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Shrink it or lose it
Balancing loss of function with shrinking genomes in the microsporidia

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Microsporidia are obligate intracellular parasites that have evolved an elaborate mechanism for invading animal host cells, but which have otherwise greatly reduced biological complexity. In particular, microsporidia possess the smallest autonomous nuclear genomes known (as opposed to nucleus derived organelles or nucleomorphs), and their 'anaerobic' core carbon metabolism is severely limited. Here we compare the extremes to which these two characteristics have evolved, and contrast how their reduction has either proceeded within the constraints of an unchanging set of functions, or has reduced the functional capabilities of the cell. Specifically, we review how the smallest known nuclear genome, the 2.3 Mbp genome of Encephalitozoon intestinalis, has arrived at this diminutive form without significantly affecting its protein-coding complexity in comparison with closely related, larger genomes. In contrast to this, Enterocytozoon bieneusi has a relatively large genome, and yet has lost all enzymes necessary to synthesize ATP from sugar—imposing a major limitation on the functional capabilities of the cell. The extremity of this reduction demands a re-evaluation of metabolic processes in other microsporidia: although pathways such as glycolysis are present, comparative genomic data suggest they may not play the cellular role that they are generally assumed to play.

Introduction

Microsporidia are an abundant and diverse lineage of obligate intracellular parasites related to fungi.1,2 Outside their host cell these organisms are only viable as highly resistant spores that are equipped with an intriguing suite of subcellular structures used to gain entry to their host cell. Most obvious of these is the polar filament, a coil wound about the inside of the spore that is ejected at high speed during germination, inverting at the same time to become a tube. Should this projectile happen to penetrate the cytoplasm of a host, it allows the contents of the spore to move through the tube and into that cytoplasm, where it can begin its infectious cycle.3 This mechanism may be used to attack a host cell from outside, to move between adjacent host cells or to escape from an endocytic vacuole to the cytoplasm.4

Aside from the machinery that mediates infection, microsporidia are surprisingly simple: so simple, in fact, they were for a time argued to be ancient, primitive eukaryotes.5 Based on molecular phylogenies, we now know this is not the case; instead, they are related to fungi and as such we now know they are derived or reduced, from more complex ancestors.2 This reduction has been the focus of some attention due to the extremes to which microsporidia have gone in losing characteristics we normally expect in a eukaryote, and it is now clear that these strange parasites have taken several routes to this end. Here, two different ways to reduce biological complexity that have been recently described in three papers6-8 will be illustrated and, by contrasting these, some of their broader implications will be discussed.
Loss of DNA—Trimming the Fat from Skinny Genomes

Though there are many odd things about the biology of microsporidia, the small size of their genomes has stood out since their genomes were first investigated. It is hard to make concrete generalizations about nuclear genomes, but we do have an imperfect concept of how they differ from those of bacteria: they are composed of multiple linear chromosomes with centromere and telomeres, and they are more often than not large by comparison. In structure, microsporidian genomes fit this conception, but in content many of them do not. The largest microsporidian genomes are 25–30 Mbp, well within the range for microbial eukaryotes, but others are much smaller, and indeed are the smallest known nuclear genomes of any eukaryote. At the extreme is the genus Encephalitozoon, a widespread parasite of vertebrates, including humans, where genome sizes smaller than 3 Mbp have been measured. To put this into context, this is more than 1,000x smaller than the human genome, and almost twice as small as that of E. coli K12. Almost ten years ago the genome of Encephalitozoon cuniculi was completely sequenced, and found to be a mere 2.9 Mbp. The genome has become a model for reduced and compacted genomes, with about 2,000 genes and a bacterial-like gene density.

But E. cuniculi does not have the smallest microsporidian genome—the smallest currently known is found in its cousin, E. intestinalis, which was estimated to be 2.3 Mbp or 20% smaller. Looking at the low number of genes and extreme density of the E. cuniculi genome, one wonders, where is the slack? If E. intestinalis really is 20% smaller, what is it missing compared to E. cuniculi?

To find out, we sequenced the complete genome of E. intestinalis. The sequence, composed of 11 chromosomes with no internal gaps, conformed closely to the original estimate of 2.3 Mbp and showed quite clearly where the 20% difference between E. cuniculi and E. intestinalis lay. Between the chromosome ‘cores’ (the region defined as spanning between the first and last identifiably homologous genes in the two species) there was little difference—more or less the same genes, in the same order, and at a similar density. Although the chromosome cores represent the vast majority of the genome, they only accounted for only 34 kbp of the size difference, suggesting it has been reduced to the limit of its functional constraints. This conclusion was supported by sequence evolution as well: intergenic regions showed a lower rate of substitution than synonymous positions in the genes surrounding them—suggesting most of the remaining intergenic sequences were constrained by selection, and so were likely functionally significant, or in other words if there was any excess, it had already been removed.

