *E. histolytica* is not yet complete it has already
2 revealed much about the biology of the parasite. There appear to be forces
3 acting to compact the genome, leading to a reduction in the coding region and
4 intron length of genes, and resulting in the loss of numerous metabolic
5 pathways. However, there are also opposing evolutionary forces as many gene
6 families have expanded. This applies particularly to genes involved in signaling
7 and trafficking that allow the parasite to sense and respond to its environment, a
8 necessary adaptation for a predatory protist. Unfortunately, it is difficult at
9 present to understand the genome structure on a macro scale due to the
10 fragmented nature of the current assembly. In other parasites, genome structure
11 has been vital to unraveling important biological processes, such as antigenic
12 variation in *T. brucei* and identification of rifin genes in *P. falciparum*. Until
13 the *E. histolytica* genome is complete we will not know what else remains to be
14 uncovered. Efforts are already underway to complete the genome by first
15 generating a HAPPY map (Dear and Cook, 1993). Over 2000 markers are being
16 designed at approximately 25 kb intervals across all contigs. Using PCR, co-
17 segregation analysis allows the identification of contigs that are physically
18 linked in the genome. This will allow the ordering and orientation of the contigs
19 and will facilitate gap closure. Shotgun genome sequencing projects of *E.*
20 *invadens* and *E. dispar* are underway (Loftus and Hall, 2005). At present the *E.*
21 *invadens* genome appears to assemble with fewer problems than were
22 encountered with that of *E. histolytica*. It is anticipated that an essentially
23 complete *E. invadens* genome sequence will be obtained, enabling extensive
24 comparative analyses to be made, and facilitating the study of pathogenicity,
25 host interaction and the evolutionary forces acting on the genome.

26

27 **ACKNOWLEDGEMENTS**

28 This review is dedicated to Louis S. Diamond without whose pioneering studies
29 the sequencing of the genome would have been impossible. The genome
30 sequencing was supported by grants from the Wellcome Trust (064057) and the
31 National Institute of Allergy and Infectious Disease (5R01AI046516-03).

1 UCMA is supported by a Wellcome Trust project grant (075796) awarded to
2 TME and RPH. Analyses by MH and MD were supported by grant P15960
3 from the Austrian Science Fund, Vienna, Austria. Analyses by CM and AL
4 were supported by grant 1RO3 TW007314-01 from the Fogarty International
5 Center of the National Institutes of Health, USA. Analyses by NG were
6 supported by an INCO-DEV grant in the fifth framework program of the
7 European Union. Analyses by IB were supported by the Deutsche
8 Forschungsgemeinschaft (DFG), BR 1744/7-1. NH would like to thank Lynn
9 Schriml and Regina Cer for preparing the web pages with the Tables and ML
10 would like to thank Julia Winkelmann for preparing table 6.

11

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1 **Table 1. Genome summary statistics for selected single celled**
 2 **organisms with sequenced genomes.**

3

	<i>Entamoeba histolytica</i>	<i>Plasmodium falciparum</i>	<i>Dictyostelium discoideum</i>	<i>Saccharomyces cerevisiae</i>	<i>Encephalitozoon cuniculi</i>
Genome Size (Mb)	23.7	22.8	33.8	12.5	2.5
G+C content (%)	24.1	19.4	22.5	38	45.5
Gene number	9,938	5268	12,500	5538	1,997
Av. gene size (bp)	1167	2,534	1,756	1428	1077
% coding DNA	49.2	52.6	ND	70.5	ND
Av. protein size (aa)	389	761	518	475	359
Av. intergenic dist. (kb)	0.8	1.7	0.8	0.6	0.1
Gene density (kb per gene)	1.9	4.3	2.5	2.2kb	1.1
% Genes with introns	25.2	54	69	5	<1
Av. intron size (bp)	102.1	179	146	ND	-
Av. number of introns/gene	1.5	2.6	1.9	1	1

4

1 **Table 2. Summary properties of the repeated DNAs.**

2 (a)

Type	Size (kb)	Estimated copy no. from genome sequence (Ref)	Estimated copy no. per haploid genome from hybridisation (Ref)	Transcript size (kb)(Ref)
EhLINE1	4.8	142 (1) 409; 49 full-length(2)	140 (3)	No full-length transcript (4)
EhLINE2	4.72	79 (1) 290; 56 full-length (2)	Not Determined	Not Determined
EhLINE3	4.81	12 (1) 52; 3 full-length(2)	Not Determined	Not Determined
EhSINE1	0.5-0.6	219 (1) 272; 81 full-length(2) 214; >90 full-length (3)	500	0.7 (6)
EhSINE2	0.65	120 (1) 117; 62 full-length(2) 122; ~50 full-length (3)	Not Determined	0.75 (7)
EhSINE3	0.58	1 (1,2)	Not Determined	Not detected (3)
Tr	0.7	1 per rDNA episome (5)	Not Determined	0.7 (5)
BspA-like	0.96	77 (8)	190 (3)	Not detected(3) ^a
Ehssp1	0.9-1.1	Not Determined	306 (9)	1.5 (9)

3

4 (b)

Family 16	GTAATGAATATAYA ACTAAGAATTTTCATTTAAAATGRATATG
Family 17	CAACAAATAAATRGKTTCAATAAAATA

5

6 (a) References for data: (1): (Van Dellen *et al.*, 2002a), (2): (Bakre *et al.*,
7 2005), (3): This analysis, (4) Bakre and Bhattacharya, unpublished
8 observations; (5): (Burch *et al.*, 1991), (6): (Cruz-Reyes *et al.*, 1995), (7): Shire
9 and Ackers, submitted, (8): (Davis *et al.*, 2006), (9): (Satish *et al.*, 2003).^a -
10 although no transcript was detected the protein has been demonstrated on the
11 cell surface and in Western blots using antibodies (Davis *et al.*, 2006).

- 1 (b) Consensus sequences of Family 16 and 17 repeats. Standard abbreviation
- 2 for degenerate sequence positions are used: R= purine, Y= pyrimidine, K= G or
- 3 T.

1 **Table 3. Number and ranking of Pfam domains across different genomes.**

		EH		EC		PF		SC		AT		CE		DD	
		#	Rank	#	Rank	#	Rank	#	Rank	#	Rank	#	Rank	#	Rank
WD40	WD domain, G-beta repeat	249	1	139	1	287	2	414	1	1137	3	694	1	719	2
LRR_1	Leucine Rich Repeat	131	2	40	2	55	12	43	17	3793	2	494	5	372	4
Pkinase	Protein kinase domain	95	3	27	5	78	8	116	2	839	4	405	8	225	7
HEAT	HEAT repeat	70	4	13	15	44	17	114	3	220	17	162	26	108	12
efhand	EF hand	58	5	7	28	80	7	29	25	422	8	213	20	153	9
RRM_1	RNA recognition motif.	57	6	30	3	95	6	86	6	375	10	223	19	134	10
Ras	Ras family	46	7	9	22	13	44	25	28	78	68	66	76	126	11
TPR_1	Tetratricopeptide repeat	42	8	23	7	48	15	103	4	334	12	180	22	168	8
Ank	Ankyrin repeat	34	9	6	34	55	12	61	9	431	6	629	2	446	3
PUF	Pumilio-family RNA binding repeat	33	10	8	23	15	34	51	13	142	32	75	68	34	62
RhoGAP	RhoGAP domain	27	11	2	118	1	520	11	80	9	559	31	138	45	39
Myb_DNA-binding	Myb-like DNA-binding domain	22	12	15	12	10	62	21	34	424	7	30	141	55	26
RhoGEF	RhoGEF domain	22	12	1	230	0	1215	3	366	0	2581	34	130	47	37
Helicase_C	Helicase conserved C-terminal domain	20	14	28	4	64	11	74	8	150	31	98	49	84	20
DEAD	DEAD/DEAH box helicase	20	14	22	9	49	14	59	10	103	50	76	67	48	35
PH	PH domain	19	16	1	230	5	123	25	28	22	255	77	63	94	16
Metallophos	Calcineurin-like phosphoesterase	19	16	6	34	16	32	21	34	66	83	78	62	31	67
Gelsolin	Gelsolin repeat	18	18	2	118	2	295	4	255	33	169	12	323	29	68
LIM	LIM domain	17	19	0	703	0	1215	8	116	16	341	103	47	56	25
CH	Calponin homology (CH) domain	16	20	4	54	1	520	7	137	26	211	57	87	49	33
Filamin	Filamin/ABP280 repeat	16	20	0	703	1	520	0	1842	2	1450	55	91	10	203

18 Columns labeled “#” give the total number of occurrences of a particular domain. Columns labeled “Rank” give the ranking of the domain where
19 the most common domain is ranked 1. The organisms shown are *Entamoeba histolytica* (EH), *Encephalitozoon cuniculi* (EC), *Plasmodium*
20 *falciparum* (PF), *Arabidopsis thaliana* (AT), *Saccharomyces cerevisiae* (SC), *Dictyostelium discoideum* (DD)..

