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The reduced genome of *Candidatus* Kinetoplastibacterium sorsogonicusi, the endosymbiont of *Kentomonas sorsogonicus* (Trypanosomatidae): loss of the haem-synthesis pathway

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Abstract

Trypanosomatids of the genera Angomonas and Strigomonas (subfamily Strigomonadinae) have long been known to contain intracellular beta-proteobacteria, which provide them with many important nutrients such as haem, essential amino acids and vitamins. Recently, Kentomonas sorsogonicus, a divergent member of Strigomonadinae, has been described. Herein, we characterize the genome of its endosymbiont, Candidatus Kinetoplastibacterium sorsogonicusi. This genome is completely syntenic with those of other known Ca. Kinetoplastibacterium spp., but more reduced in size (~742 kb, compared with 810-833 kb, respectively). Gene losses are not concentrated in any hot-spots but are instead distributed throughout the genome. The most conspicuous loss is that of the haemsynthesis pathway. For long, removing haemin from the culture medium has been a standard procedure in cultivating trypanosomatids isolated from insects; continued growth was considered as an evidence of endosymbiont presence. However, we demonstrate that, despite bearing the endosymbiont, K. sorsogonicus cannot grow in culture without haem. Thus, the traditional test cannot be taken as a reliable criterion for the absence or presence of endosymbionts in trypanosomatid flagellates. It remains unclear why the ability to synthesize such an essential compound was lost in Ca. K. sorsogonicusi, whereas all other known bacterial endosymbionts of trypanosomatids retain them.

Introduction

The study of symbiosis, and particularly mutualistic endosymbiosis (defined as one organism living inside another in a cooperative arrangement) has been gaining importance over time as such relationships grow in number and diversity. While endosymbiosis has been long considered a central phenomenon in the evolution of eukaryotic life and the origin or organelles (for a review, see Archibald, 2015), it is also implicated in more recent, but still significant, biological interactions in organisms ranging from bacteria to single-celled eukaryotes to Metazoa (usually insects) and plants.

Endosymbionts frequently specialize in complementing the metabolic capabilities of the host (Douglas, 2016). For example, in the nested endosymbiosis involving two bacteria (Gammaproteobacteria within a Betaproteobacteria) within mealybug bacteriocytes (von Dohlen *et al.* 2001), each bacterium has retained just the genes necessary for supplying the insect with essential amino acids (McCutcheon and von Dohlen, 2011), complementing each other as well as the insect host. Secondary endosymbiosis between eukaryotes has been responsible for the origin of complex organisms such as parasites of the phylum Apicomplexa, which usually contain a highly reduced algal cell (apicoplast) in the cytoplasm (Gentil *et al.* 2017). The study of endosymbiotic relationships is thus of great interest in the characterization of genetic, metabolic, ecological and evolutionary aspects.

The parasitic flagellates of the subfamily Strigomonadinae (Kinetoplastea: Trypanosomatidae) are gut-dwelling insect parasites bearing endosymbiotic bacteria from the family Alcaligenaceae (Betaproteobacteria) in their cytoplasm (Teixeira *et al.* 2011; Votýpka *et al.* 2014). Owing to this peculiarity, the Strigomonadinae are relatively well studied, although not to such extent as their relatives from the genera *Trypanosoma* and *Leishmania*, many of which are of medical or veterinary importance (Nussbaum *et al.* 2010).

As uncovered in studies of species from the genera *Angomonas* and *Strigomonas*, the collaboration between the endosymbiont and its trypanosomatid host is very close and finely tuned. There is typically only one bacterium inside each trypanosomatid cell, and the division of the two partners is synchronized (Motta *et al.* 2010). Significantly, as shown by biochemical and cell biology experiments over decades and recently characterized at the genome level, the bacterium provides the eukaryotic host with several important nutrients, such as essential amino acids (Alves *et al.* 2013*b*), vitamins (Klein *et al.* 2013) and the haem group (Kořený *et al.* 2010; Alves *et al.* 2011). The bacterium cannot survive outside its host, while the aposymbiotic (i.e. 'cured' by antibiotic treatment) trypanosomatid can grow only in a medium supplemented with those compounds previously provided by the endosymbiont.

Although these parasites have been studied for a long time, their taxonomic status has only recently been clarified through the use of molecular data (Teixeira et al. 2011), placing them into the sister genera Angomonas and Strigomonas. Each of two Angomonas (A. deanei and A. desouzai) and three Strigomonas species (S. culicis, S. galati and S. oncopelti) carries its own species of Candidatus Kinetoplastibacterium (Ca. K. crithidii, Ca. K. desouzaii, Ca. K. blastocrithidii, Ca. K. galatii and Ca. K. oncopeltii, respectively). The sole exception is A. ambiguus, which bears the same species of endosymbiont as A. deanei, in spite of its nuclear genome being more closely related to A. desouzai. Apart from that ambiguity, the phylogenetic tree of these trypanosomatids shows an identical branching pattern to that seen for the corresponding symbionts, providing compelling evidence that there was a single origin for this endosymbiosis, followed by co-speciation (Teixeira et al. 2011).

