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Molecular characterization and distribution of *Plasmodium matutinum*, a common avian malaria parasite

GEDIMINAS VALKIŪNAS*, MIKAS ILGŪNAS, DOVILĖ BUKAUSKAITĖ, VAIDAS PALINAUSKAS, RASA BERNOTIENĖ and TATJANA A. IEZHOVA

Nature Research Centre, Akademijos 2, Vilnius 2100, LT-08412, Lithuania

(Received 3 February 2017; revised 25 April 2017; accepted 26 April 2017; first published online 21 September 2017)

SUMMARY

Species of *Plasmodium* (Plasmodiidae, Haemosporida) are widespread and cause malaria, which can be severe in avian hosts. Molecular markers are essential to detect and identify parasites, but still absent for many avian malaria and related haemosporidian species. Here, we provide first molecular characterization of *Plasmodium matutinum*, a common agent of avian malaria. This parasite was isolated from a naturally infected thrush nightingale *Luscinia luscinia* (Muscicapidae). Fragments of mitochondrial, apicoplast and nuclear genomes were obtained. Domestic canaries *Serinus canaria* were susceptible after inoculation of infected blood, and the long-lasting light parasitemia developed in two exposed birds. Clinical signs of illness were not reported. Illustrations of blood stages of *P. matutinum* (pLINN1) are given, and phylogenetic analysis identified the closely related avian *Plasmodium* species. The phylogeny based on partial cytochrome *b* (cyt *b*) sequences suggests that this parasite is most closely related to *Plasmodium tejerai* (cyt *b* lineage pSPMAG01), a common malaria parasite of American birds. Both these parasites belong to subgenus *Haemamoeba*, and their blood stages are similar morphologically, particularly due to marked vacuolization of the cytoplasm in growing erythrocytic meronts. Molecular data show that transmission of *P. matutinum* (pLINN1) occurs broadly in the Holarctic, and the parasite likely is of cosmopolitan distribution. Passeriform birds and *Culex* mosquitoes are common hosts. This study provides first molecular markers for detection of *P. matutinum*.

Key words: Avian malaria, Plasmodium matutinum, molecular characterization, distribution, vectors, birds.

INTRODUCTION

Avian malaria is caused by over 50 described species of *Plasmodium* (Plasmodiidae, Haemosporida), which are of global distribution (Bennett *et al.* 1982; Atkinson *et al.* 2008; Marzal, 2012; Perkins, 2014). Diversity of these parasites is greatest in tropics, but some *Plasmodium* species are actively transmitted in temperate climates and have been recently reported even in high arctic tundra (Valkiūnas, 2005; Oakgrove *et al.* 2014). Birds suffer of malaria all over the world, but this disease is absent from Antarctica, most likely because of the absence of suitable vectors and insufficient air temperature for completing of sporogonic development.

Avian malaria is often more severe disease than mammal malaria (Garnham, 1966; Gabaldon and Ulloa, 1980; Ilgūnas *et al.* 2016). In birds, species of *Plasmodium* and related haemosporidian parasites cause not only blood pathology due to high parasitaemia, which is the main reason of pathology during malaria in mammals (Sherman, 1998), but also due to damage of various organs by exoerythrocytic meronts, which develop in reticular-endothelial cells and can be present all over the body in susceptible avian hosts (Garnham, 1966; Valkiūnas, 2005;

difficult, and the majority of recent wildlife studies deal only with the relatively easy to access blood stages of these parasites (Bensch et al. 2009; Clark et al. 2014; Perkins, 2014). However, the application of histological and chromogenic in situ hybridization methods provided evidence that common Plasmodium parasite lineages cause disease and even mortality in birds due to damage of organs by the exoerythrocytic stages (Dinhopl et al. 2015; Ilgūnas et al. 2016; Palinauskas et al. 2016). It is important to note that mortality due to internal organ pathology may occur before development of parasitemia, and such haemosporidian infections are particularly difficult to diagnose using microscopic tools (Donovan et al. 2008; Valkiūnas et al. 2014; Valkiūnas and Iezhova, 2017). If tissue stages are reported by microscopic examination in dead birds, it is often difficult or even impossible to identify species, subgenera and sometimes even families of reported haemosporidian parasites when solely morphological characters are available (Garnham, 1966; Walker and Garnham, 1972; Valkiūnas et al. 2014). The exoerythrocytic development remains insufficiently investigated in the majority of avian haemosporidian species because of the difficulties to access and identify reported parasites in organs, particularly in wild animals. Molecular markers are essential for rapid and precise diagnostic of avian malaria, but