1 **Table 4. Family C1-like cysteine endopeptidases of *E. histolytica*.**
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4	Protein	Previous	Accession No.	Protein length	Active site	Conserved motifs	Remarks
5	Name	designation		Total (pre,pro,mature)	residues		
6							
7	EhCP-A1	EhCP1	XP_650156	315 (13,80,222)	QCHN	ERFNIN, DWR	
8	EhCP-A2	EhCP2	XP_650642	315 (13,80,222)	QCHN	ERFNIN, DWR	
9	EhCP-A3	EhCP3	XP_653254	308 (13,79,216)	QCHN	ERFNIN, DWR	
10	EhCP-A4	EhCP4	XP_656602	311 (20,73,218)	QCHN	ERFNIN, DWR	
11	EhCP-A5	EhCP5	XP_650937	318 (20,72,225)	QCHN	ERFNIN, DWR, RGD	Degenerate in <i>E. dispar</i>
12	EhCP-A6	EhCP6	XP_657364	320 (17,79,224)	QCHN	ERFNIN, DWR	
13	EhCP-A7	EhCP8	XP_648996	315 (13,80,222)	QCHN	ERFNIN, DWR	
14	EhCP-A8	EhCP9	XP_657446	317 (15,82,220)	QCHN	ERFNIN, DWR	
15	EhCP-A9	EhCP10	XP_655675	297 (17,90,190)	QCHN	ERFNIN, DWR	
16	EhCP-A10	EhCP17	XP_651147	420 (18,148,254)	QCHN	ERFNIN, DWR	
17	EhCP-A11	EhCP19	XP_651690	324 (17,79,228)	QC IN ^a	ERFNIN, DWR	
18	EhCP-A12	new	XP_653823	317 (14,83,220)	(d)	ERFNIN, DWR	
19	EhCP-B1	EhCP7	XP_651581	426 (15,106,305)	QCHN	ERFNIN, PCNC	hydrophobic C-terminus
20	EhCP-B2	EhCP11	AAO03568	431 (15,106,310)	QCHS ^a	ERFNIN, PCNC	GPI cleavage site
21	EhCP-B3	EhCP12	XP_656747	474 (16,107,351)	QCHN	ERFNIN, PCNC	TMH:444-466 aa
22	EhCP-B4	EhCP13	XP_648501	379 (16,105,258)	QCHN	ERFNIN, PCNC	TMH or GPI cleavage site
23	EhCP-B5	EhCP14	XP_652671	434 (12,108,314)	QCHN	ERFNIN, PCNC	GPI cleavage site
24	EhCP-B6	EhCP15	XP_652465	300 (14,55,231)	QCHN	PCNC	hydrophobic C-terminus
25	EhCP-B7	EhCP16	XP_650400	650 (18,144,488)	QCHN	ERFNIN, PCNC	hydrophobic C-terminus, Cys-rich profile
26	EhCP-B8	EhCP18	XP_651049	473 (15,105,353)	QCHN	ERFNIN, PCNC, RGD	GPI cleavage site
27	EhCP-B9	EhCP112	XP_652993	446 (19,112,315)	QCHN	ERFNIN, PCNC, RGD	hydrophobic C-terminus, Cys-rich profile
28	EhCP-B10	new	XP_648306	372 (b)	QCHN	ERFNIN, PCNC, RGD	hydrophobic C-terminus
29	EhCP-B11	new	XP_648013	133 (b)	Q ???	PCNC	
30	EhCP-C1	new	XP_654453	586 (c)	QCIN ^a	HS(X) ₆ I ₁ CP	TMH:12-34
31	EhCP-C2	new	XP_656632	567 (c)	QCHN	HS(X) ₆ I ₁ CP	TMH:27-49
32	EhCP-C3	new	XP_655128	572 (c)	QCHN	HS(X) ₆ L ₁ CP	TMH:17-39
33	EhCP-C4	new	XP_655800	502 (c)	QCHN	LT(X) ₆ L ₁ CP	
34	EhCP-C5	new	XP_654800	557 (c)	QCHN	IS(X) ₆ I ₁ CP	TMH:20-42
35	EhCP-C6	new	XP_651553	557 (c)	QCHD ^a	HS(X) ₆ L ₁ CA	TMH:14-36
36	EhCP-C7	new	XP_657273	595 (c)	QCHN	IS(X) ₆ L ₁ CP	TMH:19-41
37	EhCP-C8	new	XP_655479	627 (c)	QCHN	IS(X) ₆ I ₁ CP	TMH:29-51

1	EhCP-C9	new	XP_655011	518 (c)	(d)	HS(X) ₆ ICP	TMH:12-34
2	EhCP-C10	new	XP_654829	530 (c)	QCHN	IS(X) ₆ ICP	TMH:15-37
3	EhCP-C11	new	XP_648083	526 (c)	(d)	HS(X) ₆ ICP	TMH:20-42
4	EhCP-C12	new	XP_650829	473 (c)	(d)	MS(X) ₆ LCG	TMH:26-48 & 449-471
5	EhCP-C13	new	XP_656556	564 (c)	QCHN	VS(X) ₆ RCG	TMH:21-43
6							

7 a - active sites that lack the canonical motif QCHN; b - incomplete sequence; c - cleavage sites to be determined; d - not conserved

1 **Table 5. Family C2-, C19-, C54-, and C65-like Cysteine endopeptidases of *E. histolytica***

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4 Name	Homology	Family	ProteinID	Protein length	Active site
6 EhCALP1	Calpain-like	C2	XP_649922	591 aa	not cons.
7 EhCALP2	Calpain-like	C2	XP_657312	473 aa	QCHN
8 EhUBHY	Ubiquitin Hydrolase-like	C19	XP_657356	444 aa	NDTN
9 EhAUTO1	Autophagin-like	C54	XP_651386	325 aa	YCHS
10 EhAUTO2	Autophagin-like	C54	XP_653798	364 aa	YCHD
11 EhAUTO3	Autophagin-like	C54	XP_652043	364 aa	YCHD
12 EhAUTO4	Autophagin-like	C54	XP_656724	348 aa	YCHD
13 EhOTU	Otubain-like	C65	XP_654013	259 aa	DCH

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1 **Table 6. Attributes of the identified SAPLIPs of *E. histolytica***

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Name	Size, aa			SAPLIP domain		Identical to annotated protein	SAPLIP domain can be found within this sequence ^g	Similar to (aa sequence identity, %) ^f	Homologous proteins in other organisms (aa sequence identity, %)
	entire	signal peptide (predicted) ^a	proform / mature ^c	position aa residues	similarity ^e name, Acc. no.				
Amoebapore A	98	21 ^b	77	22 - 98	SAPOSIN B, IPR 008139	amoebapore A precursor ACCESSION XP_653265		Amoebapore A Acc. AAA29111 (100%)	Disparpore A Acc. AAA18632 <i>E.dispar</i> (94%)
SAPLIP 1	92	15	77	16 - 92	SAPOSIN B, IPR 008139	saposin-like protein ACCESSION XP_655836		Amoebapore A Acc. AAA29111 (64%)	Disparpore A Acc. AAA18632 <i>E.dispar</i> (68%)
Amoebapore B	96	19 ^b	77	20 - 96	SAPOSIN B, IPR 008139	Pore-forming peptide amoebapore B precursor (EH-APP) ACCESSION Q24824		Amoebapore B Acc. CAA54226 (100%)	Disparpore B Acc. AAF04195 <i>E.dispar</i> (90%)
Amoebapore C	101	24 ^b	77	25 - 101	SAPOSIN B, IPR 008139	amoebapore C ACCESSION XP_656029		Amoebapore C Acc. CAA54225 (100%)	Disparpore C Acc. AAF04196 <i>E.dispar</i> (88%)
SAPLIP 2	153	15	138	71 - 153	SAPOSIN B, IPR 008139	hypothetical protein ACCESSION XP_656037		-	-
SAPLIP 3	94	16	78	18 - 94	SAPOSIN B, IPR 008139	hypothetical protein ACCESSION XP_656682		Amoebapore A Acc. AAA29111 (30%)	Invapore X Acc. AAP80381 <i>E.invadens</i> (67%)
SAPLIP 4	96	17	79	18 - 96	SAPOSIN B, IPR 008139	hypothetical protein ACCESSION XP_652159 AND hypothetical protein ACCESSION XP_652303		Amoebapore C Acc. CAA54225 (27%)	Disparpore C Acc. AAF04196 <i>E.dispar</i> (30%)
SAPLIP 5	1026	18	1008	946 - 1026	SAPOSIN B, IPR 008139	chromosome partition protein ACCESSION XP_655789		-	-
SAPLIP 6	92	15	77	14 - 92	SAPOSIN B, IPR 008139	hypothetical protein ACCESSION XP_655820		-	-
SAPLIP 7	926	17	909	855 - 926	SAPOSIN B, IPR 008139	conserved hypothetical protein ACCESSION XP_656441		-	-
SAPLIP 8	980	15	965	902 - 980	SAPOSIN B, IPR 008139	hypothetical protein ACCESSION XP_656913		-	-
SAPLIP 9	140	15	125	61 - 140	SAPOSIN B, IPR 008139	hypothetical protein ACCESSION XP_650376		-	-
SAPLIP 10	657	16	641	577 - 657	SAPOSIN B, IPR 008139	-	Genomic survey sequence ACCESSION AZ687176	-	-
SAPLIP 11	693	17	676	615 - 693 ^d	-	-	Genomic survey sequence ACCESSION AZ692153	-	-
SAPLIP 12	873	16	857	793 - 873	SAPOSIN B, IPR 008139	hypothetical protein ACCESSION XP_652721		-	-
SAPLIP 13	1009	no signal peptide predicted	1009	931 - 1005	SAPOSIN B, IPR 008139	hypothetical protein ACCESSION XP_655089		-	-
SAPLIP 14	915	17	898	834 - 915	SAPOSIN B, IPR 008139	-	Genomic survey sequence ACCESSION AZ690015	-	-

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7 SAPLIPs were named according to the similarity of their SAPLIP domain to amoebapore A:

8 ^a by the programme SignalP and manually corrected if predicted cleavage site is within the SAPLIP domain9 ^b verified by experimental data10 ^c with the exception of amoebapores it is not possible to decide whether proteins are further processed

- 1 ^d identified manually
- 2 ^e extracted from InterPro databases
- 3 ^f if no similarity is reported, there is none outside of the SAPLIP domain
- 4 ^g sequences only found in GSS section of GenBank with given identifier

1 **Table 7. The number of genes encoding representative proteins involved in vesicular trafficking in *E. histolytica*.**

Protein	<i>E. histolytica</i>	<i>S. cerevisiae</i>	<i>C. elegans</i>	<i>D. melanogaster</i>	<i>H. sapiens</i>	<i>A. thaliana</i>	References
Sar1	1	1	1	1	2	4	1,2
COPII	9	6	5	4	9	12	1
Arf	10	6	11	11	25	17	3
COPI	11	7	7	7	9	9	1
AP-1		5	7	5	8	9	
AP-2		4	5	5	5	6	
AP-3		4	4	4	7	4	
AP-4		0	0	0	4	4	
AP total	18	13	16	14	24	23	1
Rab	91	11	29	26	60	57	1,4
Qa	8	7	9	7	12	18	
Qb	10(b+c)	6	7	5	9	11	
Qc		8	4	5	8	8	
R	10	5	6	5	9	14	
SNARE total	28	24	23	20	35	54	1,5,6
NSF	1	1	1	2	1	1	1,7
SNAP	1	1	1	3	1	3	
Sec1	5	4	6	5	7	6	8

2 References (1), Bock *et al.* (2001); (2), Wennerberg *et al.* (2005); (3), Pasqualato *et al.* (2002); (4), Pereira-Leal and Seabra (2001); (5), Burri
3 and Lithgow (2004); (6), Uemura *et al.* (2004); (7), Sanderfoot *et al.* (2000); (8) Boehm *et al.* (2001).

