Multi-locus sequence analysis unveils a novel genus of filarial nematodes associated with ticks in French Guiana

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Received 24 January 2024, Accepted 27 February 2024, Published online 15 March 2024

Abstract — Filarial nematodes of the Dipetalonema lineage include tick-borne filarialoids that infect both domestic and wild vertebrate hosts, but they remain understudied in many cases. In this study, we conducted a molecular characterization of a Dipetalonema-like filarialoid (DLF) recently identified in two tick species in French Guiana, South America. While the cox1 mitochondrial gene was the sole marker initially sequenced for describing DLF, its classification and phylogenetic relationship with other members of the Dipetalonema lineage were unclear. Therefore, we better characterized DLF through the sequencing of six additional gene markers and conducted phylogenetic analyses. Based on this multi-locus typing scheme, DLF exhibited significant divergence from known genera and species of filarialoids, or other sequences available in public databases, suggesting its potential classification as a novel genus within the Dipetalonema lineage. Phylogenetic analyses further unveiled a close evolutionary relationship between DLF and all other filarialoids associated with Acari (ticks and mites) within a robust monophyletic subclade in the Dipetalonema lineage. Overall, these findings confirm the existence of a specialized, Acari-borne group of filarialoids and underscore the need for comprehensive investigations into their epidemiology and potential impact on animal health.

Key words: Filarial nematodes, Dipetalonema, tick, French Guiana.

Résumé — Une analyse de séquences multi-locus dévoile un nouveau genre de nématodes filaires, associé aux tiques en Guyane française. Les filaires de la lignée Dipetalonema comprennent des espèces transmises par les tiques qui infectent à la fois des hôtes vertébrés domestiques et sauvages, mais qui restent sous-étudiés dans de nombreux cas. Dans cette étude, nous avons réalisé une caractérisation moléculaire d’un filarioïde ressemblant à Dipetalonema (FRD) récemment identifié dans deux espèces de tiques en Guyane française. Alors que la séquence du gène mitochondrial cox1 était le seul marqueur génétique initialement séquencé pour décrire FRD, sa classification et sa relation phylogénétique avec d’autres membres de la lignée Dipetalonema étaient incertaines. Par conséquent, nous avons caractérisé plus précisément DLF en séquençant six gènes supplémentaires et en réalisant des analyses phylogénétiques. Sur la base de ce typage multi-locus, FRD présentait une divergence significative par rapport aux genres et espèces connus de filarioïdes, ou à d’autres séquences disponibles dans les bases de données publiques, suggérant sa classification potentielle en tant que nouveau genre au sein de la lignée Dipetalonema. Les analyses phylogénétiques ont en outre révélé une relation évolutive étroite entre FRD et tous les autres filarioïdes associés aux tiques et acariens au sein d’un sous-clade monophylétique dans la lignée Dipetalonema. Dans l’ensemble, ces résultats confirment l’existence d’un groupe spécialisé de filarioïdes transmis par les tiques et acariens et soulignent la nécessité d’études approfondies sur leur épidémiologie et leur impact potentiel sur la santé animale.

Introduction

Ticks are vectors of major viruses, bacteria, and protozoan parasites of medical and veterinary significance [24, 33, 34]. However, surveys of tick-borne pathogens often neglect filarial nematodes from the family Onchocercidae, commonly referred to as filariae or filarialoids, while these parasites are regularly detected in most tick genera [1, 4, 6, 10, 11, 17, 29, 36, 39, 43, 45, 46, 50, 51]. Microscopic observations and molecular typing consistently categorize most tick-associated filarialoids into the genera Acanthocheilonema, Monanema, Yatesia, and Cerco-pithifilaria [1, 4, 6, 10, 11, 17, 29, 35, 36, 39, 43, 45, 46, 50, 51], although most were initially classified in the genus Dipetalonema [7, 9, 23]. Phylogenetic analyses based on molecular and morphological data further showed that the genera Acanthocheilonema, Monanema, Yatesia, and Cerco-pithifilaria (all associated with ticks), as well as Cruorifilaria (not yet
associated with a vector, but detected in ticks [17]), Lithosnooides (associated with parasitic mites) and Dipetalonema (associated with biting midges), cluster in a monophyletic clade of filarioids, termed the Dipetalonema lineage or the ONC4 clade, within the family Onchocercidae [7, 9, 23, 36].