Vanstreels et al. 2015; Ilgūnas et al. 2016). Detection

and identification of haemosporidian tissue stages are

Parasitology (2017), **144**, 1726–1735. © Cambridge University Press 2017 doi:10.1017/S0031182017000737

^{*} Corresponding author: Nature Research Centre, Akademijos 2, LT-08412 Vilnius, Lithuania. E-mail: gedvalk@ekoi.lt

remain unknown for the majority of described *Plasmodium* and other related haemosporidian species (Bensch *et al.* 2009; Clark *et al.* 2014). That is an obstacle for detection of malaria and other wild-life haemosporidian infections. The same parasite diagnostic problems remain during examination of sporogonic stages of haemosporidians in wild-caught vectors (Valkiūnas *et al.* 2013*a, b*). Haemosporidian molecular markers are needed for the veterinary parasitology research (Dinhopl *et al.* 2015), and they are essential for evolutionary biology studies as well during determining cryptic speciation in these parasites (Outlaw and Ricklefs, 2014; Palinauskas *et al.* 2015; Bensch *et al.* 2016; Nilsson *et al.* 2016).

During an on-going research on evolutionary biology, ecology and virulence of avian blood parasites (Dimitrov et al. 2016; Ilgūnas et al. 2016; Palinauskas et al. 2016; Valkiūnas et al. 2016), Plasmodium matutinum was found in the thrush nightingale Luscinia luscinia (Muscicapidae) during spring migration in Lithuania. According to the former microscopic studies (Garnham, 1966; Valkiūnas, 2005), this parasite was reported in the Holarctic birds, but its molecular characterization is absent. The aims of this study were (1) to develop molecular characterization of *P. matutinum*, (2) to determine its phylogenetic relationships with other avian *Plasmodium* spp. and (3) to specify the host range and distribution of this parasite lineage.

MATERIAL AND METHODS

Collection of blood samples

In all, 11 thrush nightingales *L. luscinia* (Muscicapidae) were caught with mist nets and large stationary traps at the Ventės Ragas Ornithological Station, Lithuania between 11 and 27 May, 2016. About $30 \,\mu\text{L}$ of whole blood was taken in heparinized microcapillaries and stored in SET buffer (Hellgren *et al.* 2004) for molecular analysis. Three blood films were prepared, air-dried, fixed in absolute methanol and stained with Giemsa (Valkiūnas, 2005).

One blood film from each bird was rapidly stained and examined microscopically, as described by Valkiūnas *et al.* (2016). One nightingale infected with an unidentified lineage of *P. matutinum* was determined and used for the strain isolation during this study. Infected blood was collected in heparinized microcapillaries and used to expose two uninfected domestic canaries by subinoculation of about 250 μ L of freshly prepared mixture, containing the infected blood and anticoagulant in the pectoral muscle, as described by Iezhova *et al.* (2005). Blood samples from donor nightingale were also cryopreserved for future research, as described by Garnham (1966). The donor nightingale was released.

Both recipient canaries were proven to be uninfected with haemosporidian parasites by microscopic examination of blood films and polymerase chain reaction (PCR)-based testing in the laboratory (see below). These birds were kept indoors in a vector-free room for a 4-month period postexposure. Blood was taken for microscopic examination and PCR-based testing as described above 5 days postexposure (dpe) for the first time, then once per 3 or 4 days during the first 2 months postexposure and once per month during the remaining experiment time. Parasitemia developed in both exposed canaries.

