Morphological and molecular description of *Blastocritidia cyrtomeni* sp. nov. (Kinetoplastea: Trypanosomatidae) associated with *Cyrtomenus bergi* Froeschner (Hemiptera: Cydnidae) from Colombia

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A new trypanosomatid species, Blastocritidia cyrtomeni, is herein described using morphological and molecular data. It was found parasitising the alimentary tract of the insect host Cyrtomenus bergi, a polyphagous pest. The morphology of B. cyrtomeni was investigated using light and transmission microscopy and molecular phylogeny was inferred from the sequences of spliced leader RNA (SL rRNA) - 5S rRNA gene repeats and the 18S small subunit (SSU) rRNA gene. Epimastigotes of variable size with straphanger cysts adhering to the middle of the flagellum were observed in the intestinal tract, hemolymph and Malpighian tubules. Kinetoplasts were always observed anterior to the nucleus. The ultrastructure of longitudinal sections of epimastigotes showed the flagellum arising laterally from a relatively shallow flagellar pocket near the kinetoplast. SL RNA and 5S rRNA gene repeats were positive in all cases, producing a 0.8-kb band. The amplicons were 797-803 bp long with > 98.5% identity, indicating that they originated from the same organism. According to the sequence analysis of the SL-5S rRNA gene repeats and the 18S SSU rRNA gene, B. cyrtomeni is different from all other known species or isolates of Trypanosomatidae. Both analyses indicate that among known species, it is most closely related to Blastocritidia triatomae.

Key words: SL rRNA - 5S rRNA - SSU rRNA - trypanosomatids - *Cyrtomenus* - Blastocritidia cyrtomeni

Kinotoplastid flagellates (Euglenozoa and Kineto-plastea) are a mixed group of free-living organisms and mono and dixenous parasites that colonise a variety of eukaryotes, preferentially insects (Wallace 1966, Fernandes et al. 1993, Vickerman 1994, Podlipaev et al. 2004). The most studied kinetoplastids belong to the family Trypanosomatidae, which is characterised by a single flagellum and a relatively small kinetoplast, resembling a part of a single-branched mitochondrion of the cell containing a large mass of mitochondrial DNA (McGhee & Cosgrove 1980, Maslov et al. 1996, 2001, Merzlyak et al. 2001).

Trypanosomatids of the genera *Trypanosoma, Endotrypanum* and *Leishmania* are dixenous parasites of vertebrates (Fernandes et al. 1993, Merzlyak et al. 2001, Podlipaev 2001). The remaining members of the family include monoxenous parasites of invertebrates (genera *Blastocritidia, Crithidia, Leptomonas*, Herpetomonas and *Wallacea*) and dixenous parasites of plants and insects (genus *Phytomonas*) (Dollet et al. 2000, Maslov et al. 2001, Podlipaev et al. 2004). Since the early 1960s, life-cycle information combined with morphological features such as cell shape and dimension and the relative positions of the nucleus and kinetoplast have been used to develop the current taxonomic system (Hoare 1964, Hoare & Wallace 1966, Wallace 1966).

Although morphological characteristics are often reliable for identifying a genus, they are inadequate for identifying the species, mainly due to the high variability of morphology and their sensitivity to environmental factors or culture conditions (Merzlyak et al. 2001). As stated by Yurchenko et al. (2006), the validity of most of the described trypanosomatid species is questionable, with the notable exception of those of medical and veterinary importance (*Trypanosoma* and *Leishmania*), which have been studied extensively. Moreover, as shown by recent molecular phylogenetic studies (Maslov et al. 1996, Merzlyak et al. 2001, Podlipaev et al. 2004, Simpson et al. 2006, Yurchenko et al. 2006), most morphological genera are polyphyletic.