Experimental infection assays and field observations have confirmed tick vector competence for filarioids of the genera Acanthocheilonema, Monanema, Yatevia, and Cercopithifilaria. Ticks feeding on infected vertebrates ingest microfilariae, which can develop up to the viable infective stage in a few weeks and are further excreted with saliva during biting, establishing specific tick-borne infection cycles [3–5, 8, 10, 13–15, 20, 32, 38, 40, 41, 44, 49, 52, 53]. This vector competence extends to major tick genera, including Ixodes, Rhipicephalus, Amblyomma, Haeaphyphalis, Hyalomma, and Ornithodoros, further emphasizing the effectiveness of ticks as vectors for filarioids of the Dipetalonema lineage [3–5, 8, 10, 13–15, 20, 32, 38, 40, 41, 44, 49, 52, 53]. These filarioids also survive transstadially in ticks since the development from microfilariae to infective larvae occurs only while the tick is off-host, that is, during ecdisis from tick larva to nymph or from nymph to adult [41, 49].

In a recent survey of ticks in French Guiana, South America, molecular analysis and phylogenetic studies revealed the presence of novel filarioids belonging to the Dipetalonema lineage in several tick species [17]. Based on cytochrome c oxidase subunit I (cox1) mitochondrial gene sequences, all but one of these filarioids are distinct to already known species of Dipetalonema lineage. Indeed, in the Cayenne tick Amblyomma cajennense (Fabricius, 1787) and in the opossum tick Ixodes luciae Sénévet, 1940, one filarioid, provisionally named Dipetalonema-like (DLF hereafter), showed a cox1 gene sequence substantially divergent from other species and genera of the Dipetalonema lineage [17]. DLF could be of health concern since it was divergent from other species and genera of the Dipetalonema lineage [17]. The predominant tick species bit- ing humans in South America [16]. This feature may not apply to I. luciae, as it is a specialized tick species with a primary feeding preference for opossums [16]. However, while DLF has been detected in 6% of field specimens of A. cajennense [17], no further data are currently available on this filarioid.

In this study, we conducted an extended molecular characteri- zation of DLF previously detected in A. cajennense and I. luciae in French Guiana. The cox1 gene sequence was the only genetic marker used for its description [17], but this marker exhibits limited resolution for inferring the evolutionary history in the family Onchocercidae [36]. Using infected field specimens of A. cajennense and I. luciae, we thus characterized DLF through the sequencing of six additional genes (MyoHC, hsp70, rhp1, 12S rRNA, 28S rRNA, and 18S rRNA) previously used for inferring the Onchocercidae phylogeny [36]. We further examined their genetic proximity with other filarioid species, including all known members of the Dipetalonema lineage, under a phylogenetic framework.

## Materials and methods

### Tick collection

A collection of 10 DNA templates from A. cajennense (n = 8) and I. luciae (n = 2) infected by DLF was used for the present analysis. All templates were obtained from field specimens collected on vegetation through flagging (questing ticks) or on opossums (engorged ticks) in French Guiana in 2016 and 2017 (Table 1). Each DNA template was obtained from individual extraction of tick whole body using a DNeasy Blood and Tissue Kit (QIAGEN, Hilden, Germany), following manufacturer instructions. For each DNA template, infection by DLF had previously been confirmed through cox1 gene sequencing [17]. Use of the genetic resources was approved by the French Ministry of the Environment under reference #TREL.19028117S/156, in compliance with the Access and Benefit Sharing procedure implemented by the Loi pour la Reconquête de la Biodiversité.