1 Table 8. Reassessment of the 96 candidate LGT cases identified in the original genome publication

Acc. ^a RefSeq	Acc. ^a	EhL ^b	Top Prokaryotic blast hit	PL ^c	%ID ^d	Top Eukayotic blast hit	EL ^c	%ID ^d	PE-score ^e	EE-score ^f	P/E Ratio ^g
41 LGT cases that remain strongly supported according to our criteria											
EAL43201	XP_648590.1	487	<i>Treponema denticola</i>	507	57	<i>Trichomonas vaginalis</i>	398	43	1.00E-167	5.00E-88	2.00E-80
EAL43619	XP_649008.1	621	<i>Vibrio vulnificus</i>	673	41	<i>Saccharomyces cerevisiae</i>	664	40	1.00E-132	1.00E-125	1.00E-07
EAL43678	XP_649067.1	538	<i>Fusobacterium nucleatum</i> <i>Mannheimia</i>	562	47	<i>Trichomonas vaginalis</i>	477	34	1.00E-135	2.00E-60	5.00E-76
EAL43850	XP_649240.1	880	<i>succiniciproducens</i>	898	63	<i>Mastigamoeba balamuthi</i>	882	45	0	0	N/A
EAL44182	XP_649570.1	260	<i>Bacteroides thetaiotaomicron</i>	273	34	<i>Yarrowia lipolytica</i>	298	29	2.00E-35	4.00E-10	5.00E-26
EAL44226	XP_649612.1	262	<i>Bacteroides thetaiotaomicron</i>	267	28	<i>Tetrahymena thermophila</i>	1476	30	2.00E-25	0.11	1.82E-24
EAL44778	XP_650165.1	188	<i>Bacteroides thetaiotaomicron</i>	188	43	<i>Neurospora crassa</i>	546	34	8.00E-41	1.8	4.44E-41
EAL45076	XP_650453.1	358	<i>Bacteroides fragilis</i>	362	46	<i>Trichomonas vaginalis</i>	562	22	1.00E-87	0.24	4.17E-87
EAL45145	XP_650531.1	825	<i>Staphylococcus aureus</i>	1036	30	<i>Trichomonas vaginalis</i>	2468	20	3.00E-59	0.016	1.88E-57
EAL45220	XP_650606.1	479	<i>Clostridium tetani</i>	471	45	<i>Arabidopsis thaliana</i>	581	31	1.00E-114	1.00E-54	1.00E-60
EAL44744	XP_650131.1	160	<i>Bacteroides fragilis</i>	424	41	<i>Yarrowia lipolytica</i>	169	31	3.00E-24	7.00E-11	4.29E-14
EAL46110	XP_651498.1	157	<i>Bacteroides fragilis</i>	166	49	<i>Arabidopsis thaliana</i>	627	35	5.00E-35	3.2	1.56E-35
EAL45378	XP_650765.1	311	<i>Haloarcula marismortui</i>	299	43	<i>Leishmania major</i>	411	43	3.00E-54	1.00E-32	3.00E-22
EAL45618	XP_651004.1	159	<i>Bacteroides thetaiotaomicron</i>	157	46	<i>Plasmodium vivax</i>	1275	33	2.00E-28	0.69	2.90E-28
EAL46311	XP_651697.1	248	<i>Synechococcus elongates</i>	270	36	<i>Trichomonas vaginalis</i>	3075	18	1.00E-30	0.38	2.63E-30
EAL46679	XP_652065.1	218	<i>Methanosarcina mazei</i>	230	37	<i>Candida glabrata</i>	461	24	8.00E-31	0.079	1.01E-29
EAL46975	XP_652361.1	370	<i>Bordetella bronchiseptica</i>	368	46	<i>Cryptococcus neoformans</i>	372	40	8.00E-83	3.00E-71	2.67E-12
EAL47525	XP_652912.1	380	<i>Clostridium perfringens</i>	296	23	<i>Plasmodium falciparum</i>	390	34	2.00E-13	1.3	1.54E-13
EAL47905	XP_653291.1	227	<i>Clostridium perfringens</i>	259	33	<i>Tetrahymena thermophila</i>	1425	24	4.00E-19	0.32	1.25E-18
EAL48587	XP_653973.1	425	<i>Desulfovibrio vulgaris</i>	442	60	<i>Yarrowia lipolytica</i>	572	37	1.00E-149	9.00E-57	1.11E-93
EAL48979	XP_654365.1	732	<i>Thermotoga neapolitana</i>	740	40	<i>Cryptococcus neoformans</i>	735	28	1.00E-135	3.00E-64	3.33E-72
EAL49084	XP_654474.1	350	<i>Methanococcus jannaschii</i>	241	29	<i>Anopheles gambiae</i>	784	40	1.00E-24	5.00E-06	2.00E-19
EAL49209	XP_654596.1	247	<i>Bacteroides fragilis</i>	243	38	<i>Thalassiosira pseudonana</i>	269	22	7.00E-43	0.0002	3.50E-39
EAL49277	XP_654665.1	737	<i>Bacteroides thetaiotaomicron</i>	781	31	<i>Cryptococcus neoformans</i>	935	24	1.00E-111	6.00E-44	1.67E-68
EAL49613	XP_654999.1	168	<i>Sulfolobus solfataricus</i>	237	34	<i>Tetrahymena thermophila</i>	487	38	1.00E-16	6.00E-06	1.67E-11
EAL49813	XP_655200.1	186	<i>Escherichia coli</i>	200	31	<i>P. brasiliensis</i>	257	26	2.00E-13	0.47	4.26E-13
EAL49869	XP_655257.1	390	<i>Campylobacter jejuni</i>	407	56	<i>Ashbya gossypii</i>	490	39	1.00E-124	8.00E-73	1.25E-52
EAL50263	XP_655646.1	390	<i>Porphyromonas gingivalis</i>	408	48	<i>Yarrowia lipolytica</i>	428	38	1.00E-98	3.00E-60	3.33E-39
EAL50440	XP_655826.1	344	<i>Bacillus anthracis</i>	491	54	<i>Rhizopus oryzae</i>	510	40	1.00E-101	2.00E-67	5.00E-35
EAL50508	XP_655888.1	348	<i>Wolinella succinogenes</i>	340	55	<i>Mus musculus</i>	168	40	1.00E-106	2.00E-18	5.00E-89
EAL50603	XP_655988.1	567	<i>Bacteroides thetaiotaomicron</i>	622	45	<i>Trichomonas vaginalis</i>	632	39	1.00E-141	2.00E-99	5.00E-43 3.33E-
EAL50801	XP_656185.1	499	<i>Bacteroides thetaiotaomicron</i>	513	52	<i>Trichomonas vaginalis</i>	514	28	1.00E-145	3.00E-40	106
EAL50992	XP_656375.1	140	<i>Archaeoglobus fulgidus</i>	184	40	<i>Trichomonas vaginalis</i>	195	46	1.00E-27	0.018	5.56E-26

EAL50997	XP_656380.1	656	<i>Bacteroides thetaiotaomicron</i>	718	53	<i>Cryptococcus neoformans</i>	770	32	0	2.00E-69	0.00E+00
EAL51149	XP_656535.1	343	<i>Bacteroides fragilis</i>	359	43	<i>Pichia ofunaensis</i>	378	34	8.00E-84	1.00E-53	8.00E-31
EAL51236	XP_656622.1	259	<i>Symbiobacterium thermophilum</i>	274	45	<i>Oryza sativa</i>	315	21	3.00E-51	0.003	1.00E-48
EAL51348	XP_656749.1	171	<i>Methanopyrus kandleri</i>	204	37	<i>Tetrahymena thermophila</i>	2872	22	3.00E-21	0.1	3.00E-20
EAL51525	XP_656903.1	316	<i>Bacteroides thetaiotaomicron</i>	300	29	<i>Candida boidinii</i>	314	32	8.00E-27	0.0007	1.14E-23
EAL51565	XP_656946.1	415	<i>Clostridium perfringens</i>	900	43	<i>Trichomonas vaginalis</i>	897	40	1.00E-89	5.00E-81	2.00E-09
EAL51925	XP_657304.1	448	<i>T. tengcongensis</i>	481	43	<i>Giardia lamblia</i>	937	33	3.00E-96	2.00E-60	1.50E-36
EAL52001	XP_657387.1	303	<i>Oceanobacillus iheyensis</i>	306	27				2.00E-15	0.00E+00	