The chromosome ends, on the other hand, could hardly have been more different. The sequenced sub-telomeric regions of the E. intestinalis chromosomes were all significantly smaller than the corresponding ends in E. cuniculi, so that a total of at least 224 kbp encoding 174 genes were absent from E. intestinalis. Broadly, this answers the question of how an already compressed genome can be further reduced—simply snip off the ends of all the chromosomes. Sounds easy, but it raises as many questions as it answers. E. cuniculi is no slouch in terms of genome reduction, so why does it have hundreds of genes at its chromosome ends if they are dispensable in a closely related species? Clearly they are, under similar circumstances, non-essential, so why would such a tightly-packed genome have many kilobases of such genes, and why are they all clustered at the chromosome ends? The nature of the genes in question is obviously central to this, and not surprisingly many of them are unidentified ORFs or part of repeated families or both. It could be that these genes and families are rapidly turned over in these genomes, and E. cuniculi just happens to represent a temporary ‘bloated’ state due to a recent expansion. To determine if the genes were gained in E. cuniculi or lost in E. intestinalis, the genome of a third species will be necessary. It is also worth noting that that this difference may be adaptation in response to some pressure (i.e., there is some advantage to having a small genome if you are an Encephalitozoon). While intuitive, this has not been demonstrated convincingly, and it is also possible that even extreme reduction is due to some intrinsic characteristic of the genome that is non-adaptive. For instance, a bias towards deletions over insertions with limited recombination could pack the genome very tightly, and in small populations (the population structure of various microsporidia remains an open question) this could allow even mildly deleterious changes to drift to fixation.

As interesting as these changes are for genome evolution at its most radical extremes, it is noteworthy that they potentially have very little effect on the molecular or cell biology of the organisms themselves because of the nature of the missing genes and the fact that many of those that are part of duplicated families with homology elsewhere in the E. intestinalis genome. Overall, E. intestinalis is not missing any known pathway or functional cluster of genes that is present in E. cuniculi; so this difference in genome size could be the result of an intrinsic property of the genome itself as opposed to a reaction to any external pressure or change.

Loss of Function—
The Deceptive Simplicity of Microsporidian Metabolism

In contrast to the trimming of the E. intestinalis genome, other microsporidia have taken other reductive routes just as far; and here metabolism stands out. Microsporidian metabolism is at the best of times very restricted. Direct biochemical analyses are hampered by their obligate intracellular nature, but in the species where we have abundant sequence data, the metabolism can be modeled based on what genes are present. In all such species, many small molecule biosynthetic pathways are completely absent, and core carbon metabolism is limited to glycolysis, the pentose phosphate pathway, and synthesis and degradation of the storage carbohydrate trehalose—even the smallest E. intestinalis genome has the full complement of genes for these pathways. In Enteroctozoon bieneusi, however, virtually all these genes have been lost. This species cannot be maintained indefinitely in vitro so sequencing its entire genome is a challenge, and without a complete genome it is difficult to conclude a gene has really
been lost. To get around this problem, two separate, deep genome sequence surveys have been conducted on *E. bieneusi* and the list of genes generated by these two surveys compared.8,18 Remarkably, nearly identical lists of genes were found, suggesting the genome is very well sampled, though incomplete. More specifically, an analysis of over 300 genes representing 30 functional pathways suggested that over 90% of the genes found in *E. cuniculi* had also been sampled in *E. bieneusi*. In contrast, of the 21 *E. cuniculi* genes for core carbon metabolism, only two were found in *E. bieneusi* and, significantly, the same two were found in both surveys.8,18 Moreover, fatty acid metabolism was also severely underrepresented and, fascinatingly, both spliceosomal introns and all the proteins and RNAs associated with spliceosomal splicing were completely absent, suggesting this genome has also lost all traces of intron splicing.8,18 Overall, it appears that this species completely lacks any capacity to carry out several normal eukaryotic activities, the most surprising of which is the ability to generate energy from sugars. Instead, it must rely on importing ATP directly from its host. Unlike *E. intestinalis*, the reductive process in *E. bieneusi* has not led to a particularly small genome (it is thought to be about 6 Mbp)8,18 but, instead, to a major change of function since what are normally important functional pathways have been entirely lost.