### Multi-locus typing of the Dipetalonema-like filarioid

Segments of six genes (MyoHC, hsp70, rhp1, 12S rRNA, 28S rRNA, and 18S rRNA) were amplified using simple, semi-nested or nested PCR assays adapted from Lefoulon et al. [36]. Gene features, primers and PCR conditions are detailed in Table S1. Simple PCR amplifications were performed in a total volume of 25 μL containing ca. 20 ng of genomic DNA, 8 mM of each dNTP (Thermo Scientific, Waltham, MA, USA), 10 mM of MgCl₂ (Thermo Scientific), 7.5 μM of each of the internal primers, 2.5 μL of 10× PCR buffer (Thermo Scientific), and 1.25 U of Taq DNA polymerase (Thermo Scientific). Nested and semi-nested PCR amplifications were performed as follows: the first PCR run with the external primers was performed in a 10 μL volume containing ca. 20 ng of genomic DNA, 3 mM of each dNTP (Thermo Scientific), 8 mM of MgCl₂ (Thermo Scientific), 3 μM of each primer, 1 μL of 10× PCR buffer (Roche Diagnostics), and 0.5 U of Taq DNA polymerase (Roche Diagnostics). A 1 μL aliquot of the PCR product from the first reaction was then used as a template for the second round of amplification. The second PCR was performed in a total volume of 25 μL and contained 8 mM of each dNTP (Thermo Scientific), 10 mM of MgCl₂ (Thermo Scientific), 7.5 μM of each of the internal primers, 2.5 μL of 10× PCR buffer (Thermo Scientific), and 1.25 U of Taq DNA polymerase (Thermo Scientific).

## Table 1. List and origin of DLF-infected tick specimens examined in this study.

<table>
<thead>
<tr>
<th>Host species</th>
<th>Sampling location, date</th>
<th>Stage, sex</th>
<th>Feeding status</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cayenne tick, A. cajennense</td>
<td>Kourou, 2016</td>
<td>1 nymph, 1 adult (female)</td>
<td>Questing ticks</td>
</tr>
<tr>
<td></td>
<td>Matoury, 2016</td>
<td>6 nymphs</td>
<td>Questing ticks</td>
</tr>
<tr>
<td>Opossum tick, I. luciae</td>
<td>Cayenne, 2017</td>
<td>2 adults (1 male, 1 female)</td>
<td>Engorged ticks (collected on opossum)</td>
</tr>
</tbody>
</table>
Table 2. Best nucleotide identities of DLF MyoHC, hsp70, rbp, 12S rRNA, 28S rRNA, and 18S rRNA gene sequences obtained in this study with sequences available in GenBank.

<table>
<thead>
<tr>
<th>Gene</th>
<th>Best matches in public databases (% nucleotide identity, GenBank accession number)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MyoHC</td>
<td>Acanthocheilonema viteae (91.77%, KP760213); Acanthocheilonema odendhali (91.07%, KP760212)</td>
</tr>
<tr>
<td>hsp70</td>
<td>Monanema martini (85.01%, KP760443); Acanthocheilonema viteae (84.71%, KP760411)</td>
</tr>
<tr>
<td>rbp1</td>
<td>Dipetalonema robini (92.57%, KP760280); Dipetalonema caudispina (92.37%, KP760274)</td>
</tr>
<tr>
<td>12S rRNA</td>
<td>Micipsella ibérica (83.18%, MW928503); Onchocercidae sp. GK-2015 (82.52%, KR676614)</td>
</tr>
<tr>
<td>28S rRNA</td>
<td>Dipetalonema gracile (97.45%, KP760371); Dipetalonema robini (97.09%, KP760374)</td>
</tr>
<tr>
<td>18S rRNA</td>
<td>Yatesia hydrochoeris (98.94%, KP760166); Cercopithifilaria rugosicauda (98.94%, KP760124)</td>
</tr>
</tbody>
</table>

All PCR amplifications were performed under the following conditions: initial denaturation at 94 °C for 3 min, cycles of denaturation (35–40 cycles, depending on gene fragment size) (94 °C, 30 s), annealing (Tm = 50–55 °C, depending on primers, 30 s), extension (72 °C, 1 min–1 min 30 s, depending on gene fragment size), and a final extension at 72 °C for 5 min (Table S1). To prevent possible contamination, first and second PCR runs were physically separated from one another, in entirely separate rooms. Negative (water) controls were included in each PCR assay. All PCR products were visualized through electrophoresis in a 1.5% agarose gel. All amplicons were purified and sequenced in both directions (EUROFINS, Luxembourg). Sequence chromatograms were cleaned with Chromas Lite (http://www.techneilum.com.au/chromas_lite.html), and alignments were performed using ClustalW, implemented in the MEGA software package (https://www.megasoftware.net/). New sequences obtained in this study were deposited in GenBank under accession numbers PP182382–PP182391 (MyoHC), PP182371–PP182380 (hsp70), PP182391–PP182401 (rbp1), PP196371–PP196380 (12S rRNA), PP196417–PP196426 (28S rRNA), and PP196384–PP196393 (18S rRNA).