DNA extraction, PCR and sequencing

Total DNA was extracted from whole blood using the ammonium-acetate protocol (Sambrook et al. 1989). Quantification of the DNA was performed by using a NanoDrop spectrophotometer (Implen NanoPhotometer P330). Four established PCR assays were used for detection of P. matutinum infection from the blood of experimentally exposed canaries. Presence of parasites was confirmed by microscopic examination in the corresponding blood films. Two of these PCR assays used primer sets, which amplify DNA fragments of mitochondrial genome, one of apicoplast genome and one of nuclear genome. The following partial sequences of mitochondrial genome were obtained: cytochrome b (cvt b) gene (primer set HaemNFI/NR3 and HAEMF/R2, Bensch et al. 2000; Hellgren et al. 2004) and cytochrome oxidase subunit I (COI) gene (primer set COIF/R and COIF2/R2, Martinsen et al. 2008). Apicoplast gene (clpc) sequences were obtained using the primer set ClpcF/R and ClpcF2/R2 (Martinsen et al. 2008). Nuclear DNA apical membrane antigen-1 (AMA1) gene fragment sequences were obtained using primer set Pg_AMA1F1/R1 and Pg_AMA1F2/R2 (Lauron et al. 2014). All reactions were performed in 25 μ L total volumes, including 50 ng of a total genomic DNA template $(2 \mu L)$, $12.5 \mu L$ of DreamTaq Master Mix (Thermo Fisher Scientific, Lithuania), $8.5 \,\mu\text{L}$ nuclease-free water and $1 \,\mu\text{L}$ of each primer. One negative control (nuclease-free water) and one positive control (extracted parasite DNA from a blood sample, which was confirmed positive by microscopic examination of blood films) were used to control for false amplifications. Temperature profiles in all PCRs were the same as in the original protocol descriptions. The amplification success was evaluated after running of $2 \mu L$ of PCR product on 2% agarose gels. PCR products $(21 \,\mu\text{L})$ were purified by adding $11 \,\mu\text{L}$ of $8 \,\text{M}$ NH 4Ac, $37 \,\mu\text{L}$ of 96% and 150 μL of 70% ethanol. After centrifugation, air-dried DNA pellets were dissolved in 16 µL of ddH2O. Positive amplifications were sequenced using Big Dye Terminator V3.1 Cycle Sequencing Kit and then loaded on an ABI PRISM[™] 3100 capillary sequencing robot (Applied Biosystems, Foster City, California). The



Fig. 1. Bayesian phylogeny of 24 mitochondrial cytochrome *b* lineages of *Plasmodium* species, seven lineages of *Haemoproteus* spp. and two lineages of *Leucocytozoon* spp. One lineage of *Plasmodium falciparum* is used as outgroup. GenBank accession numbers of lineages are given, followed with codes of the lineages (according to MalAvi database, http://mbio-serv2.mbioekol.lu.se/Malavi) and parasite names. Posterior probabilities >0.69 are indicated near the nodes. The branch lengths are drawn proportionally to the amount of change (scale bar is shown). Bold font indicates *Plasmodium matutinum*.

sequences were edited and aligned using BioEdit software (Hall, 1999). The BLAST algorithm (http://blast.ncbi.nlm.nih.gov/Blast.cgi) was used to compare detected sequences with those deposited in GenBank. All detected sequences were deposited in GenBank (see parasite description).

Phylogenetic analysis

The phylogenetic relationship of the lineage pLINN1 of *P. matutinum* and other avian malaria parasites was inferred using 24 mitochondrial cyt *b* gene sequences of avian *Plasmodium* species, seven of *Haemoproteus* spp. and two of *Leucocytozoon* spp. The sequences were selected from GenBank and

MalAvi database (Bensch *et al.* 2009). *Plasmodium falciparum* was used as outgroup. The GenBank accession numbers of the used sequences and codes of the lineages are given in Fig. 1.