More recently, a third genus in this subfamily was discovered, containing so far only one described species: *Kentomonas sorsogonicus* (Votýpka *et al.* 2014). This trypanosomatid, like all other Strigomonadinae, contains an intracytoplasmic endosymbiont, *Ca.* K. sorsogonicusi. Preliminary phylogenetic analyses of the host and the bacterium using their SSU rRNA genes showed conflicting results, with *Kentomonas* being sister to the rest of Strigomonadinae, but its symbiont clustering with those from *Angomonas*, thereby making the bacteria of *Strigomonas* spp. the earliest branch of the genus *Ca.* Kinetoplastibacterium. However, the relationships between endosymbionts were poorly supported (Votýpka *et al.* 2014).

Herein, we characterize the complete genome of *Ca.* K. sorsogonicusi, showing that it is even more reduced in size than the genomes of other *Ca.* Kinetoplastibacterium spp., having lost the haem-synthesis pathway until now regarded as a hallmark of endosymbionts in trypanosomatids. We also resolve the phylogenetic position of this organism within its genus by phylogenomic methods.

Materials and methods

Organism cultivation and DNA isolation

For genomic DNA preparation, *K. sorsogonicus* strain MF08 was grown in the BHI medium as previously described (Votýpka *et al.* 2014). Log-phase trypanosomatid cells were collected by centrifugation for 10 min at 1000 *g*, washed once with PBS and frozen at -20 °C until DNA isolation. The latter was performed using DNeasy Blood & Tissue Kit (Qiagen, Hilden, Germany) according to the manufacturer's instructions.

Haem requirement tests were performed in the M199 medium (Life Technologies, Carlsbad, USA) without fetal bovine serum and supplemented with biopterin (2 μ g mL⁻¹), HEPES (25 mM) as well as a mix of microelements as described previously (Porcel

et al. 2014). Log-phase PBS-washed trypanosomatid cells were seeded into flat-sided tubes with 2 mL of this medium both with and without 10 μ g mL⁻¹ of haemin (M199+ and M199-, respectively). After 3 days, 100 μ L of the culture from the M199- tube were placed into new tubes with M199+ and M199-.

Genome sequencing and assembly

Shotgun genome sequencing of *K. sorsogonicus* and its symbiont was performed jointly, without separating the two organisms. Genomic sequencing (Macrogen Inc., Seoul, South Korea) was performed using an Illumina paired-end library (2×100 , inserts of on average 600 bp) constructed using the TruSeq DNA PCR Free kit.

Reads were trimmed using cutadapt (Martin, 2011) to remove adapters and low-quality regions, and assembled using the Newbler v.2.9 assembler (distributed by 454 Roche).

Contigs and scaffolds corresponding to the endosymbiont were initially separated from the host sequences by BLASTN (Camacho *et al.* 2009) sequence similarity searches with other, fully sequenced *Ca.* Kinetoplastibacterium genomes. A minimum of 80% sequence similarity along most of the contig was used as the threshold for considering it as part of the endosymbiont's genome.

The final order of the contigs on the bacterial genome was identified by PCR-amplification using PCRBIO Taq Mix Red (PCR Biosystems Ltd., London, UK) and custom primers annealing at the ends of contigs: Ks1F (5'-gttctcttatgatacctccagt-3'), Ks1R (5'-ctccctcctatatattgct-3'), Ks2F (5'-agcacattgtaatccaggta-3'), Ks2R (5'-ggattgctagtatggtagaagg-3'), Ks3F (5'-cacaatcaggtgtagcatgt-3'), Ks3R (5'-tcaaatactcatacacgtgc-3'), Ks4F (5'-accatagtagcaaaaccag-3'), Ks56R (5'-cacgttccagatactattactcac-3'). All generated fragments were sequenced directly by the Sanger method (Macrogen Europe, Amsterdam, The Netherlands) using the amplification primers as well as those annealing to 16S and 23S rRNA genes (see Votýpka *et al.* 2014).

Genomic annotation and analysis

The finished genome was annotated automatically by the Prokka pipeline v. 1.12 (Seemann, 2014) and annotations were sampled for manual validation against previously annotated *Ca*. Kinetoplastibacterium genomes whose annotations had been manually curated (see below).

Pseudogenes and missing predictions were identified by BLASTX of intergenic regions against the NCBI non-redundant database, followed by a manual examination.

The assembled genomic sequence and its annotation are available under accession number CP025628 (NCBI BioProject PRJNA414463). Other genomic sequences used in this work are: *Ca.* K. blastocrithidii TCC012E (accession number CP003733.1), *Ca.* K. crithidii TCC036E (CP003804.1), *Ca.* K. desouzaii TCC079E (CP003803.1), *Ca.* K. galatii TCC219 (CP003806.1), *Ca.* K. oncopeltii TCC290E (CP003805.1) (all five from Alves *et al.* 2013*a*), *Achromobacter arsenitoxydans* SY8 (AGUF00000000.1) (Li *et al.* 2012) and *Taylorella equigenitalis* MCE9 (CP002456.1) (Hebert *et al.* 2011).