One of the main concerns in the field of trypanosomatid research is the scarcity of information about the diversity of this group. It has been estimated that of the approximately one million insect species described, only ~2,500 have been examined for the potential presence of trypanosomatids (Momen 2001, Podlipaev 2001). For instance, of approximately 23,000 identified species of Hemiptera, one of the most studied orders, only 500-600 have been examined for flagellate parasites (Teixeira et al. 2000). Moreover, the current system does not take into account genetic and evolutionary parameters; consequently, a great number of synonyms or descriptions of organisms of dubious origin have been generated (Podlipaev 2001).
To overcome the aforementioned taxonomic limitations, biochemical, ultrastructural, serological and nutritional approaches to discriminate and define taxa were proposed 20 years ago (Wallace et al. 1983). The most recent characterisations and analyses of biodiversity of new trypanosomatid species are based on the use of molecular methods and phylogenetic analysis of gene sequences such as spliced leader RNA (SL rRNA), 5S rRNA and glycosomal glyceraldehyde phosphate dehydrogenase, which have proven to be powerful for discriminating among genera and species (Podlipaev et al. 2004, Yurchenko et al. 2006, Maslov et al. 2007, Svobodová et al. 2007). Recently, small subunit (SSU) rRNA gene-based phylogenies have been used to establish major natural groups within a family (Merzlyak et al. 2001). Although useful in individual cases, these new criteria have not been applied uniformly (Podlipaev et al. 2004, Yurchenko et al. 2006). Therefore, it is highly advisable to use and compare at least two of these methods when a new species is described.

In recent studies, we observed that Cyrtomenus bergi Froeschner, a polyphagous Cydnidae pest, is naturally infected with trypanosomatid flagellates in their salivary glands, intestinal tract and hemolymph (AM Caicedo, unpublished observations). In this study, we report the isolation of the trypanosomatid parasites from C. bergi as well as their morphological description and molecular characterisation. Based on these analyses, this organism has been identified as a new species of Blastocrithidia.

**MATERIALS AND METHODS**

_**Maintenance of insect hosts** - Approximately 5,000 specimens of the burrowing bug C. bergi were collected in onion fields in Pereira (Risaralda, Colombia) from 2004-2008. The insects were maintained in an insectarium (23 ± 2°C, relative humidity 65 ± 5%, L12:D12) in sterilised loamy clay soil and sand, mixed in a 3:1 proportion, at a moisture level approximating field conditions (33.5% gravimetric soil water) and fed on sprouting maize kernels.

_**Light microscopy** - Hemolymph taken from fifth instar nymphs and adults was re-suspended in phosphate-buffered saline (PBS). Drops of the suspension were smeared on slides, methanol dried and stained with Wright-Giemsa at room temperature (RT) for 24 h. Slides were examined at 100X with a Nikon Microphot. Measures and photographs of different morph types were taken using an NIS elements Nikon camera (v. 2.3). Wright-Giemsa-stained microscope slides, representing gut smears of the original host infected with the Trypanosomatidae isolate and described herein as Blastocrithidia cyrtomeni, were deposited with the Department of Microbiology of the Universidad del Valle, Cali (Valle de Cauca Province, Colombia).

_**Electron microscopy** - Flagellates collected from the hindgut of C. bergi adults were centrifuged at 10,000 g, washed in 0.1 M PBS and fixed in 2.5% glutaraldehyde in 5 mM HCl and 0.1 M cacodylate buffer, first for 1 h at RT and then for at least 24 h at 4°C. Samples were post-fixed in 2% osmium tetroxide in the same buffer for 2 h at RT. After dehydration in an ethanol gradient series, the cells were embedded in resin. Thin ultramicrotomed sections (0.7-0.8 µm) were stained with lead citrate and uranyl acetate and examined under a JEM 1010 microscope.

_**DNA extraction, polymerase chain reaction (PCR) amplification and sequencing** - Pooled gut and hemolymph samples from five infected insects were homogenised and the debris was removed by centrifugation. Total DNA was extracted according to Rotureau et al. (2005) and Westenberger et al. (2004).