Molecular phylogenetic analyses

Phylogenetic analyses were based on sequence alignments of the filarioid MyoHC, hsp70, rbp1, 12S rRNA, 28S rRNA, and 18S rRNA gene sequences obtained in this study. Analyses also included the filarioid cox1 gene sequences (GenBank accession numbers OR030080–OR030087, OR030094, and OR030095) previously obtained from the same A. cajennense and I. luciae specimens by Binetruy and Duron [17]. Sequences of other filarioids obtained from GenBank, including representative members of the Dipetalonema lineage (Acanthocheilonema, Yatesia, Cercopithifilaria, Cuorifilaria, Litomosoides, and Dipetalonema) and of other filarial nematodes were also included in the phylogenetic analyses (Table S2). The Basic Local Alignment Search Tool (BLAST; https://blast.ncbi.nlm.nih.gov/blast/Blast.cgi) was used to find additional sequences available on the GenBank. The Gblocks program with default parameters was used to obtain non-ambiguous sequence alignments [22]. Tree-based phylogenetic analyses were performed using maximum-likelihood (ML) analyses using the MEGA software package (https://www.megasoftware.net/). The evolutionary models that best fit the sequence data were determined using the Akaike information criterion. Clade robustness was assessed by bootstrap analysis using 1,000 replicates. We further conducted a phylogenetic network analysis based on uncorrected P distances using the Neighbor-net algorithm [21] implemented in SPLITSTREE [30]. The resulting phylogenetic networks generalize the trees by allowing cross-connections between branches, which might display conflicting signals in the phylogenetic data set [21].

Results

Multi-locus typing of the Dipetalonema-like filarioid

The DLF MyoHC, hsp70, rbp1, 12S rRNA, 28S rRNA, and 18S rRNA gene sequences were amplified from the 10 DNA templates (A. cajennense, n = 8; I. luciae, n = 2). All sequences were easily readable without double peaks, indicating a confident degree of primer specificity for filarioid PCR amplifications. On the basis of DNA sequencing, we characterized only one allele for each of the six genes. The DLF MyoHC, hsp70, rbp1, 28S rRNA, and 18S rRNA gene sequences were distinct from sequences available in public databases, and showed 83.2–98.9% pairwise nucleotide identities (depending on gene sequence) with other members of the Dipetalonema lineage, including Acanthocheilonema, Monanema, Yatesia, and Cercopithifilaria spp. (Table 2). Comparisons with filarioids other than members of the Dipetalonema lineage showed lower pairwise nucleotide identities for these gene sequences although the 12S rRNA sequences exhibited the highest pairwise nucleotide identities with filarioids of uncertain phylogenetic position (Table 2).

Phylogeny of the Dipetalonema-like filarioid

ML and phylogenetic network analyses based on MyoHC (717 bp), hsp70 (561 bp), rbp1 (500 bp), 12S rRNA (470 bp), 28S rRNA (436 bp), 18S rRNA (660 bp), and cox1 (649 bp) nucleotide sequences were further conducted to examine the phylogenetic proximity of DLF with other filarioids. For any given gene, ML estimations gave similar tree topologies with minor differences, but also harbored some polytomies at any given gene, ML estimations gave similar tree topologies with minor differences (Figs. 1 and 2). All phylogenetic reconstructions revealed a clustering of DLF with the genera Cercopithifilaria, Cuorifilaria, Litomosoides, Yatesia, Acanthocheilonema,
Monanema, and Dipetalonema in a single monophyletic clade, the Dipetalonema lineage, distinct from other members of the family Ochocercidae. Phylogenetic reconstructions further revealed the division of the Dipetalonema lineage into two monophyletic subclades supported by high bootstrap values (Figs. 1 and 2).

1. The first subclade comprised DLF and members of the genera Acanthocheilonema, Monanema, Cercopithifilaria, Yatesia, Cruorifilaria, and Litomosoides. Within this subclade, DLF formed a branch substantially divergent from all other genera, but was more related to Acanthocheilonema species. Remarkably, all these filarioids are naturally associated with Acari, either with ticks (for DLF, Acanthocheilonema, Monanema, Cercopithifilaria, Yatesia, and Cruorifilaria), or with parasitic mites (for Litomosoides).

2. The second subclade comprised only Dipetalonema spp., which are filarioids specifically associated with biting midges.