Overall structural comparison of *Ca.* K. sorsogonicusi and selected *Ca.* Kinetoplastibacterium spp. was performed by genome alignment in MAUVE v. 2015-02-13 (Darling *et al.* 2010). Gene-level comparisons were based on the orthology inference results of OrthoMCL (Li *et al.* 2003) analyses. Comparative metabolic annotation analysis of the endosymbionts from *A. desouzaii, K. sorsogonicus, S. culicis* and *S. oncopelti* was performed with ASGARD (Alves and Buck, 2007), using the KEGG (Ogata *et al.* 1999) and UniRef100 (Suzek *et al.* 2007) databases as

references, followed by manual curation of results of interest. Circos (Krzywinski *et al.* 2009) was used for generating genome comparison plots.

Phylogenetic analysis

All predicted proteins from either just the *Ca*. Kinetoplastibacterium spp. or all eight genomes mentioned above were analysed by OrthoMCL (Li *et al.* 2003) in order to find all single-copy genes that were present in all organisms considered in each analysis. All such orthologous groups (OGs) identified were then aligned by MUSCLE v. 3.8.31 (Edgar, 2004), with subsequent removal of ambiguously aligned positions by Gblocks v. 0.91b (Castresana, 2000). The resulting filtered alignments were concatenated into a supermatrix with FASconCAT-G v.1.04 (Kück and Meusemann, 2010), after conversion of alignment files to the requirements of that program using in-house Perl scripts.

Partitioned phylogenetic analysis of the resulting supermatrix was performed by maximum likelihood using RAxML v. 8.2.11 (Stamatakis, 2014) and running 100 bootstrap pseudoreplicates. Protein substitution models were selected automatically by the program, estimated separately for each partition using a maximum-likelihood criterion. Additionally, partitioned Bayesian inference was performed with MrBayes v. 3.2.6 (Ronquist et al. 2012) using two runs of four chains (three heated) each for 100 000 generations (while monitoring run convergence parameters), using the 'mixed' amino acid substitution model and discarding 25% of the generations as burn-in. Substitution rate heterogeneity for each partition was modelled with gammadistributed rates for both maximum likelihood and Bayesian inference methods.

Phylogenetic trees were drawn with MEGA v. 7 (Kumar *et al.* 2016) and cosmetic adjustments were performed in the Inkscape vector editor (https://inkscape.org).

Results

Genomic characterization

Genomic sequencing by Illumina paired-end technology yielded about 22.8 million pairs of 100 bp reads, for a total of 4.57 billion base pairs (around 114-fold coverage of a 40 million base pair genome). The assembly resulted in seven contigs that could be identified, by similarity to other *Ca.* Kinetoplastibacterium 1289

genomes, as belonging to the endosymbiont. These sequences were used for the design of PCR primers to amplify the adjacent genomic regions. The obtained fragments were sequenced, leading to the generation of an unambiguously complete genome assembly.

The genome of *Ca.* K. sorsogonicusi contains 741 697 base pairs and an overall GC content of 25.22%. Coding regions have similar composition, at 25.49% GC on average, while ribosomal and transfer RNA genes present the much higher average GC content of 50.36% (Table 1). The overall genome organization of *Ca.* K. sorsogonicusi can be seen in Fig. 1. The total number of genes predicted to be present in the genome is 722, with 670 protein-coding genes, three pseudogenes, 39 genes for tRNAs, nine genes for rRNAs (distributed in three clusters), and the gene for the ssrA transfer-messenger RNA.

The ribosomal gene clusters are not identical, with only one of them (the last one, starting at around position 590 000) containing two tRNAs between the large subunit rRNA (LSU) and the small subunit rRNA (SSU) genes. The SSU, LSU and 5S rRNA genes are, in this order, present in each of the three ribosomal gene clusters.

At least one tRNA gene has been identified for each of the 20 amino acids present in natural proteins. Eleven of the amino acids can be ligated to only one predicted tRNA. Leucine is the amino acid with the most tRNA genes predicted, with five, while arginine and serine have four tRNA genes each.

OG inference of the six *Ca*. Kinetoplastibacterium spp. genomes yielded 769 OGs, of which 132 were absent from *Ca*. K. sorsogonicusi. Among these 132 OGs, 68 (Supplementary File 1) were found in all other five *Ca*. Kinetoplastibacterium spp. genomes analysed here.

Figure 1 shows that the overall gene distribution in the *Ca.* K. sorsogonicusi genome is typical of those of *Ca.* Kinetoplastibacterium spp., with genes tightly spaced. Its comparison with the genomes of *Ca.* K. desouzaii and *Ca.* K. oncopeltii (the longest and shortest ones, respectively, among the previously sequenced species of the genus) demonstrates their overall synteny, which can be also extrapolated to all other *Ca.* Kinetoplastibacterium genomes studied so far. The GC skew plots for the three genomes are similar to those previously seen for *Ca.* K. blastocrithidii and *Ca.* K. crithidii (Alves *et al.* 2013*a*). A little more than half of each genome shows predominantly positive GC skew whilst the remainder exhibits mostly negative skew (Fig. 1).