27 LGT cases that are more weakly supported than before according to our criteria

EAL45152	XP_650539.1	122	<i>Shewanella oneidensis</i>	132	34	<i>Trypanosoma brucei</i>	385	24	5.00E-10	6.6	7.58E-11
EAL43347	XP_648734.1	848	<i>Burkholderia pseudomallei</i>	779	38	<i>Plasmodium falciparum</i>	2463	32	1.00E-136	4.00E-44	2.50E-93
EAL44257	XP_649643.1	407	<i>Clostridium acetobutylicum</i>	406	25	<i>Homo sapiens</i>	468	24	6.00E-23	1.00E-14	6.00E-09
EAL45586	XP_650972.1	460	<i>Clostridium tetani</i>	476	47	<i>Xenopus laevis</i>	513	38	1.00E-116	5.00E-84	2.00E-33
EAL46313	XP_651699.1	118	<i>Prochlorococcus marinus</i>	163	42	<i>Hordeum vulgare</i>	223	22	2.00E-21	1.4	1.43E-21
EAL46399	XP_651785.1	218	<i>Clostridium perfringens</i>	235	65	<i>Trypanosoma brucei</i>	295	52	3.00E-73	9.00E-54	3.33E-20
EAL46421	XP_651808.1	205	<i>Clostridium acetobutylicum</i>	230	40	<i>Arabidopsis thaliana</i>	241	33	7.00E-34	6.00E-12	1.17E-22
EAL46701	XP_652087.1	294	<i>Bacteroides fragilis</i>	308	45	<i>Thalassiosira pseudonana</i>	348	27	4.00E-63	1.00E-14	4.00E-49
EAL46757	XP_652143.1	95	<i>Lactococcus lactis</i>	103	31	<i>Tetrahymena thermophila</i>	112	32	3.00E-09	1.00E-07	3.00E-02
EAL46858	XP_652245.1	192	<i>Pseudomonas aeruginosa</i>	195	41	<i>Caenorhabditis briggsae</i>	229	40	6.00E-36	2.00E-17	3.00E-19
EAL47026	XP_652397.1	164	<i>Bacillus subtilis</i>	181	30	<i>Trichomonas vaginalis</i>	182	26	3.00E-10	2.00E-08	1.50E-02
EAL47464	XP_652839.1	504	<i>Treponema denticola</i>	509	39	<i>Piromyces sp.</i>	555	27	5.00E-88	2.00E-30	2.50E-58
EAL47648	XP_653034.1	259	<i>Methanosarcina mazei</i>	272	36	<i>Arabidopsis thaliana</i>	345	25	2.00E-39	3.00E-11	6.67E-29
EAL47787	XP_653173.1	546	<i>Spirochaeta thermophila</i>	571	56	<i>Solanum tuberosum</i>	552	46	1.00E-175	1.00E-135	1.00E-40
EAL48186	XP_653572.1	232	<i>Bacillus cereus</i>	279	34	<i>Thalassiosira pseudonana</i>	271	32	2.00E-10	2.00E-08	1.00E-02
EAL49309	XP_654698.1	358	<i>Methanosarcina mazei</i>	379	42	<i>Leishmania major</i>	373	31	5.00E-77	9.00E-44	5.56E-34
EAL48568	XP_653954.1	113	<i>Chlamydia pneumoniae</i>	271	38	<i>Debaryomyces hansenii</i>	699	38	5.00E-14	7.00E-16	7.14E+01
EAL48767	XP_654156.1	165	<i>Bacteroides fragilis</i>	177	40	<i>Trichomonas vaginalis</i>	189	28	7.00E-28	2.00E-05	3.50E-23
EAL48783	XP_654172.1	217	<i>Pseudomonas putida</i>	225	46	<i>Giardia lamblia</i>	239	35	2.00E-43	7.00E-24	2.86E-20
EAL49703	XP_655090.1	396	<i>Clostridium acetobutylicum</i>	398	34	<i>Tetrahymena thermophila</i>	445	29	4.00E-64	3.00E-44	1.33E-20
EAL49996	XP_655383.1	358	<i>Bacteroides thetaiotaomicron</i>	368	60	<i>Brachydanio rerio</i>	367	43	1.00E-121	5.00E-76	2.00E-46
EAL50325	XP_655711.1	447	<i>Clostridium tetani</i>	448	30	<i>Trichomonas vaginalis</i>	871	29	4.00E-46	1.00E-37	4.00E-09
EAL50521	XP_655905.1	285	<i>Streptococcus agalactiae</i>	323	29	<i>Leishmania major</i>	452	24	2.00E-22	3.00E-06	6.67E-17
EAL50620	XP_656005.1	261	<i>Wolinella succinogenes</i>	655	27	<i>Trichomonas vaginalis</i>	261	28	6.00E-21	1.00E-06	6.00E-15
EAL50838	XP_656225.1	299	<i>Anabaena sp.</i>	287	27	<i>Trichomonas vaginalis</i>	336	29	4.00E-15	0.0009	4.44E-12
EAL50986	XP_656369.1	219	<i>Bacteroides thetaiotaomicron</i>	240	31	<i>Xenopus laevis</i>	309	29	2.00E-20	1.00E-12	2.00E-08
EAL52121	XP_657511.1	220	<i>T. tengcongensis</i>	222	36	<i>Caenorhabditis elegans</i>	255	26	1.00E-30	1.00E-07	1.00E-23

14 cases where increased sampling has weakened that case for LGT

EAL42539	XP_647925.1	213	<i>Bacteroides thetaiotaomicron</i>	319	47	<i>Entodinium caudatum</i>	411	43	3.00E-53	1.00E-32	3.00E-21
EAL42738	XP_648124.1	313	<i>Campylobacter jejuni</i>	324	40	<i>Trichomonas vaginalis</i>	313	36	1.00E-63	4.00E-42	2.50E-22
EAL44270	XP_649657.1	179	<i>Methanococcus maripaludis</i>	193	37	<i>Anopheles gambiae</i>	186	21	2.00E-27	2.00E-09	1.00E-18

EAL44593	XP_649979.1	220	<i>Vibrio vulnificus</i>	244	24	<i>Trichomonas vaginalis</i>	238	21	0.0002	2.6	7.69E-05
EAL45320	XP_650707.1	154	<i>Geobacillus kaustophilus</i>	183	53	<i>Thalassiosira pseudonana</i>	182	43	8.00E-38	2.00E-32	4.00E-06
EAL45332	XP_650718.1	392	<i>Methanosarcina acetivorans</i>	420	48	<i>Trichomonas vaginalis</i>	396	47	8.00E-99	2.00E-93	4.00E-06
EAL45528	XP_650913.1	349	<i>Sulfolobus acidocaldarius</i>	343	28	<i>Cyanophora paradoxa</i>	313	27	1.00E-24	5.00E-17	2.00E-08
EAL45907	XP_651293.1	380	<i>Streptomyces coelicolor</i>	603	32	<i>Dictyostelium discoideum</i>	457	30	2.00E-39	2.00E-35	1.00E-04
EAL46026	XP_651412.1	176	<i>Bacteroides fragilis</i>	184	51	<i>Tetrahymena thermophila</i>	323	32	2.00E-44	8.00E-08	2.50E-37
EAL46116	XP_651488.1	662	<i>Bacillus clausii</i>	684	48	<i>Solanum tuberosum</i>	761	48	0	1.00E-172	0.00E+00
EAL46656	XP_652044.1	419	<i>Dictyoglomus thermophilum</i>	579	30	<i>S. pombe</i>	493	41	2.00E-35	2.00E-19	1.00E-16
EAL50605	XP_655990.1	392	<i>Thermotoga maritima</i>	417	38	<i>Cryptococcus neoformans</i>	445	30	2.00E-69	1.00E-33	2.00E-36
EAL51270	XP_656656.1	251	<i>Porphyromonas gingivalis</i>	261	50	<i>Anopheles gambiae</i>	272	39	6.00E-53	1.00E-35	6.00E-18
EAL52102	XP_657492.1	345	<i>Bacteroides thetaiotaomicron</i>	358	54	<i>Thalassiosira pseudonana</i>	354	47	1.00E-105	7.00E-86	1.43E-20
Nine cases where <i>Entamoeba</i> is now recovered with a recently sequenced gene from another microbial eukaryote											
EAL44213	XP_649600.1	710	<i>Bdellovibrio bacteriovorus</i>	698	37	<i>Trichomonas vaginalis</i>	713	35	1.00E-127	1.00E-127	1.00E+00
EAL44435	XP_649823.1	250	<i>Bacteroides fragilis</i>	395	40	<i>Trichomonas vaginalis</i>	395	33	1.00E-43	3.00E-35	3.33E-09
EAL44766	XP_650152.1	401	<i>Porphyromonas gingivalis</i>	419	36	<i>Trichomonas vaginalis</i>	445	32	3.00E-65	1.00E-51	3.00E-14
EAL47785	XP_653171.1	234	<i>Bacillus anthracis</i>	242	32	<i>Trichomonas vaginalis</i>	256	39	2.00E-30	3.00E-33	6.67E+02
EAL47859	XP_653246.1	337	<i>Clostridium acetobutylicum</i>	322	50	<i>C. reinhardtii</i>	352	44	9.00E-74	0	N/A
EAL49158	XP_654544.1	397	<i>T. tengcongensis</i>	412	49	<i>Trichomonas vaginalis</i>	416	46	1.00E-100	4.00E-99	2.50E-02
EAL49488	XP_654874.1	320	<i>Geobacter sulfurreducens</i>	336	34	<i>Leishmania major</i>	357	31	1.00E-38	4.00E-30	2.50E-09
EAL49791	XP_655177.1	164	<i>Oceanobacillus iheyensis</i>	177	42	<i>Thalassiosira pseudonana</i>	96	38	8.00E-30	6.00E-09	1.33E-21
EAL50404	XP_655790.1	718	<i>T. tengcongensis</i>	717	37	<i>Trichomonas vaginalis</i>	721	34	1.00E-139	1.00E-118	1.00E-21
Five cases where vertical inheritance is now the simplest explanation for the new tree											
EAL44346	XP_649732.1	314	<i>Oceanobacillus iheyensis</i>	239	47	<i>Dictyostelium discoideum</i>	278	65	1.00E-52	3.00E-95	3.33E+42
EAL45466	XP_650849.1	209	<i>Agrobacterium tumefaciens</i>	254	31	<i>Thalassiosira pseudonana</i>	227	35	3.00E-23	1.00E-27	3.00E+04
EAL45548	XP_650934.1	259	<i>Bacillus cereus (strain ZK)</i>	233	29	<i>Candida glabrata</i>	270	30	7.00E-06	5.00E-05	1.40E-01
EAL45595	XP_650981.1	284	<i>Pyrobaculum aerophilum</i>	293	27	<i>Ashbya gossypii</i>	343	27	1.00E-23	7.00E-16	1.43E-08
EAL50185	XP_655571.1	186	<i>Aeropyrum pernix</i>	192	31	<i>Thalassiosira pseudonana</i>	149	30	4.00E-13	5.00E-06	8.00E-08