The SL rRNA gene repeats were amplified with the primers ME-1 5' - TTCTGATCTTTATGGTA and ME-2 5' - CAATAAAGTACAGAATCT (Podlipaev et al. 2004). The 5S rRNA gene repeats were amplified with the primers 5S-L 5' - CGTCCGATTGTGAAGTAAGC and 5S-R 5' - TAACCTCACAAATCGGAGGGAT. Amplifications were done with Taq polymerase. The thermal cycling profile for both 5S rRNA and SL RNA genes consisted of the following steps: initial denaturation at 94°C for 2 min, 40 cycles of denaturing at 94°C for 30 s, annealing at 50°C for 30 s and extension at 72°C for 30 s, followed by a final extension at 72°C for 12 min. SSU rRNA genes were amplified with the primers S-762 5’ - GACTTTGTGCCCTTCTA(T)TG and S-763 5’ - CATATGCTTGTCTCAGGAC (Maslov et al. 1996). Taq DNA polymerase was used for amplification and thermal cycling conditions were as follows: initial denaturation at 95°C for 5 min, 5 cycles at 95°C for 1 min, annealing at 45°C for 30 s and extension at 65°C for 1 min, 35 cycles at 95°C for 1 min, 50°C for 30 s and 72°C for 1 min, and a final extension at 65°C for 30 min (Maslov et al. 1996).

The amplified SL and SS products were extracted from agarose gel using a QIAquick gel extraction kit (QUIGEN, Valencia, CA) and cloned into the pGEM-T Easy Vector (Promega, Madison, WI). The clones were sequenced at a commercial facility (Macrogen, Korea). The SSU rRNA gene amplification product was gel purified by the same procedure and directly sequenced using a set of conserved internal primers (Maslov et al. 1996) at the University of California-Riverside IIGB Core Instrumentation facility.

The sequences were deposited in GenBank™ under the following accessions: SL RNA gene and SS rRNA gene repeat unit: FJ916991 and FJ916990 and SSU rRNA gene: FJ916992.

_**Phylogenetic analyses** - Prior to the analyses of SL RNA gene repeats, the regions corresponding to the amplification primers and most of the hypervariable intergenic regions were excluded (Westenberger et al. 2004). The remaining ~150 nt, including the exon, the intron and a relatively conserved sequence at positions -100 to -1 upstream of the exon, were aligned using CLUSTALX v. 1.81 (Thompson et al. 1997). Cluster analysis was performed using the neighbour-joining method and Kimura 2-parameter distances and PAUP* 4.0, beta version (Swofford 1998). The phylogenetic analyses of the SSU rRNA gene sequences included the most conserved regions, which were selected arbitrarily from the initial multiple alignment generated by CLUSTAL. The best-fitting model of the sequence evolution was selected us-
ing Modeltest, v. 3.06 (Posada & Crandall 1998), which corresponded to the general time-reversible model with a proportion of invariable sites I = 0.5029 and the gamma distribution shape parameter $\Gamma = 0.5416$. The analyses involved maximum likelihood, distance and parsimony implemented with PAUP* 4.0 (Swofford 1998).

RESULTS

Trypanosomatid infection (100% prevalence) was detected in \textit{C. bergi} adult populations collected in onion fields in La Florida, Pereira. However, no obvious morphological or behavioural symptoms of infection were observed in individual hosts. Light microscopy revealed trypanosomatids within the intestinal tract, hemolymph and Malpighian tubules. Attempts were made to establish a stable culture starting from hemolymph samples in several media, including NNN, LIT and Schneider, but these were not successful.

Light microscopy - Hemolymph and midgut smears revealed epimastigotes of variable size with straphanger cysts adhering to the middle of the flagellum. The cells varied in size and shape, from slender to rounded forms with short and long flagella. Some promastigotes could also be seen (Figs 1A, B, 8). The kinetoplast was always observed anterior to the nucleus.

Three forms of epimastigotes were seen: Form I, with a short flagellum and rounded body (Fig. 2A, B), Form II, with a slender body, pointed ends, a short flagellum and two to three spiral twists in the body (Fig. 3), and Form III, representing apparently dividing epimastigotes, with a second long flagellum separated from the cell body (Fig. 4).