Table 1. Overal	l genome	and ge	ene com	position	statistics
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	CKsor	CKcri	CKdes	CKbla	CKgal	CKonc	Tequi
Genome length (bp)	741 697	821 930	833 125	820 029	822 140	810 172	1 695 860
CDS genes ^a	670	728	742	724	727	694	1556
tRNA genes	39	44	43	43	43	43	38
rRNA genes	9	9	9	9	9	9	9
Pseudogenes	3	1	1	7	4	20	0
GC%	25.22%	30.96%	30.17%	32.55%	32.36%	31.23%	37.42%
GC% CDS	25.49%	31.38%	30.64%	33.04%	32.79%	31.87%	37.78%
Longest CDS (bp)	5556	5451	5427	4185	5559	5520	9996
GC% RNAs	50.36%	52.29%	52.37%	52.13%	51.96%	51.98%	52.26%
% genes ^b	92.46%	91.63%	91.68%	90.59%	91.52%	89.09%	93.45%

Genome codes: CKsor: Candidatus Kinetoplastibacterium sorsogonicusi; CKcri: Ca. K. crithidii; CKdes: Ca. K. desouzaii; CKbla: Ca. K. blastocrithidii; CKgal: Ca. K. galatii; CKonc: Ca. K. oncopeltii; Tequi: Taylorella equigenitalis MCE9.

^aPercentage of the total genome length that is in genes.

^bNumbers include only apparently intact coding sequences, excluding pseudogenes.

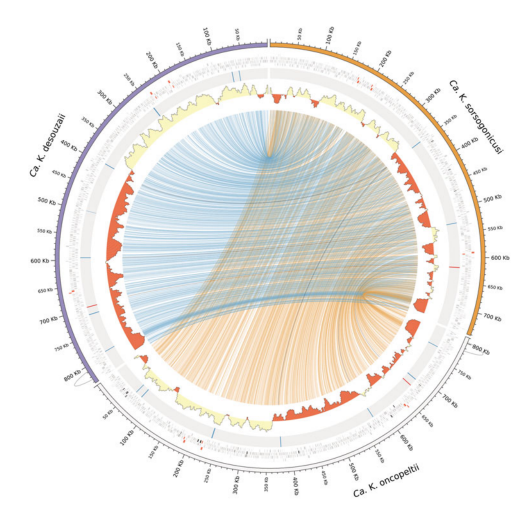


Fig. 1. Schematic representation of the structure of, and comparison between, the genomes of *Ca.* K. sorsogonicusi, *Ca.* K. desouzaii and *Ca.* K. oncopeltii. Starting from the outside, the different rings mean, in order: genomic coordinate representations; predicted genes (in grey, protein-coding genes; in red, rRNA genes; in blue, tRNA genes; in green, the ssrA genes; and in black, pseudogenes); highlights of the position of the genes participating in haem synthesis in each organism (in red, the gene for glutamyl-tRNA synthetase, which is not specific to haem synthesis; in blue, all others); GC skew plots (negative skew in red and positive skew in yellow). The central area of the schema shows lines connecting the different orthologous groups found in both *Ca.* K. sorsogonicusi and *Ca.* K. desouzaii (blue lines) or *Ca.* K. oncopeltii (orange lines); black lines highlight the linkage between haem-synthesis genes present in all three organisms.

The only protein-coding gene duplication present in the *Ca.* Kinetoplastibacterium spp. genomes involves the EF–Tu gene. It is shown in Fig. 1 by lines running outside the genome coordinate circle and connecting the gene copies, close to the end of each sequence. This duplication is not present in *Ca.* K. sorsogonicusi, in a situation similar to that seen in *Ca.* K. galatii, where one of the copies was pseudogenized. In *Ca.* K. sorsogonicusi, however, even the pseudogene is absent.

arsenitoxydans in accordance with the previous phylogenomic inference of the Betaproteobacteria (Alves *et al.* 2013*a*). All bootstrap support values and posterior probabilities in the tree reached the maximum values of 100 and 1.0, respectively.

The three endosymbionts from *Strigomonas* spp. formed a single clade, while those from *Angomonas* composed another one. *Ca.* K. sorsogonicusi was placed as the earliest group to split from the rest of *Ca.* Kinetoplastibacterium spp. (Fig. 2).

Phylogenetic analysis

The inference of OGs by OrthoMCL yielded 506 single-copy genes present in the six *Ca.* Kinetoplastibacterium spp. and two species from other genera of the family Alcaligenaceae. Individual OG alignments were concatenated into a single data matrix comprising 186 276 columns. After removal of ambiguously aligned positions, 159 196 columns (86% of the original amount) remained in the alignment.

Phylogenomic analysis of all eight organisms by maximum likelihood resulted in the tree presented in Fig. 2. The Bayesian inference consensus tree is identical in topology and has nearly identical branch lengths (not shown). The outgroups (*A. arsenitoxydans* and *T. equigenitalis*) were used for rooting the tree, with the root being placed on the branch leading to *A.*

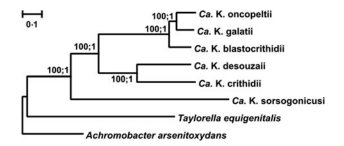


Fig. 2. Maximum-likelihood protein supermatrix phylogeny of 506 orthologous groups from *Ca.* Kinetoplastibacterium spp. and two other Alcaligenaceae bacteria as outgroups. Semi-colon-separated numbers on each node represent the maximum-likelihood bootstrap support value (up to 100) and the Bayesian inference's posterior probability (up to 1) for that node.