1

2 All 96 trees reanalysed here can be downloaded (in pdf format) from the following web site: http://www.ncl.ac.uk/microbial_eukaryotes/

3

4 ^aGenBank accession numbers and RefSeq accession numbers, respectively, for the 96 original candidates LGT identified by phylogenetic
5 analysis (Loftus *et al.*, 2005)6 ^bEhL, the length of the *E. histolytica* protein7 ^cPL/EL, the protein length of the prokaryotic or eukaryotic top BlastP hit, respectively8 ^d%ID, the percent identity between the *E. histolytica* protein and the top prokaryotic or eukaryotic protein in BlastP alignments (in respective
9 columns)

1 °PE-score, the e-score of the top prokaryotic hit

2 °EE-score, the e-score of the top eukaryotic hit

3 °P/E Ratio, the e-score ratio between the top prokaryotic hit and top eukaryotic hit

4 Abbreviated taxon names (to fit the columns):

5 *Chlamydomonas reinhardtii*: *C. reinhardtii*; *Paracoccidioides brasiliensis*: *P. brasiliensis*; *Schizosaccharomyces pombe*: *S. pombe*;

6 *Thermoanaerobacter tengcongensis*: *T. tengcongensis*

7

1 **Table 9. Examples of microarray-detected transcriptional changes in some gene families and the conditions tested**2
3

Gene family	Total number of genes in gene family	Number of genes transcriptionally regulated under condition tested	
		Heat shock ^a (1,131 genes on array)	Host colonisation and invasion ^b (9,435 genes on array)
Cysteine proteinases	29 ^c	2 upregulated (CPs 6, 4); 7 down-regulated (CPs 1, 2, 3, 8, 13, 17,	21 genes on array; 4 up-regulated (CPs 1, 9, 4, 6);1 down-regulated (CP8)
Lectin (Heavy, Light, and Intermediate subunits)	12	1 up-regulated (Hgl-2); 5 down-regulated (Lgl-1 and 3, Igl 1 and 2, Hgl-3)	No change in heavy or intermediate subunits; Light subunit Igl2 and Igl3 down-regulated)
Amoebapore	3	1 down-regulated (amoebapore C)	No substantial changes
Transmembrane receptor kinases	>80	NA	6 up-regulated (TMKs 69, 53, 95, 105, 63, 56) 2 down-regulated (TMKs 03 and 17)
AIG-1 (similar to plant antibacterial proteins)	15	NA	5 up-regulated at day 1; 6 down-regulated at day 29 (all non-overlapping)

4
5
6

^a Adapted from Weber *et al.* (2006); ^b Adapted from Gilchrist *et al.* (2006);

^cNumber of cysteine proteinase gene families in genome annotation at time studies were performed

1 Figure Legends

2

3 *Figure 1.*

4 **Positions of introns in the vacuolar ATPase subunit D gene in *P. falciparum*, *D.***
5 ***discoideum*, and *E. histolytica***

6

7 *Figure 2. Comparison of protein sizes in *E. histolytica* and *D. discoideum*.*

8 a: The graph shows the distribution of predicted amino acid length across sequenced
9 genomes from single celled eukaryotes: *D. discoideum* (DD) *Encephalitozoon*
10 *cuniculi* (EC), *P. falciparum* (PF), *E. histolytica* (EH), and *S. cerevisiae* (SC). *E.*
11 *histolytica* and *E. cuniculi* have a distribution that is skewed toward smaller proteins
12 relative to the other species.

13 b: The histogram displays the degree of size change of genes in *E. histolytica* relative
14 to *D. discoideum* when comparing orthologous genes identified by reciprocal best
15 blast hits. The black bars show genes that are smaller in *E. histolytica* where as the
16 grey bars are smaller in *D. discoideum*.

17

18 *Figure 3. Domain diagram of the Hgl subunit of the Gal/GalNAc lectin. CW-*
19 *Cysteine-Tryptophan region; CF- Cysteine free region; C-Rich- Cysteine rich region.*
20 The black vertical box near the carboxy-terminus of the protein represents the single
21 transmembrane domain. The horizontal black bars above the diagram indicate the
22 location of a carbohydrate recognition domain (CRD), the region with similarity to
23 the hepatic growth factor receptor, **c-Met**, and the region that has similarity to the
24 **CD59**, the membrane inhibitor of the complement membrane attack complex. The
25 numbers in parentheses indicate the location of these regions in the Hgl1 isoform
26 (Mann *et al.*, 1991), where the methionine of the immature protein is residue 1.

27

28 *Figure 4. Structural domains of the 3 different types of family C1-like cysteine*
29 **endopeptidases EhCP-A, EhCP-B and EhCP-C.** Shown are the location and length
30 of domains specific for each the 3 types as well as the conserved active site and
31 cysteine residue

32

33 *Figure 5. Predicted antioxidant system of *Entamoeba histolytica*.* A. Superoxide
34 radical anions are detoxified by an iron-containing superoxide dismutase (FeSOD).

1 Molecular oxygen is reduced to hydrogen peroxide by a NADPH:flavin
 2 oxidoreductase (thioredoxin reductase, p34). Hydrogen peroxide is converted to water
 3 by rubrerythrin (Rbr). The nature of its redox partner is unknown. Hydrogen peroxide
 4 can also be converted to water via a classical thioredoxin redox system consisting of
 5 thioredoxin reductase (TrxR, p34), thioredoxin (Trx) and peroxiredoxin (Prx). B.
 6 Nitric oxide is reduced by an A-type flavoprotein (FprA) to nitrous oxide and water.
 7 For this reaction FprA receives electrons from NADH oxidase (Far).

8

9 **Figure 6. A phylogenetic tree of Rab proteins from *Entamoeba histolytica*,**
 10 **human, and yeast.** The number on the nodes represent the bootstrap proportions (%)
 11 of 1000 pseudo samples; only bootstrap proportions >30% are shown. *E. histolytica*
 12 Rab proteins are indicated in bold. Tentative subfamilies that revealed significant
 13 similarity (>40% identity) to their human or yeast counterpart are shaded dark, while
 14 *Entamoeba*-specific subfamilies have light shading. The scale bar indicates 0.1
 15 substitutions at each amino acid position. *: *EhRab* proteins that lack the conserved
 16 effector region, switch regions, or GTP-binding boxes. **: *EhRab* proteins that
 17 possess a non-conventional carboxyl-terminus or lack carboxyl-terminal cysteines.
 18 ***: Rab proteins that were not classified as isotypes based on <40% identity to other
 19 members of the subfamily. References on tree: (1), Temesvari *et al.* (1999); (2),
 20 Rodríguez *et al.* (2000); (3), Saito-Nakano *et al.* (2001); (4), Juarez *et al.* (2001); (5),
 21 Saito-Nakano *et al.* (2004); and (6), Okada *et al.* (2005).

22

23 **Figure 7. Synthesis of N-glycan precursors by *S. cerevisiae* (A) and *E. histolytica***
 24 **(B).** The N-glycan precursor of *S. cerevisiae* contains 14 sugars (Glc₃Man₉GlcNAc₂),
 25 each of which is added by a specific enzyme. The *E. histolytica* N-glycan precursor
 26 contains just seven sugars (Man₅GlcNAc₂), as the protist is missing enzymes that add
 27 mannose and glucose in the lumen of the ER. The figure is redrawn from Figure 1 of
 28 Samuelson *et al.* (2005). Glc = Glucose; GlcNAc = N-acetyl glucosamine; Man =
 29 Mannose.

30

31 **Figure 8. Selected N-glycans of mammals (A-E) and *Entamoeba* (F-H).** Precursors
 32 transferred to nascent peptide (A and F). Glycosylated products involved in N-
 33 glycan-associated QC of protein folding (B and G). Mannosidase product involved in

1 N-glycan-associated protein degradation (mammals only) (C). Trimmed product that
 2 is building block for complex N-glycans (mammals and *Entamoeba*) (D). Complex
 3 N-glycans made in the Golgi (E and H). Glc = Glucose; GlcNAc = N-acetyl
 4 glucosamine; Man = Mannose; Gal = Galactose; Fuc = Fucose.

5

6 **Figure 9. Model of quality control of protein folding in *Entamoeba*.** 1. N-glycan-
 7 dependent QC of protein folding. 2. N-glycan-independent QC of protein folding. 3.
 8 N-glycan-independent ERAD. 4. Ire1 and unfolded protein response (see text for
 9 details).

10

11 **Figure 10. Structure of cysteine-rich plasma membrane proteins of *E. histolytica*.**

12 These proteins include the various subunits of the Gal/GalNAc lectin, a cysteine
 13 protease, and numerous receptor kinases. Ire1, which is involved in the unfolded
 14 protein response, is also a receptor kinase but has no Cys-rich domain.

15

16 **Figure 11. Model for the *Entamoeba* cyst wall derived primarily from**

17 **experiments with *E. invadens*.** A. The cyst wall consists of chitosan fibrils, which
 18 are made by chitin synthase and chitin deacetylase. Wall proteins include Jacob
 19 lectins with tandem arrays of 6-Cys chitin-binding domains (CBDs), as well as
 20 chitinase and Jessie lectins that have a single 8-Cys CBD. The Gal/GalNAc lectin in
 21 the plasma membrane binds sugars on the Jacob and Jessie lectins. B. Structures of
 22 representative lectins illustrated in A.