The morphometry of the three main morph types observed is summarised in Table I. The most noticeable difference between Forms I and III was flagellum length, with the latter being longer than the former. These forms also showed at least two-three twists, and the posterior end of Form I was more pointed than Form III (Figs 1A, 3, 4).

Furthermore, the presence of flagellar cysts (strap-hangers) was observed in Forms I and III, with groups of four-five cysts adhering to the middle of the flagellum (Figs 1A, 5). In addition, relatively small cells, apparently generated by unequal division, were also observed as flagellar attachments (Figs 1A, 5), suggesting that they represent immature cysts. Cyst forms varied from small and round, to bacillus-like cells, and to those with anterior pointed ends. In all forms, the kinetoplast was anterior to the nucleus. The width of the mature strap-hangers ranged from 2.42-4.43 µm, the length ranged from 5.86-11.93 µm.

Transmission (TEM) and scanning (SEM) electron microscopy - The ultrastructure of longitudinal sections of epimastigotes isolated from midgut showed that the flagellum arises laterally from a relatively shallow flagellar pocket near the kinetoplast. The typical disk-shaped kinetoplast was located next to the bottom of the flagellar pocket (Fig. 6). We observed the longitudinal division of epimastigotes with kinetoplasts and bi-flagellum (Fig. 7). Few promastigotes were seen with the flagellum arising in front of the kinetoplast in the midgut (Fig. 8). Different forms of amastigotes with some acidocalcisomes and basophilic granules were also observed throughout the cell body (Fig. 9). The flagellum was supported by a nonprominent paraflagellar rod (Fig. 10). The parasites were attached to the intestine by the flagellum and different forms of parasites were observed (Fig. 11). In general, the nucleus was found mainly in the middle or close to the posterior end of the cell body, covering most of the cell width. The apparent absence of an undulated membrane, one of the features of epimastigotes, is noteworthy.
Using SEM, it was possible to observe numerous parasites adhered to the intestinal wall by their flagella (Fig. 12). Typical elongated epimastigotes with straphanger cysts were also observed on the midgut of *C. bergi* adults (Fig. 13).

**PCR amplification and phylogenetic analysis** - DNA was extracted from the midgut of *C. bergi* adults, fourth and fifth instar nymphs and eggs. PCR amplification of SL RNA gene repeats was positive in all cases, producing a 0.8 kb band corresponding to a monomeric repeat unit, often accompanied by higher molecular weight bands, possibly representing repeat dimers and trimers. Similarly, PCR amplification with SS rRNA primers resulted in a monomeric amplicon of the same size and higher molecular weight products that were apparently oligomeric (Fig. 14).

Cloning and sequencing of the SL RNA and SS rRNA monomeric amplicons showed that they have the same repeat unit, wherein individual SL RNA genes were interspersed with SS rRNA genes, forming a linkage arrangement often observed in trypanosomatids (Westenberger et al. 2004). Individual repeats, derived from more than 30 sequenced amplicons, were 797-803 bp long and >98.5% identical to each other, indicating that they all originated from the same organism (Thomas et al. 2005).

Seven individual SL repeat units were compared with the database of the repeats available from other Trypanosomatidae (Maslov et al. 2007) using neighbour-joining cluster analysis. The trypanosomatid from *C. bergi* clustered with members of the *B. triatomae-Blastocrithidia leptocoridis* clade (not shown). A part of this analysis, covering the *Blastocrithidia* group, is shown in Fig. 15. The repeats from the *C. bergi* parasite are unique and clearly distinct from *B. triatomae* (59.2% identity level) and *B. leptocoridis* (55.8% identity). The *C. bergi* parasite is most closely related to the organisms from Typing Unit 25, an unnamed trypanosomatid species found in two species of *Leptocorid* (Hemiptera: Coreidae) from Costa Rica and Ecuador. However, the relatively low repeat identity level (~63%) can clearly separate these two organisms.