Genomic analyses of metabolic pathways

Comparison of metabolic pathways revealed only one of them was significantly disrupted in *Ca.* K. sorsogonicusi, whereas most other differences in enzyme presence or absence were scattered across the metabolic network. The haem pathway contains only two out of nine expected specific enzymes: oxygen-independent coproporphyrinogen-III oxidase (*hemN*) and ferrochelatase (*hemH*), respectively the eighth and the tenth (and last) enzymes of the pathway (Fig. 3 and Supplementary File 2, map 00 860, Porphyrin and chlorophyll metabolism). The first enzyme shown in the pathway, glutamyl-tRNA synthetase (*gltX*), not specific for the haem-synthesis pathway, is also present in the genome.

We have also searched the unpublished draft sequence of the trypanosomatid host, *K. sorsogonicus*, for the haem-synthesisspecific genes, but found none.

Other biosynthetic pathways that have been previously shown, in other *Ca*. Kinetoplastibacterium spp., to be provided by the bacterium to complement the trypanosomatid host's metabolism were examined in *Ca*. *K*. sorsogonicusi and compared to other bacteria from the genus (Supplementary File 2). Pathways for the synthesis of essential amino acids as well as vitamins B_6 , pantothenic acid, folic acid and riboflavin are essentially identical between *Ca*. K. sorsogonicusi and the other *Ca*. Kinetoplastibacterium spp. compared.

Haem requirement test

In the first passage, the growth of *K. sorsogonicus* in culture was observed in both M199+ and M199– media. However, in the latter case, it was much slower. Taking into account potential left-overs of haemin on the cell surface or in intracellular depots, we passaged the trypanosomatids from M199– to new tubes containing both variants of media. It this case, the growth was observed only in M199+, whereas in M199– it stopped and after one week all cells in this medium were dead.

Discussion

Given a number of characteristic morphological features as well as the presence of the beta-proteobacterial endosymbiont (Votýpka *et al.* 2014), *Kentomonas* is definitely a member of the subfamily Strigomonadinae. As mentioned above, in this subfamily the hosts and their respective endosymbionts almost always present (with the exception being *A. ambiguus*) a perfect pattern of co-speciation (Teixeira *et al.* 2011). However, in the first description of the *Kentomonas* genus, the trypanosomatid host and its endosymbiont had discordant phylogenetic positions, as judged

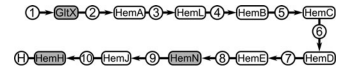


Fig. 3. Schematic representation of the haem-synthesis pathway in *Ca.* Kinetoplastibacterium spp. Compounds are represented within circles, and enzymes are within rounded-corner rectangles. Grey background highlights the enzymes for which genes were found in the *Ca.* K. sorsogonicusi genome. Compounds are 1: L-glutamate; 2: L-glutamyl-tRNA; 3: glutamate-1-semialdehyde; 4: aminolevulinic acid; 5: porphobilinogen; 6: hydroxymethylbilane; 7: uroporphyrinogen III; 8: copro-porphyrinogen III; 9: protoporphyrinogen IX; 10: protoporphyrin IX; H: haem. Enzymes are GIX: glutamyl-tRNA synthetase (EC:6.1.1.17); HemA: glutamyl-tRNA reductase (EC:1.2.1.70); HemL: glutamate-1-semialdehyde 2,1-aminomutase (EC:5.4.3.8); HemB: aminolevulinic acid dehydratase (EC:4.2.1.24); HemC: porphobilinogen deaminase (EC:2.5.1.61); HemD: uroporphyrinogen III synthase (EC:4.2.1.75); HemE: uroporphyrinogen III decarboxilase (EC:4.1.1.37); HemN: oxygen-independent coproporphyrinogen-III oxidase (EC:4.99.1.1).

by the inference using SSU rRNA genes. *Kentomonas* was the earliest branch within its subfamily, while *Ca.* K. sorsogonicusi, albeit with low support values, appeared as the sister taxon to the endosymbionts of *Angomonas* spp. Thus, the bacteria from *Strigomonas* spp. seemed to be the first diverging lineage within the genus *Ca.* Kinetoplastibacterium. Our multigene phylogenetic analyses of 506 protein sequences of the endosymbiont, including *Ca.* K. sorsogonicusi, the five previously sequenced *Ca.* Kinetoplastibacterium genomes, and two other Alcaligenaceae (*Achromobacter arsenitoxydans* and *Taylorella equigenitalis*) as outgroups has shown with strong support that the *Kentomonas* endosymbiont is indeed at the base of the tree for this genus. Thus, the phylogenies of the host and endosymbiont agree, leaving *A. ambiguus* as the only exception to the strict co-speciation between Strigomonadinae and *Ca.* Kinetoplastibacterium spp.