23

24 **Figure 12. Phylogenetic relationships of *E. histolytica* glutamine synthase.** The
 25 gene encoding glutamine synthase (EC 6.3.1.2) is now shared by *E. histolytica* and the
 26 diatom *Thalassiosira*. This gene is mainly restricted to prokaryotic genomes
 27 (eukaryotes are highlighted by arrows). *T. vaginalis* also contains a homologue but in
 28 this case it clusters weakly with *Fusobacterium*. The scale bar represents 10% of
 29 inferred sequence divergence. Both the GenBank and RefSeq accession numbers are
 30 given for the *E. histolytica* entry.

31

32 **Figure 13. Phylogenetic relationships of *E. histolytica* tryptophanase.** This tree
 33 suggests that the *E. histolytica* gene encoding a tryptophanase was acquired by LGT
 34 from a relative of the anaerobic bacterium *Fusobacterium*. In contrast, the *T.*

1 *vaginalis* gene appears to have a separate origin with a LGT from a relative of the
2 anaerobic *Bacteroides* group. The scale bar represents 10% of inferred sequence
3 divergence. Both the GenBank and RefSeq accession numbers are given for the *E.*
4 *histolytica* entry. The EC number is also shown.

5

6 **Figure 14. Pie chart of functional categories for the 68 strongest LGT cases.** The
7 cases are those discussed in the text and listed in Table 8. Most entries encode
8 metabolic enzymes (KEGG annotation).

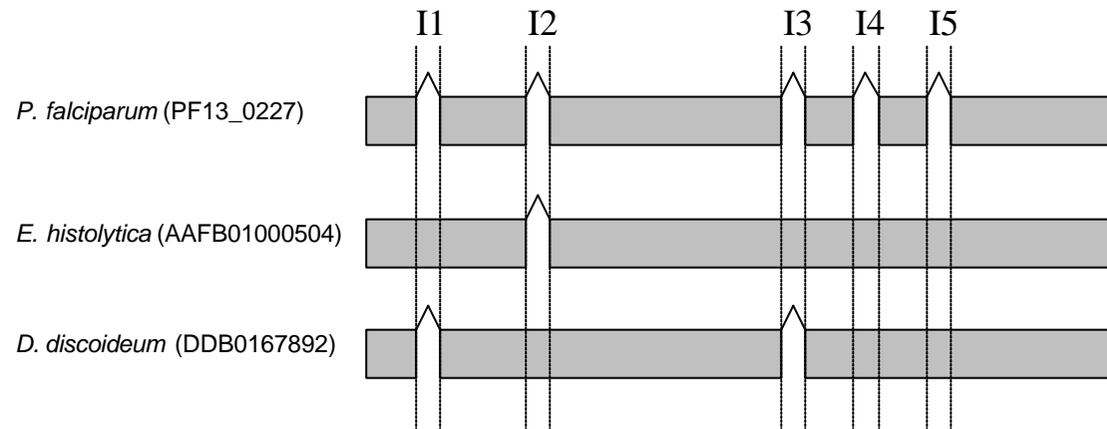
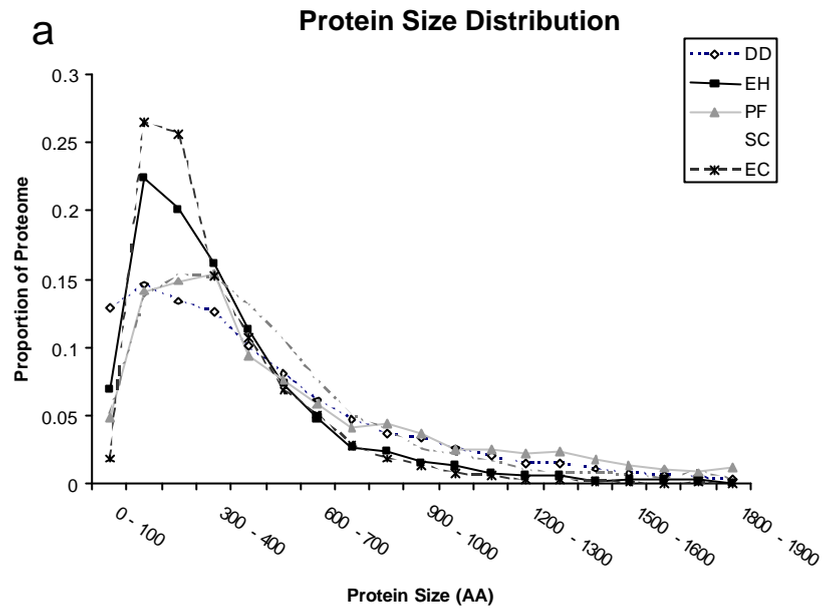


Fig. 1



Degree of reduction in length of proteins in *E. histolytica* compared to *D. discoideum*