The identification of the new parasite as *Blastocrithidia* was verified by the analyses of SSU rRNA gene sequences and comparison with selected members of the major known phylogenetic clades of the Trypanosomatidae. The general topology of the resulting trees was analogous to that derived in previous studies (Merzlyak et al. 2001, Yurchenko et al. 2006, 2008) (data not shown). In all analyses, the *C. bergi* parasites were found to be closely associated with *B. triatomae* (the only available representative of the aforementioned clade), with 100% bootstrap support in all the methods used (Fig. 16).

Based on the presence of epimastigotes and flagellar cysts and on the molecular phylogenetic analyses, we concluded that the parasite from *C. bergi* is a new trypanosomatid species that should be assigned to the genus *Blastocrithidia*.


**Type host** - Salivary glands, intestinal tract and hemolymph of *C. bergi* Froeschner, 1960 (Hemiptera: Cydnidae).

**Type locality** - La Florida, Pereira, Risaralda Province, Colombia (4°45'38.65"N 75°36'57.49"W).

**Type data** - Microscope slides (MUV 001-006), designated as hapantotype, were deposited in the type collection of the School of Microbiology, Universidad del Valle, Cali, Colombia.

**Genus assignment** - The presence of epimastigotes and promastigotes defines the placement of the new species in the genus *Blastocrithidia* Laird, 1959.

![Figs 12-13: scanning electronic microscopy of Blastocrithidia cyrtomeni sp. nov. in midgut of Cyrtomenus bergi adults. Most elongated forms adhered by flagellum to the intestine wall (12). Epimastigotes form with straphanger cysts and round body cell with long flagellum (13).](image)

### TABLE I

<table>
<thead>
<tr>
<th>Parasite</th>
<th>Length (µ) mean SE (n)</th>
<th>Width (µ) mean SE (n)</th>
<th>Posterior-nucleus (µ) mean SE (n)</th>
<th>Kinetoplast-nucleus (µ) mean SE (n)</th>
<th>Flagellum (µ) mean SE (n)</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>24.10 ± 9.24 (33)</td>
<td>4.24 ± 1.43 (33)</td>
<td>11.24 ± 6.28 (33)</td>
<td>4.37 ± 2.50 (33)</td>
<td>28.09 ± 13.18 (33)</td>
</tr>
<tr>
<td>II</td>
<td>53.55 ±12.92 (30)</td>
<td>3.62 ±0.64 (30)</td>
<td>20.70 ± 0.96 (30)</td>
<td>10.23 ± 3.60 (30)</td>
<td>36.33 ± 9.59 (30)</td>
</tr>
<tr>
<td>III</td>
<td>50.2 ±4.92 (31)</td>
<td>3.24 ±0.44 (31)</td>
<td>27.8 ± 4.63 (31)</td>
<td>10.07 ± 2.56 (31)</td>
<td>46.61 ± 9.23 (31)</td>
</tr>
<tr>
<td>Cyst and straphangers</td>
<td>9.36 ±1.5 (70)</td>
<td>3.50 ± 0.40 (70)</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>
Species characters - The species is mainly defined by molecular phylogenetic criteria. The new species is clearly distinguished from the other known trypanosomatids, including the closely related *B. triatomae*, by the SL RNA - 5S rRNA gene repeat sequences (GenBank™ accessions FJ916991, FJ916990).

Remarks - Epimastigotes often display straphanger cysts adhered to the middle of the flagellum in clusters of two-four. The species has proven to be refractory to cultivation in the media tested.

Etymology - The species name is derived from the insect host.

**DISCUSSION**

The new species of *Blastocrithidia* Laird, 1959, named *B. cyrtomeni* sp. nov., was isolated from hemolymph and the intestinal tract of *C. bergi*. Identification of this trypanosomatid was based on morphological descriptions using light and electron microscopic techniques and supported by analyses of molecular markers and phylogeny (Maslov et al. 1996, 2007, Merzlyak et al. 2001, Yurchenko et al. 2006).