The phylogenetic tree was constructed using Bayesian phylogenetic analysis as implemented in mrBayes version 3.1 (Ronquist and Huelsenbeck, 2003). The General Time Reversible Model including invariable sites and variation among sites (GTR + I + G) was used as suggested by the software mrModeltest 2.2 (http://www.abc.se/~nylander/ mrmodeltest2/MrModelblock). Two independent runs were performed. Each analysis was run for a total of 5 million generations with a sampling frequency of every 100 generations. Before constructing a majority consensus tree, 25% of the initial trees in each run were discarded as 'burn in' periods. The tree was visualized using the software Tree View 1.6.6. (http://evolution.genetics.washington. edu/phylip/software.html).

The sequence divergence between different lineages was calculated according to the Jukes–Cantor model of substitution; the programme MEGA 6.0 was used (Tamura *et al.* 2013). Presence of possible co-infections of malaria parasites in samples from exposed canaries was ruled out due to the absence of doublebase calling in sequence electropherograms (Pèrez-Tris and Bensch, 2005).

Microscopic examination of blood films and voucher material

An Olympus BX61 light microscope equipped with imaging software AnalySIS FIVE were used in morphological analysis (for details see Valkiūnas *et al.* 2016). Intensity of parasitemia was calculated as a percentage by actual counting of the number of parasites per 10 000 red blood cells. The analyses were carried out using the 'Statistica 7' package. Voucher preparations of blood stages of *P. matutinum* (pLINN1) from nightingale as well as the whole blood samples fixed in SET buffer and the cryopreserved infected blood were deposited in Nature Research Centre, Vilnius, Lithuania (see description of the parasite).

Ethical statement

Experimental procedures were approved by the Lithuanian State Food and Veterinary Office (permit 2015-05-07, no. G2-27) and Environmental Protection Agency, Vilnius (permit 2016-05-05, no. 23).

RESULTS

Single malaria infection of P. matutinum was identified in the donor thrush nightingale; other Plasmodium species were not reported neither by microscopic examination of blood films, nor PCRbased tools. However, a Haemoproteus balmorali (Haemosporida, Haemoproteidae) co-infection was present. Both exposed canaries were susceptible and developed single infection of the same malaria parasite, as determined both by microscopic examination of blood films and PCR-based testing. Canaries were resistant to Haemoproteus infection, which was absent from the both exposed birds, providing opportunities to deal with single P. matutinum (pLINN1) infection during parasite morphological and molecular characterization. The lineage pLINN1 of P. matutinum was identified; it clustered with other lineages of malaria parasites and was well distinguishable in the phylogenetic tree (Fig. 1).

Plasmodium (Haemamoeba) matutinum Huff, 1937 (Fig. 2A-L).

DNA sequences: Mitochondrial cyt b lineage pLINN1 (479 bp, GenBank accession no. KY287235). Additional sequences: COI gene (881 bp), clpc (495 bp) and nuclear AMA1 (455 bp), with GenBank accession numbers KY471295, KY471294 and KY471296, respectively.

Avian hosts and distribution: The lineage pLINN1 has been reported in North America, Europe, Asia and New Zealand. It was found in 20 species of birds belonging to 14 families and five orders (Table 1).

Vectors: Remain unidentified. The lineage pLINN1 was found in *Culex pipiens* mosquitoes in Iberian Peninsula and the USA (Table 1). This mosquito is a probable vector of this parasite lineage both in Europe and America.

Site of infection: Red blood cells; no other data.

Representative blood films: Voucher specimens (accession numbers 48772–48775 NS, *L. luscinia*, May 2016, collected by D. Bukauskaitė) were deposited in Nature Research Centre, Vilnius, Lithuania.

Additional material: Representative blood samples from the thrush nightingale L. luscinia were fixed in SET buffer (accessions numbers 9/16R, 101/16v– 103/16v) and also cryopreserved (accessions biobank numbers Cv458/16v–Cv460/16v); this material was deposited in Nature Research Centre, Vilnius.

Prevalence: The overall prevalence was 1 of 11 (9.1%) in thrush nightingales at the study site.