The absence of these pathways in *E. bieneusi* has striking implications for its metabolism, but also significant implications on the role played by these pathways in other microsporidia. Glycolysis is generally regarded as the universal backbone of energy metabolism in anaerobic eukaryotes; and microsporidia are no different.19,20 However, although the pathway is intact in other microsporidian genomes, the *E. bieneusi* genome and other findings described below give us reason to doubt its role in these species. In addition to generating ATP, glycolysis generates reducing potential in the form of NADH. Without an outlet for this potential, it must be dissipated or the pathway becomes energetically unfavorable. Some eukaryotes dissipate this excess potential by transferring electrons to the mitochondrial via the glycerol-3-phosphate (G3PDH) shuttle where they are absorbed by the alternative oxidase (AOX).21,22 Recently, homologs of AOX have been found in some, but interestingly not all, microsporidia.6 *Antonospora locustae*, for example, has genes for all the enzymes of glycolysis, AOX, both partners for the G3PDH shuttle, and proteins necessary to target the mitochondrial G3PDH to the mitochondrion;6,23,24 altogether making up a complete system to make ATP from glucose (aerobically, which is noteworthy). Not surprisingly, Enterocytozoon has none of these systems intact,8 but other species such as Encephalitozoon, fall somewhere between these two extremes. Encephalitozoon genomes lack AOX6 and the G3PDH shuttle has been shown to be broken in *E. cuniculi*, since the mitochondrial partner is localized to the cytosol25 and it lacks a peptidase responsible for processing the targeted product in other eukaryotes.24 Overall, Encephalitozoon might have glycolysis but apparently lacks any obvious capacity to keep the pathway going continuously, as one might imagine it would do if it were being used as a significant source of ATP.

This suggests Encephalitozoon metabolism may be closer to that of Enterocytozoon than their genomes superficially suggest. Encephalitozoon genomes, and indeed all known microsporidian genomes, encode several ATP transporters,26 some that import ATP from the host,27 and others that import it into their reduced mitochondrion.28 So even if a given species of microsporidian is not entirely dependent on host ATP, as Enterocytozoon seems to be, all microsporidia appear to at least have the capacity to be so dependent. Perhaps the real questions should not be why *E. bieneusi* has lost glycolysis, but rather why all the other microsporidia that possess ATP transporters have not lost it as well? If ATP can be imported, then the synthesis of ATP (via glycolysis or any other means) needs to be explained in the context of some particular life stage or environmental clue making it critical for survival.

Lacking the necessary components to keep glycolysis running continuously, it appears possible that Encephalitozoon does not actually use glycolysis for energy: like Enterocytozoon, Encephalitozoon may also import all its ATP from its host. If this is true, and Encephalitozoon lacks the ability to use glycolysis as a continuous source of energy, then why has it retained all the genes for this pathway when they have been lost in Enterocytozoon? One possibility is, ironically, that Encephalitozoon is using glycolysis to make NADH. While excess NADH is a potential problem with glycolysis running continuously, a small amount of NADH could be made by metabolizing glucose (sparsely) by glycolysis, and the ATP also produced would be a useful by-product of this NADH production.

**Concluding Remarks— Microsporidia Entering the Era of Comparative Genomics**

Microsporidia are a phylogenetically diverse group of parasites and, although they share the same fundamental infection mechanism, they also display a significant amount of diversity in how they use that mechanism, how they affect their hosts, their life cycle variations, and in other traits relating to their life histories.29 Indeed, their genetic diversity and ubiquity across the diversity of animals leads one to speculate that they may have originated at about the same time as the animal lineage and represent a nearly equivalent amount of diversity. Although impossible to quantify, anecdotal data suggest that any animal species investigated in sufficient detail will eventually be found to be infected by one or more species of microsporidia and that many of these have limited host range. One averages, one could even mount a reasonable argument that there are a similar number of species of microsporidian as there are of animals. For these reasons, we have paid a great deal of attention to the variation between different microsporidia, but it is worth pointing out that one of the emerging themes of comparative genomics of microsporidia is the remarkable lack of variation at other levels. Despite the differences detailed above, the overall gene content of diverse microsporidia is remarkably homogeneous.7,8,11,13,18,30,31 Indeed, the overall degree of conservation in gene content serves to highlight what differences we do find. Nevertheless, these differences...
are real and many are of functional or evolutionary significance, so comparative genomics has also served to remind us that no one ‘model’ will adequately represent the lineage as a whole.

Comparing the genomes of microsporidia has also served to illustrate that the genome of one species may be difficult to interpret at face value. The presence of genes for a complete glycolytic pathway in *E. cuniculi* can present an obvious interpretation on its own, but superimposing on this the implications of findings from the *E. bieneusi* and *A. locustae* genomes significantly alters this interpretation. Given the relatively small size of even the largest microsporidian genomes, next-generation sequencing technology will soon enable the generation of a great deal of comparative data from many microsporidian species: this kind of data has already changed much of how we understand this group (e.g., many of them should no longer be considered ‘anaerobes’ as they once were described because they metabolize oxygen using AOX) and how this transition continues will be interesting to observe. One level where such technology will be of obvious interest is that of the population. The population structure of most microsporidia is all but unknown, so even basic questions such as whether many species are sexual or not are mysteries. Just as comparing genomes between distantly related microsporidian genera or between different species of the same genus addressed distinct kinds of questions, so too comparing genomes from multiple strains of a microsporidian species has the potential to tell us a great deal about other levels of diversity, evolution and even protein and genome function.

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