As recently shown (Alves et al. 2013a; Motta et al. 2013), the members of the genus Ca. Kinetoplastibacterium have highly reduced genomes in comparison with other bacteria from the same family. For instance, Taylorella, a parasite of the genital tract of horses, demonstrates a genome more than twice as big as those of Ca. Kinetoplastibacterium spp., whereas in Achromobacter, a typically free-living bacterium, the genome is around nine times larger. In the current work, we have characterized the most reduced Ca. Kinetoplastibacterium genome found to date, that of Ca. K. sorsogonicusi. With the length of 741 697 bp, it is about 10% (70-90 kbp) shorter than other known genomes within this genus (810 172-833 125 bp). No obvious explanation for the extra loss of sequence presents itself, given that Kentomonas and its endosymbiont conceivably possess a similar life cycle and overall environment as Angomonas and Strigomonas. Further in vivo and in vitro studies of this subfamily, and in particular this new genus, might shed light on the possible reasons for the differences seen here.

In spite of genome size reduction, with loss of many genes that are essential for free-living and sometimes even parasitic bacteria, the genomes of endosymbionts preferentially retain genes encoding proteins involved in the interaction with the trypanosomatid host, as typically seen in other endosymbiotic relationships (Nowack and Melkonian, 2010). Overall, analysis of the predicted metabolic maps shows that this also may be the case in *Ca*. K. sorsogonicusi, with a sole exception (see below).

As seen in the comparative analyses between other Kinetoplastibacterium genomes, gene loss in Ca. K. sorsogonicusi has been spread throughout the genome, without obvious hotspots of missing sequence (Fig. 1). Most gene losses observed do not significantly affect whole metabolic pathways, with the pointed exception of the haem-synthesis. As characterized in many biochemical and cell biology experiments over several decades (Chang et al. 1975; de Souza and Motta, 1999), and more recently by genomic methods (Alves et al. 2011; 2013b; Klein et al. 2013; de Azevedo-Martins et al. 2015), the endosymbiont provides the trypanosomatid host with a number of important compounds such as haem, essential amino acids, vitamins, and lipids. Accordingly, as seen in the comparative metabolic analyses, the Kentomonas endosymbiont possesses many such genes. The only exception is the synthesis of haem from glutamate (the C5 or Beale pathway), for which almost all genes were lost. The first enzyme in this pathway, glutamyl-tRNA synthetase (gltX), is not specific for the pathway and participates in protein synthesis by ligating L-glutamate to the corresponding tRNAs, its presence being therefore unsurprising. The other two retained genes (hemN and hemH) are specific to the haem-synthesis pathway.

Haem plays a very important role in the metabolism of virtually all cellular organisms, especially the aerobic ones, being a co-factor used by a number of haem proteins (Panek and O'Brian, 2002; Mense and Zhang, 2006). Accordingly, regular

Downloaded from https://www.cambridge.org/core. Shafer Library, on 31 Aug 2018 at 10:51:24, subject to the Cambridge Core terms of use, available at https://www.cambridge.org/core/terms. https://doi.org/10.1017/S003118201800046X trypanosomatids obtain haem or one of its precursors from their hosts, whereas most Strigomonadinae, as well as the unrelated bacteria-containing trypanosomatid *Novymonas esmeraldas*, consume the haem produced by their endosymbionts (Kořený *et al.* 2013; Kostygov *et al.* 2016; 2017; Horáková *et al.* 2017).

The genetic makeup of these organisms reflects their need for haem. Some trypanosomatids (from subfamilies Leishmaniinae and Strigomonadinae), possess the final three proteins of the haem-synthesis pathway; others (such as *Phytomonas* and *Herpetomonas*) contain only the last one, namely ferrochelatase; and, finally, *Trypanosoma* lost all the haem-synthesis-related genes (Kořený *et al.* 2013).

Of the haem-synthesis-specific genes, the *Kentomonas* endosymbiont has apparently kept only those coding for the oxygen-independent enzymes coproporphyrinogen oxidase III (*hemN*) and ferrochelatase (*hemH*). The function of *hemN* in this organism is unclear, given that it is an enzyme restricted to the haem-synthesis pathway, to the best of our knowledge. The *hemH* enzyme, on the other hand, can be used to add iron to (or remove it from) the protoporphyrin ring. Our search of the draft nuclear genome of *K. sorsogonicus* has not revealed any of the haem-synthesis-specific genes or even their fragments. Thus, the genomic situation for this organism is most similar to *Trypanosoma* spp. *In vitro*, our haem requirement experiment confirmed the genomic inference, demonstrating that *K. sorsogonicus* is unable to grow in the absence of a haem source. Hence, haem is an essential compound for this organism.