Parasitemia and virulence: All blood stages were reported in the peripheral circulation in the exposed canaries. Prepatent period was about 5 dpe. Light parasitemia developed, with maximum intensity of 0.01% in both exposed birds 10 dpe. The parasitemia remained light or even declined into latency within 1 month postexposure, with a few parasites appearing in the circulation occasionally during entire observation time. The exposed canaries survived to the end of this study; they looked healthy. The clinical signs of disease were not observed during this study.

Morphology of blood stages was the same in the thrush nightingale and the experimentally exposed canaries. The parasite description is based on preparations with parasitemia of 2% in thrush nightingale (Fig. 2).

Trophozoites (Fig. 2A and B) develop in polychromatic and mature erythrocytes. Multiple infection of red blood cells is common (Fig. 2A). Earliest trophozoites are variable in form, often irregular or amoeboid in outline (Fig. 2A); they do not displace or only slightly displace nuclei of infected erythrocytes laterally. Growing and advanced trophozoites possess prominent nuclei, and a vacuole was often seen in the cytoplasm of the largest parasites (Fig. 2B).



Fig. 2. *Plasmodium (Haemamoeba) matutinum* (cytochrome *b* lineage pLINN1) from the blood of thrush nightingale *Luscinia luscinia:* a and b, trophozoites; c–h, erythrocytic meronts; i–k, macrogametocytes; l, microgametocyte. Long arrows, nuclei of parasites; short arrows, vacuoles, arrow heads, pigment granules. Note presence of large circular vacuoles in the majority of growing erythrocytic meronts (c, d, f and g), and their absence in mature meront (h). Giemsa-stained thin blood films. Bar = $10 \,\mu$ M.

Pigment granules are small ($<0.5 \mu$ M), few, darkbrown and usually grouped (Fig. 2B). Largest trophozoites slightly displace nuclei of host cells (Fig. 2B).

Erythrocytic meronts (Fig. 2C-H) develop in polychromatic and mature erythrocytes. Growing meronts possess the plentiful cytoplasm and large nuclei, which size markedly decreases as the parasites mature (compare Fig. 2D with Fig. 2H). Clearly visible circular vacuoles, which may be up to $1.5 \,\mu\text{M}$ in largest diameter are present in many maturing meronts (Fig. 2C, D, F and G), and that is the most readily distinguishable character of this parasite species. Up to five such vacuoles may occur in large maturing meronts. Vacuoles gradually disappear in mature meronts (Fig. 2H). Pigment granules are small (<0.5 μ M), dark-brown or black, usually clumped and are difficult to calculate (Fig. 2E); they do not gather around vacuoles, a characteristic feature of this parasite species. Numerous mature meronts were reported in the morning,

indicating a 24-h merogony cycle. Number of merozoites in mature meronts is 14-28 (on average $18\cdot2\pm$ 4,4); the majority of meronts produce 16–18 merozoites, which are arranged haphazardly (Fig. 2H). Growing and mature meronts markedly displace nuclei of infected erythrocytes (Fig. 2C–H) and sometimes even enucleate the host cells.

Macrogametocytes (Fig. 2I–K) develop mainly in mature erythrocytes, but occasionally were also seen in polychromatic erythrocytes. Growing and mature gametocytes are of roundish form, possess prominent nuclei; small vacuoles were seen occasionally. Pigment granules are small ($<0.5 \mu$ M), black or dark-brown, roundish, scattered in the cytoplasm or sometimes grouped (Fig. 2I–K). Gametocytes markedly deform infected red blood cells and displace their nuclei toward one of poles of the host cells; they can enucleate the infected cells. Fully-grown gametocytes are relatively small; they do not exceed 10 μ M in their largest diameter, a characteristic feature of this *Haemamoeba* species.