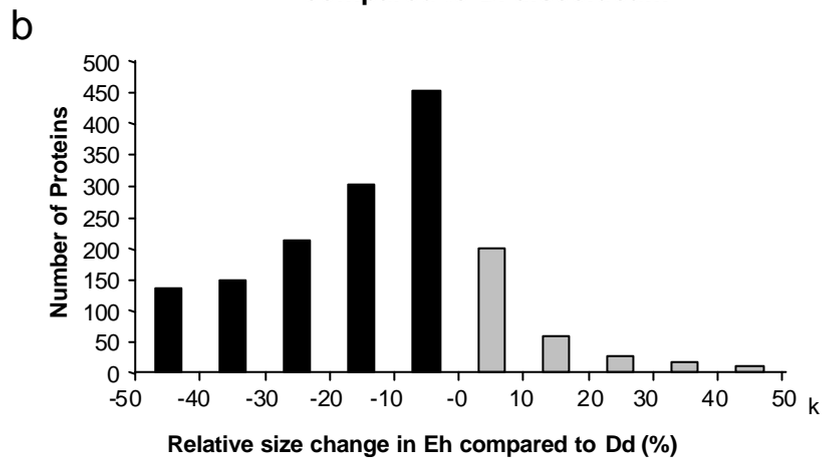


Fig. 2

1



Fig. 3

2

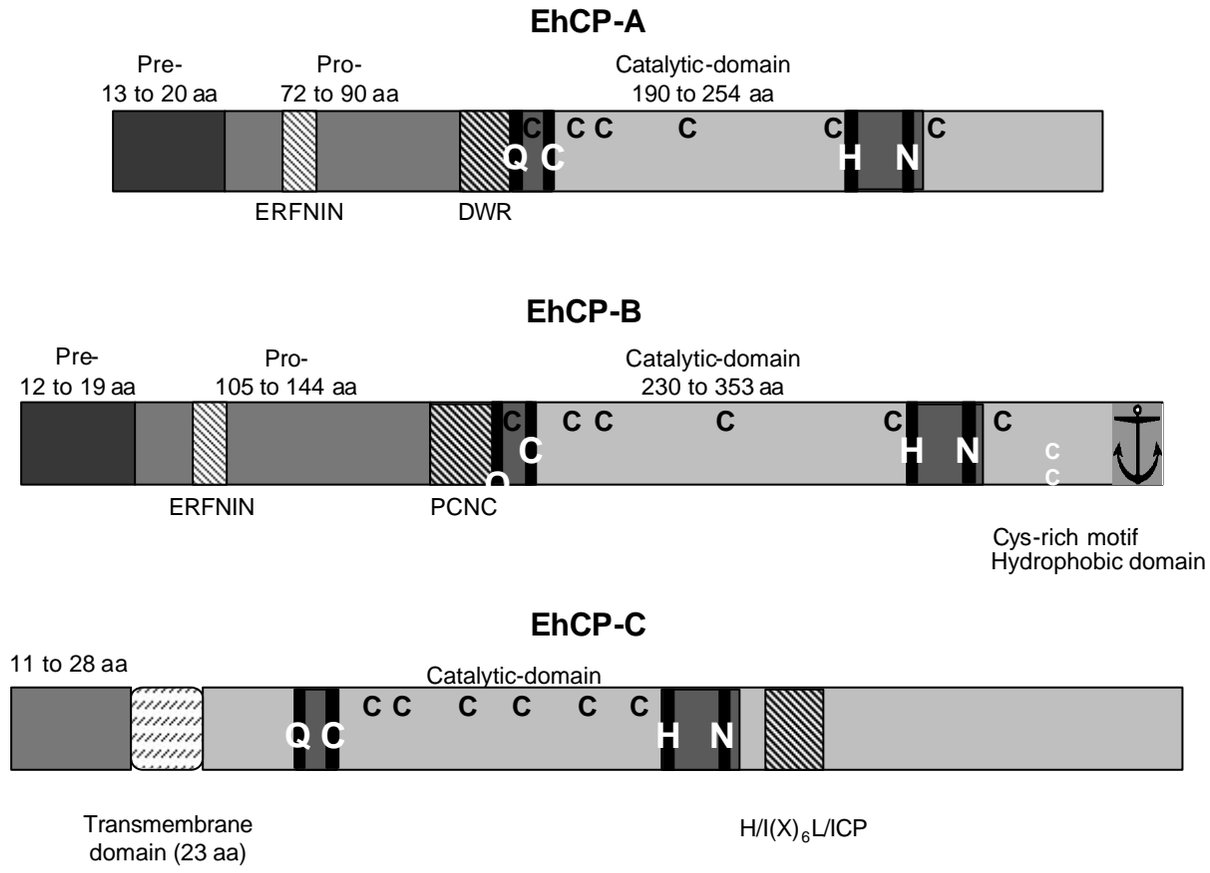


Fig. 4

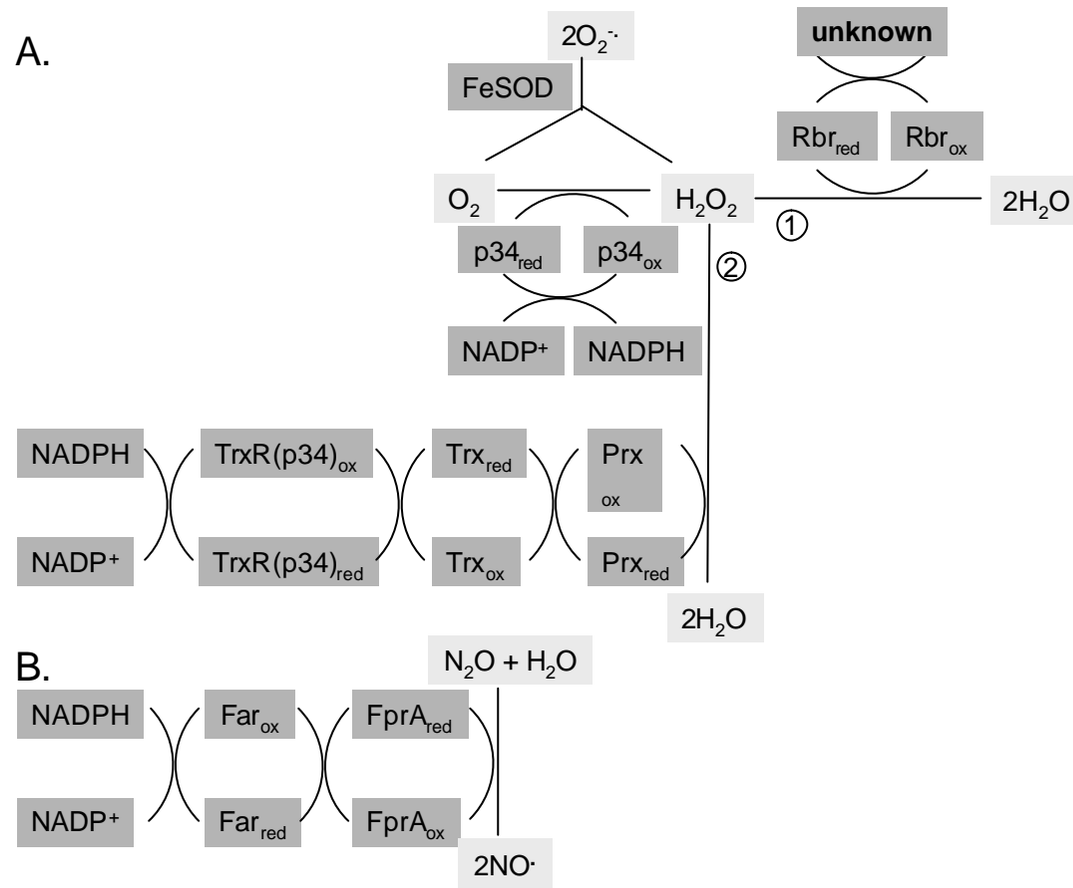


Fig. 5

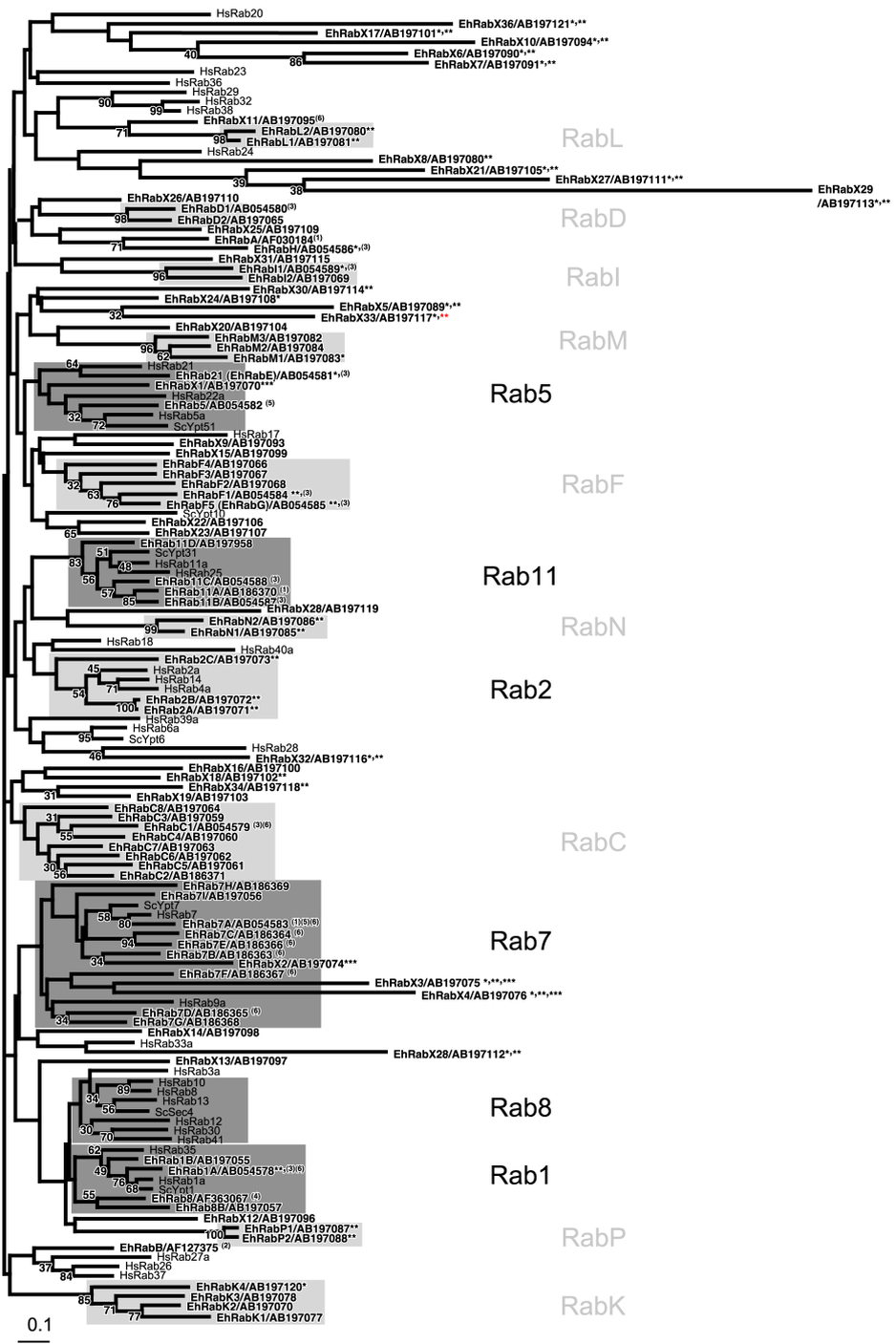
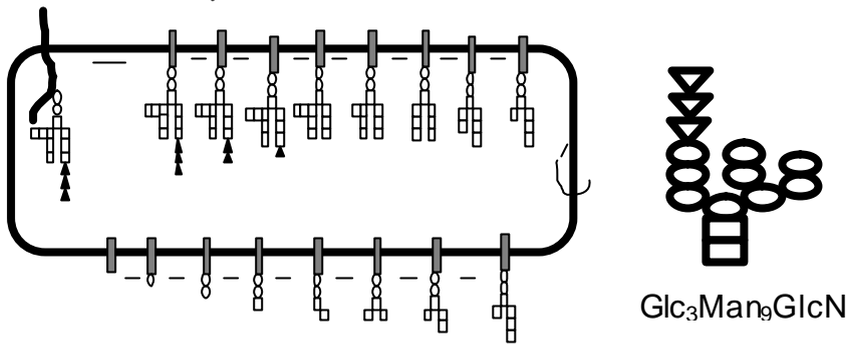
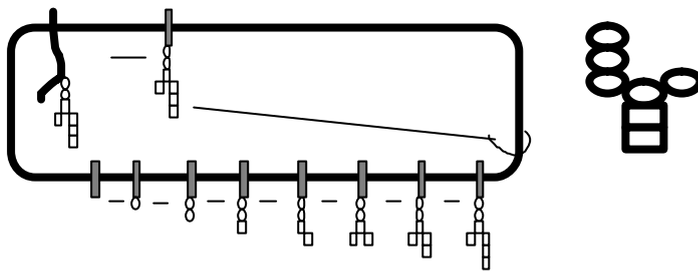