Although there are very few known organisms that do not require haem, the trypanosomatid Phytomonas is of particular interest. This plant parasite has been recently demonstrated as the first and so far the only eukaryote that is able to live in the absence of haem (Kořený et al. 2012). The only cellular process in which haem is necessary in Phytomonas is the synthesis of ergosterol, a component of the plasma membrane that is not obligatory for the cell's viability (its precursor, lanosterol, is used in its stead). Interestingly, Phytomonas still possesses the hemH gene that is lacking from K. sorsogonicus. However, the latter organism has been shown here to fail growing without a source of haem, and thus must acquire the compound from its insect host, as the other trypanosomatids routinely do. Therefore, Phytomonas continues to be the only known eukaryote that can propagate in culture without any source of haem. The loss of haem-synthesis in the Kentomonas system suggests that either the extra haem provided to Strigomonas and Angomonas by their endosymbionts might be dispensable (or essential) only for these two genera or that there is some particularity of Kentomonas that makes haem less critical to its survival and reproduction, allowing it to thrive just with the haem provided by its invertebrate host, as is the case with almost all other trypanosomatids.

In the absence of more detailed information on the ecological and physiological conditions faced by K. sorsogonicus and its endosymbiont, it is difficult to make any functional and evolutionary interpretation of the loss of the haem biosynthesis pathway in these organisms. We can only speculate about this based on comparative data. Kinetoplastids do not possess this pathway, likely because of the toxicity of haem precursors. The loss of haem synthesis is considered advantageous for many parasites (as reviewed in Kořený et al. 2013). The case of Kentomonas highlights the plasticity and adaptability of parasite genomes, which may change in response to environmental cues. The common ancestor of Strigomonadinae gained haem synthesis through endosymbiosis. While the bacterial symbionts of Angomonas and Strigomonas kept this capability, those of Kentomonas, which separated early from the rest of the subfamily, lost it. Such evolutionary plasticity, which is further exemplified by the ability of *Phytomonas* to live completely without haem, demonstrates that it is possible to lose, regain, and lose again the metabolic pathways for compounds currently considered essential.

For long, it has been a standard procedure in cultivating trypanosomatids isolated from insects to remove haemin from the culture medium; continued growth was considered as an evidence of endosymbiont presence. However, as we demonstrate here, despite bearing the intracellular bacterium, *K. sorsogonicus* cannot grow in culture without haem. Thus, the traditional test cannot be taken as a reliable criterion of the absence or presence of endosymbionts in trypanosomatid flagellates.

Supplementary material. The supplementary material for this article can be found at https://doi.org/10.1017/S003118201800046X

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References

- Alves JMP and Buck GA (2007) Automated system for gene annotation and metabolic pathway reconstruction using general sequence databases. *Chemistry & Biodiversity* 4, 2593–2602.
- Alves JMP et al. (2011) Identification and phylogenetic analysis of heme synthesis genes in trypanosomatids and their bacterial endosymbionts. PLoS ONE 6, e23518.
- Alves JMP et al. (2013a) Genome evolution and phylogenomic analysis of *Candidatus* Kinetoplastibacterium, the betaproteobacterial endosymbionts of *Strigomonas* and *Angomonas*. *Genome Biology and Evolution* **5**, 338–350.
- Alves JM et al. (2013b) Endosymbiosis in trypanosomatids: the genomic cooperation between bacterium and host in the synthesis of essential amino acids is heavily influenced by multiple horizontal gene transfers. BMC Evolutionary Biology 13, 190.
- Archibald JM (2015) Endosymbiosis and eukaryotic cell evolution. Current Biology 25, R911–R921.
- Camacho C et al. (2009) BLAST+: architecture and applications. BMC Bioinformatics 10, 421.
- Castresana J (2000) Selection of conserved blocks from multiple alignments for their use in phylogenetic analysis. *Molecular Biology and Evolution* 17, 540–552.
- Chang KP, Chang CS and Sassa S (1975) Heme biosynthesis in bacteriumprotozoon symbioses: enzymic defects in host hemoflagellates and complemental role of their intracellular symbiotes. *Proceedings of the National Academy of Sciences of the United States of America* 72, 2979–2983.
- Darling AE, Mau B and Perna NT (2010) Progressivemauve: multiple genome alignment with gene gain, loss and rearrangement. PLoS ONE 5, e11147.
- de Azevedo-Martins AC et al. (2015) Biochemical and phylogenetic analyses of phosphatidylinositol production in *Angomonas deanei*, an endosymbiont-harboring trypanosomatid. *Parasites & Vectors* 8, 247.
- de Souza W and Motta MC (1999) Endosymbiosis in protozoa of the Trypanosomatidae family. *FEMS Microbiology Letters* **173**, 1–8.
- Douglas AE (2016) How multi-partner endosymbioses function. Nature Reviews Microbiology 14, 731-743.
- Edgar RC (2004) MUSCLE: a multiple sequence alignment method with reduced time and space complexity. *BMC Bioinformatics* 5, 113.
- Gentil J et al. (2017) Review: origin of complex algae by secondary endosymbiosis: a journey through time. *Protoplasma* 254, 1835–1843.
- Hebert L et al. (2011) Genome sequence of Taylorella equigenitalis MCE9, the causative agent of contagious equine metritis. Journal of Bacteriology 193, 1785.
- Horáková E *et al.* (2017) The *Trypanosoma brucei* TbHrg protein is a heme transporter involved in the regulation of stage-specific morphological transitions. *Journal of Biological Chemistry* **292**, 6998–7010.