Urder and family	sanado	LUCATION	
Diptera Culicidae Antervaiformes	Culex pipiens	Italy, USA	Kimura et al. (2010); Martínez-de la Puente et al. (2015)
Apterygidae	Apteryx haastii Apteryx mantelli	New Zealand	Howe et al. (2012); Sijbranda et al. (2016)
Gruiformes	•		
Rallidae	Fulica cristata	Spain	Ferraguti et al. (2013)
Passeriformes			
Callaeidae	Philesturms carunculatus	New Zealand	Castro $et al. (2011)$
Carduelidae	Carduelis cannabina	Sweden	Hellgren et al. (2007) ; Nilsson et al. (2016)
Corvidae	Corvus cornix cornix	Italy	GenBank (unpublished)
Meliphagidae	An thorn is melanura	New Zealand	Ewen <i>et al.</i> (2012)
Muscicapidae	Erithacus rubecula	North-western Iberia (NWI), WGC,	Drovetski et al. (2014); Mata et al. (2015); Santiago-Alarcon et al.
		Germany	(2016)
	Luscinia luscinia ^a	Lithuania	This study
	Luscinia svecica	Czech Republic	Svoboda et al. (2015)
Paridae	Baeolophus bicolor	USA	Fast <i>et al.</i> (2016)
	Cyanistes caeruleus	UK, WGC	Wood et al. (2007); Cosgrove et al. (2008); Szöllősi et al. (2011);
		-	Drovetski et al. (2014); Mata et al. (2015)
Passeridae	Passer montanus	Italy ^b	Martínez-de la Puente <i>et al.</i> (2015)
Petroicidae	Petroica longipes	New Zealand	Sijbranda et al. (2016)
Turdidae	Turdus merula	Portugal, New Zealand, NWI,	Hellgren et al. (2011); Howe et al. (2012); Ewen et al. (2012);
		WGC, Italy ^b , Germany, Russia ^c	Drovetski et al. (2014); Mata et al. (2015); Martínez-de la Puente
			et al. (2015) ; Santiago-Alarcon et al. (2016) ; Sijbranda et al. (2016)
	I uraus philometos	New Zealand, IN W1, South Caucasus	Ewen <i>et al.</i> (2012); Drovetski <i>et al.</i> (2014); Mata <i>et al.</i> (2015); Siibranda <i>et al.</i> (2016)
Zosteropidae	Zosterops lateralis	New Zealand	Sijbranda et al. (2016)
Sphenisciformes			
Spheniscidae	Spheniscus humboldti Eudv⊅tula minor	Japan, New Zealand	Sijbranda (2015); GenBank (unpublished)
Strigiformes			
Strigidae	$A thene \ noctua$	$Italy^{b}$	Martínez-de la Puente $et al. (2015)$

Table 1. Hosts and locations where the pLINN1 lineage of *Plasmodium matutinum* has been reported

investive stages develop in these hosts. ^b Reports from mosquito blood meals. Bird species were identified using PCR-based diagnostic, which identified DNA of corresponding avian hosts in tested mosquitoes. ^c Report from the Biological Station Rybachy, Kaliningrad District, Russia (unpublished data).

Microgametocytes (Fig. 2L). General configuration and other features are as for macrogametocytes, with usual haemosporidian sexual dimorphic characters, which are the pale staining of the cytoplasm and the diffuse large nuclei.

Morphometric features of blood stages of *P. matutinum* (pLINN1) were similar to those described formerly in this species (Valkiūnas, 2005).

Remarks

Plasmodium matutinum can be readily distinguished from the majority of species of subgenus Haemamoeba due to presence of large circular vacuoles in its growing erythrocytic meronts (Fig. 2G). Large vacuoles were also described in the erythrocytic meronts of *Plasmodium griffithsi* (Garnham, 1966), Plasmodium tejerai (Gabaldon and Ulloa, 1977) and Plasmodium giovannolai (Corradetti et al. 1963). In P. griffithsi, erythrocytic meronts contain residual bodies, and elongate gametocytes develop. Both these characters are not features of P. matutinum (Fig. 2). In *P. tejerai*, the vacuoles often exceed 2 μ M in diameter, and pigment granules gather around the vacuoles; that are not characters of P. matutinum. Plasmodium giovannolai is most similar to P. matutinum (Valkiūnas, 2005). The latter parasite can be distinguished due to the smaller size of pigment granules in its gametocytes and the presence of vacuoles in growing trophozoites (Fig. 2B). Molecular characterization of *P. giovannolai* is absent.