Fig. 6

A. *Saccharomyces*:B. *Entamoeba*:

1

2 Fig. 7

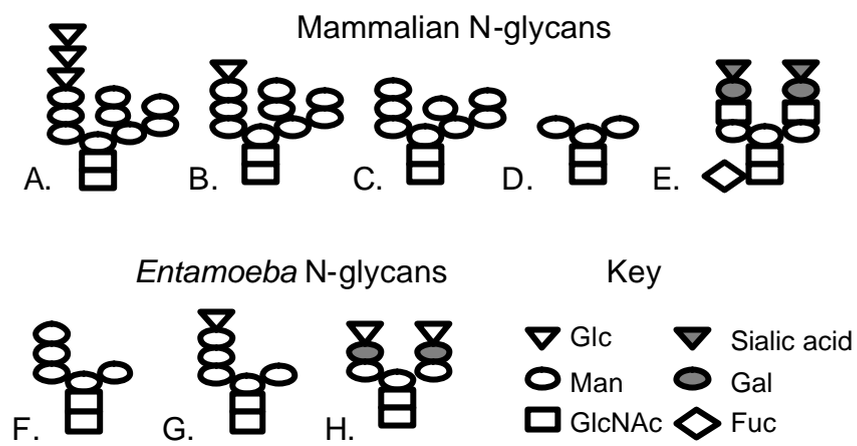


Fig. 8

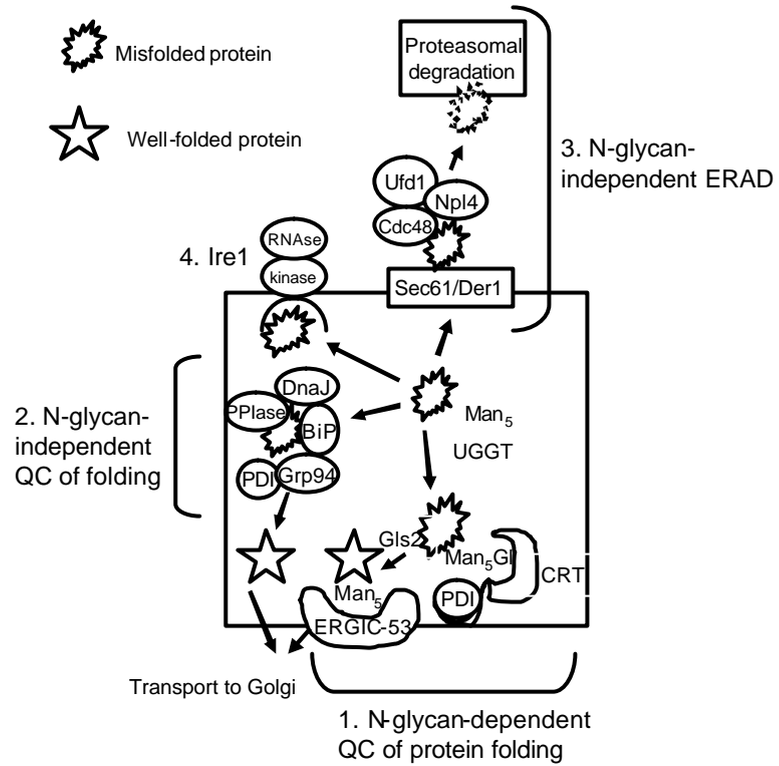


Fig. 9

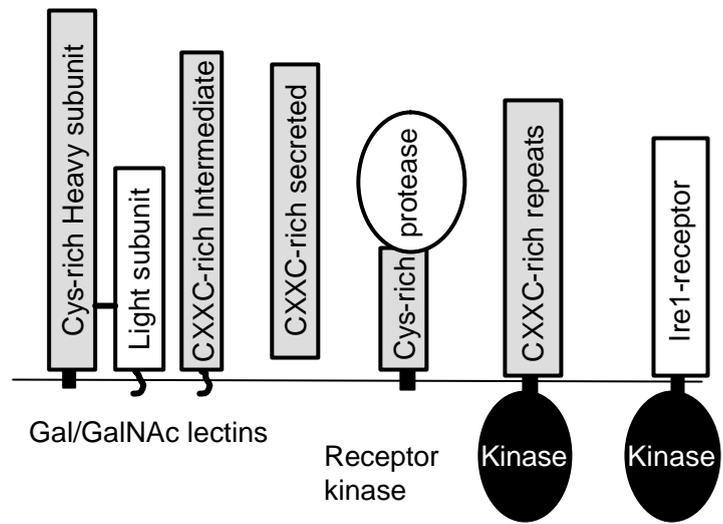
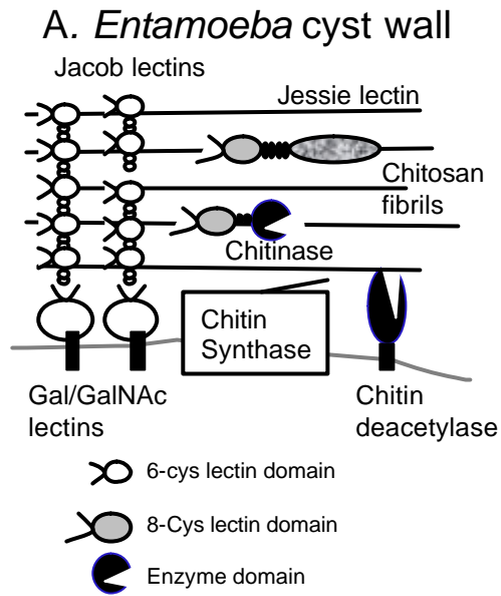


Fig. 10



B. *Entamoeba* cyst wall-associated lectins

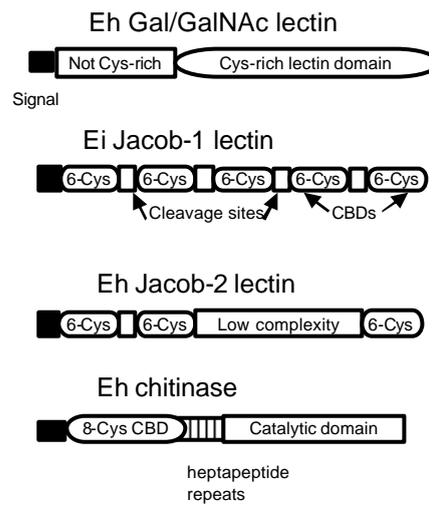


Fig. 11

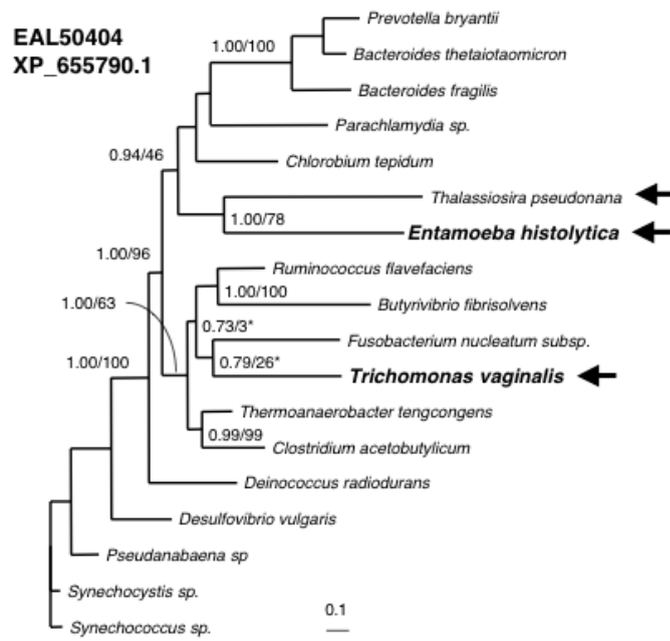


Fig. 12

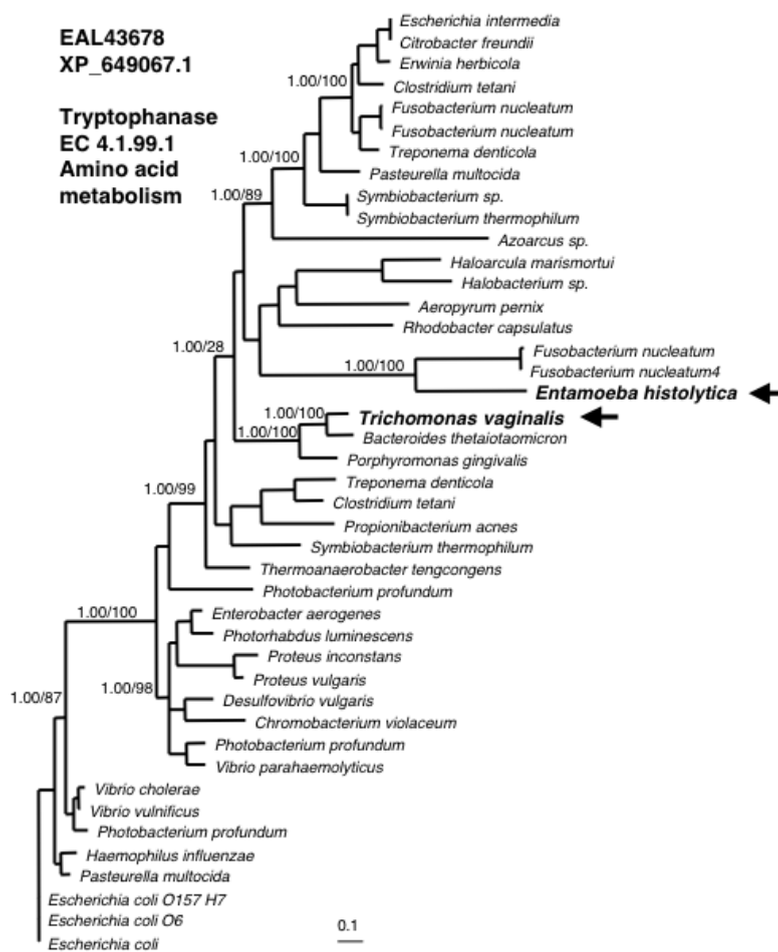


Fig. 13

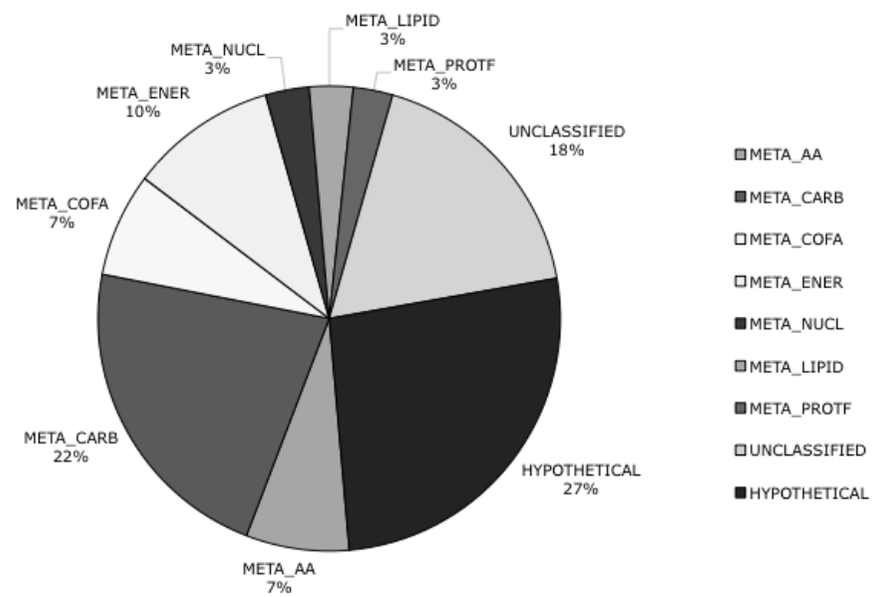


Fig. 14