As a result, the phylogenetic partitioning of the Dipetalonema lineage into two monophyletic subclades correlates with specialization for distinct types of arthropod vectors, Acari vs. dipterans.

Discussion

In this study, we show that DLF displays substantial differences in its gene sequences compared to known genera and species within the family Ochocercidae, as well as any other sequences available in public databases. This observation lends support to the hypothesis that it could be a novel genus of
filarioid with the Dipetalonema lineage. Furthermore, phylogenetic analyses unveil a close evolutionary relationship between DLF and all other filarioids associated with Acari (ticks and mites): these filarioids cluster together in a robust monophyletic subclade within the Dipetalonema lineage. These findings, consistent with earlier observations by Lefoulon et al. [36], suggest the presence of a monophyletic group of filarioids that has evolved a specialization for Acari as specific vectors.

Analyses of DNA gene sequence similarities and phylogenetics both confirm that DLF is divergent from other members of the Dipetalonema lineage. No morphological data are currently available for DLF but it may share morphological similarities with other members of the Dipetalonema lineage. Adults of these species have a long tail, a buccal capsule divided into two (or three) segments, more or less atrophied for specialized species, and a caudal extremity with two subterminal lappets [7, 23, 32]. Interestingly, opossums could be vertebrate hosts of DLF since I. luciae is a specialized tick species that feeds primarily on opossums [16]. Under this assumption, DLF may have previously been observed in opossums: previous studies have identified four filarioid species, all showing typical morphological features of the Dipetalonema lineage, i.e., Acanthocheilonema pricei, Cercopithifilaria didelphis, Skrjabinofilaria skrjabini, and Cherylia guianensis, in South American opossums [8, 27]. However, only morphological data, and no molecular data, are currently available for these four filarioid species, which prevents us from concluding whether one of these already described species is a DLF.

The clustering of all Acari-associated filarioids in a monophyletic subclade, separate from those transmitted by blood-feeding dipterans, strengthens the conclusion that ticks serve as specific vectors for certain filarioids. It also implies that these filarioids are well adapted to tick physiology, life-cycle and behavior. Earlier experimental assays have confirmed that ticks are competent vectors of filarioids of the Dipetalonema lineage [3, 5, 20, 33, 39, 40, 43, 48, 51, 52]. These observations include the filarioid Cherylia guianensis, primarily isolated from a gray and black four-eyed opossum, which can normally develop in Ixodes ticks [8]. Furthermore, the detection of DLF from questing (unfed) A. cajennense ticks that have already digested their previous blood meals, have further moulted, and are seeking vertebrates for their next blood meal suggests that A. cajennense can acquire and stably maintain infection through transstadial transmission [17], as also observed for other members of the Dipetalonema lineage [40, 48]. For animals, the risk of acquiring a DLF infection is currently unknown, but surveys of dogs and capybaras infected by other filarioids of the Dipetalonema lineage revealed skin issues,
chronic polyarthritis, anemia, and kidney and pulmonary damage [18, 19, 25, 26, 28, 41].

In conclusion, ticks transmit a broader range of infectious agents than any other arthropod vector, but their role as vectors of filarial diseases is less well-documented. The recurring identification of the *Dipetalonema* lineage species in major tick genera on most continents [2, 12, 17, 19, 31, 35, 37, 42, 46, 47] confirms that these are widespread but overlooked tick-borne parasites. Further research is needed to understand their pathogenicity, epidemiology, developmental cycles, and transmission mechanisms by ticks, including DLF in *A. cajennense* and *I. luciae*.

Acknowledgements

This work was supported by “Investissements d’Avenir” managed by the Agence Nationale de la Recherche (ANR, France: Laboratoire d’Excellence CBEA, ref. ANR-10-LABX-25-01).

Conflicts of interest

The authors declare no conflict of interest.

Supplementary material

The Supplementary material of this article are available at https://www.parasite-journal.org/10.1051/parasite/2024015/olm.

Table S1: List and description of primers used in this study for DLF molecular typing (adapted from Lefoulon et al., 2015).

Table S2: List of GenBank accession numbers for *MyoHC*, bsp70, rlp, 12S rRNA, 28S rRNA, and 18S rRNA gene sequences used in phylogenetic analyses.

References


23. Chabaud AG, Bain O. 1976. *L. en* *laria d*S. *laria d* *D* pi *laria d* *D* auca *laria d* *D* 552.