- Klein CC *et al.* (2013) Biosynthesis of vitamins and cofactors in bacteriumharbouring trypanosomatids depends on the symbiotic association as revealed by genomic analyses. *PLoS ONE* **8**, e79786.
- Kořený L, Lukes J and Oborník M (2010) Evolution of the haem synthetic pathway in kinetoplastid flagellates: an essential pathway that is not essential after all? *International Journal for Parasitology* **40**, 149–156.
- Kořený L et al. (2012) Aerobic kinetoplastid flagellate Phytomonas does not require heme for viability. Proceedings of the National Academy of Sciences of the United States of America 109, 3808–3813.
- Kořený L, Oborník M and Lukeš J (2013) Make it, take it, or leave it: heme metabolism of parasites. PLoS Pathogens 9, e1003088.
- Kostygov AY *et al.* (2016) Novel trypanosomatid-bacterium association: evolution of endosymbiosis in action. *mBio* 7, 1–12. doi: 10.1128/mBio.01985-15.
- Kostygov AY et al. (2017) Genome of Ca. Pandoraea novymonadis, an endosymbiotic bacterium of the trypanosomatid Novymonas esmeraldas. Frontiers in Microbiology 8, 1940.
- Krzywinski MI et al. (2009) Circos: an information aesthetic for comparative genomics. Genome Research 19(9), 1639–1645.
- Kück P and Meusemann K (2010) FASconCAT: convenient handling of data matrices. *Molecular Phylogenetics and Evolution* 56, 1115–1118.
- Kumar S, Stecher G and Tamura K (2016) MEGA7: molecular evolutionary genetics analysis version 7.0 for bigger datasets. *Molecular Biology and Evolution* 33, 1870–1874.
- Li L, Stoeckert Jr CJ and Roos DS (2003) OrthoMCL: identification of ortholog groups for eukaryotic genomes. *Genome Research* 13, 2178–2189.
- Li X et al. (2012) Genome sequence of the highly efficient arsenite-oxidizing bacterium Achromobacter arsenitoxydans SY8. Journal of Bacteriology 194, 1243–1244.
- Martin M (2011) Cutadapt removes adapter sequences from high-throughput sequencing reads. *EMBnet Journal* 17, 10–12.
- McCutcheon JP and von Dohlen CD (2011) An interdependent metabolic patchwork in the nested symbiosis of mealybugs. *Current Biology* 21, 1366–1372.
- Mense SM and Zhang L (2006) Heme: a versatile signaling molecule controlling the activities of diverse regulators ranging from transcription factors to MAP kinases. *Cell Research* 16, 681–692.

- Motta MCM et al. (2010) The bacterium endosymbiont of *Crithidia deanei* undergoes coordinated division with the host cell nucleus. *PLoS ONE* 5, e12415.
- Motta MCM et al. (2013) Predicting the proteins of Angomonas deanei, Strigomonas culicis and their respective endosymbionts reveals new aspects of the Trypanosomatidae family. PLoS ONE 8, e60209.
- Nowack ECM and Melkonian M (2010) Endosymbiotic associations within protists. *Philosophical Transactions of the Royal Society of London Series B, Biological Sciences* **365**, 699–712.
- Nussbaum K et al. (2010) Trypanosomatid parasites causing neglected diseases. Current Medicinal Chemistry 17, 1594–1617.
- Ogata H et al. (1999) KEGG: Kyoto encyclopedia of genes and genomes. Nucleic Acids Research 27, 29–34.
- Panek H and O'Brian MR (2002) A whole genome view of prokaryotic haem biosynthesis. *Microbiology* 148, 2273–2282.
- **Porcel BM** *et al.* (2014) The streamlined genome of *Phytomonas* spp. relative to human pathogenic kinetoplastids reveals a parasite tailored for plants. *PLoS Genetics* **10**, e1004007.
- Ronquist F et al. (2012) Mrbayes 3.2: efficient Bayesian phylogenetic inference and model choice across a large model space. *Systematic Biology* 61, 539–542.
- Seemann T (2014) Prokka: rapid prokaryotic genome annotation. Bioinformatics (oxford, England) 30, 2068–2069.
- Stamatakis A (2014) RAxML version 8: a tool for phylogenetic analysis and post-analysis of large phylogenies. *Bioinformatics (oxford, England)* 30, 1312–1313.
- Suzek BE et al. (2007) Uniref: comprehensive and non-redundant UniProt reference clusters. Bioinformatics (Oxford, England) 23, 1282–1288.
- Teixeira MMG et al. (2011) Phylogenetic validation of the genera Angomonas and Strigomonas of trypanosomatids harboring bacterial endosymbionts with the description of new species of trypanosomatids and of proteobacterial symbionts. Protist 162, 503–524.
- von Dohlen CD et al. (2001) Mealybug beta-proteobacterial endosymbionts contain gamma-proteobacterial symbionts. Nature 412, 433–436.
- Votýpka J et al. (2014) Kentomonas gen. n., a new genus of endosymbiontcontaining trypanosomatids of Strigomonadinae subfam. n. Protist 165, 825–838.