The predominant form infecting fifth instar nymph and adults of *C. bergi* was elongated epimastigotes with straphanger cysts adhered to the middle of the flagellum. This characteristic is mentioned by Wallace (1966) in only five of thirty species of *Blastocrithidia* described:
Blastocrithidia familiaris Gibbs (1950), B. leptocoridis McCulloch (1915), Blastocrithidia orthae Uribe (1926), Blastocrithidia sandoni Gibbs (1951) and Blastocrithidia euchisti Hanson and McGhee, 1961 (Cerisola et al. 1971).

In B. triatomae, Cerisola et al. (1971) mentioned the cysts as a remarkable feature, describing them as small bodies of variable size hanging from the flagellum in most specimens.

As mentioned by Merzlyak et al. (2001), the original identification of Blastocrithidia gerricola from the gut of Gerris lacustris was based on the epimastigotes observed with some rare promastigotes. This is also the case with B. cyrtomeni, with epimastigotes being the predominant forms and few promastigotes observed.

We did not observe the “cytoplasmic bridge” mentioned by Schaub and Böker (1986) in their microscopic studies of B. triatomae, nor were we able to find cysts connected to the posterior end of epimastigotes as reported by the authors in three cases.

The posterior end of members of the Blastocrithidia species is a variable feature. In B. cyrtomeni, it was observed as needle-like in light SEM and TEM images (Figs 1B, 13). In contrast, Schaub and Böker (1986) observed that the broad posterior end was rolled up in a spiral in the first SEM images of B. triatomae and B. gerridis.

B. cyrtomeni has nearly the same body length as B. gerridis (Wallace 1966) and is almost double the average size of B. triatomae (Cerisola et al. 1971). The flagellum length is the most noticeable difference among these three species; it is shortest in B. gerridis and longest in B. cyrtomeni (Table II).

Common features among these three Blastocrithidia species are the position of the kinetoplast, which is anterior to the nucleus, and the width of the nucleus, which is almost the same as that of the body (Wallace 1966, Cerisola et al. 1971).

An undulated membrane in the epimastigotes was not observed. Thus far, only small forms of B. leptotrypanoides (Hollande 1922) have been reported without an undulating membrane (Wallace 1966).

In Leptomonas wallacei, a trypanosomatid from the lygaeid bug Oncopeltus fasciatus, straphanger cysts adhere to the beginning of the base of the flagellar pocket (Romeiro et al. 2001). The dimensions of the straphangers of B. cyrtomeni are almost three times larger in length and width (9.36 and 3.50 µm, respectively), than those of B. triatomae (3.4 and 1.9 µm, respectively) (Cerisola et al. 1971).

Although we observed oval cells of B. cyrtomeni adhered to the epithelium of the insect’s midgut (Fig. 11) but free in the hemolymph, we were unable to elucidate whether these forms represent intermediate developmental stages toward the formation of draught-resistant cysts, such as in B. triatomae, a parasite of Triatominae bugs (Schaub et al. 1990).

We also do not know whether the host mounts a response aimed at reducing the growth rate of the parasite population. There is no evidence of a successful response to eliminate the parasite from the body of the host (complete recovery), but it is likely that the insect regulates the parasite population at some permanent steady-state level. We found a special case of direct transmission (transovarial), in which the infection is transferred by a parent to its unborn offspring. Therefore, despite the fact that B. cyrtomeni has an extremely high rate of direct reproduction within the host, it is unlikely that this microparasite is a pathogen.

Using PCR amplification of SL rRNA genes from 16 samples of parasites isolated from the intestinal tract and eggs of C. bergi, it was possible to identify B. cyrtomeni sp. nov. using the approach proposed by Maslov et al. (2007), who established that a single repeat of SL is sufficient to serve as a marker for a natural clone or species of trypanosomatid and to distinguish among closely related trypanosomatids.

It is critical to establish the presence and prevalence of these monoxenous trypanosomatids in other insect species and to evaluate in greater detail whether the association between insect pests and trypanosomatids could be regulating the action of microorganisms used for biological control.

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