Phylogenetic relationships of parasites

Partial DNA sequences of cyt *b* gene are available for two species of subgenus *Haemamoeba* with readily distinguishable vacuoles in erythrocytic meronts, i.e. *P. matutinum* and *P. tejerai*. Genetic difference in cyt *b* gene between them is large (4·1%), indicating that these species are valid. Both parasites clustered together in our phylogenetic analysis (Fig. 1), suggesting that the feature of vacuolization of erythrocytic meronts may be informative phylogenetically. Investigation of other *Plasmodium* species containing large vacuoles in erythrocytic meronts is needed to test this observation.

DISCUSSION

The key result of this study is the development of molecular markers to detect, identify and examine phylogenetic relationships of P. matutinum. Partial sequences of mitochondrial, apicoplast and nuclear genomes were identified. Currently, the cyt b gene sequences are particularly useful because they have been identified in many avian haemosporidian species and can be used in comparative parasitology research (Perkins, 2014; Bensch *et al.* 2016; Mantilla *et al.* 2016; Ricklefs *et al.* 2017). The obtained COI,

clpc and AMA1 gene sequences extend information about molecular characterization of P. matutinum, providing opportunities to use these data in multigene analysis of phylogenetic status of malaria parasites (Martinsen et al. 2008). The available information about nuclear genome genes of avian malaria parasites remain scarce (Lauron et al. 2014), but is needed in research aiming studies of mechanisms of speciation and diversity maintenance in Plasmodium spp. and related haemosporidian parasites (Valkiūnas et al. 2013a, b). Additionally, information about clpc gene sequences is helpful in taxonomic research. For example, the genetic distance in cyt b gene between closely related lineages of Haemoproteus jenniae and Haemoproteus iwa is only 0.6% or 1 bp, but 18 nucleotide differences or 4% sequence divergence between these parasites was recorded in a segment of clpc gene, indicating existence of different parasite species and the validity of both species names (Levin et al. 2012).

The former microscopic studies showed that P. matutinum is present in the USA and Europe and probably is cosmopolitan in countries with warm climates (Garnham, 1966; Bennett et al. 1982; Valkiūnas, 2005). It was particularly often reported in thrushes (Turdidae) and columbiform birds by microscopic examination of blood samples. The recent molecular data support these conclusions and also add species of Muscicapidae to the list of relatively common hosts (Table 1). Available data indicate that birds of the Turdidae, Muscicapidae and Columbidae are often infected. Additionally, the endemic Great spotted kiwi (Apteryx haastii), Brown kiwi (Apteryx mantelli) and several other rare bird species (Table 1) are affected by this infection in New Zealand (Castro et al. 2011; Howe et al. 2012; Sijbranda, 2015; Sijbranda et al. 2016). It is noteworthy that reports of *P. matutinum* (pLINN1) in non-migrating birds and/or mosquitoes in USA, Italy, Spain, Japan and New Zealand certainly indicate the local transmissions and the global distribution. However, the range of competent avian host, in which this parasite completes life cycle and produces invasive stages (gametocytes), remains unclear because the presence of the gametocytes in the circulation of PCR-positive birds was reported only during this study (Table 1). It is difficult to rule out that some positive PCR-based diagnostic results might be due to the amplification of DNA from sporozoites, which might be unable to initiate development in birds, or due to templates, which may come from abortive development of tissue stages in non-competent avian hosts (Valkiūnas et al. 2014; Moens et al. 2016; Palinauskas et al. 2016; Valkiūnas and Iezhova, 2017). Further studies combining PCR-based and microscopic approaches are needed to determine the certain range of competent avian hosts of this malaria infection.

The close phylogenetic relationships between P. matutinum and P. tejerai (Fig. 1) likely indicate American origin of these infections because both the parasites are present in American birds, but P. tejerai is absent from European birds. In other words, P. matutinum might be of secondary origin in Europe. Interestingly, both these parasites are quite different in cyt b gene from other common species of subgenus Haemamoeba: Plasmodium relictum, Plasmodium cathemerium, Plasmodium gallina*ceum*, which appeared in the different clade (Fig. 1). For example, the genetic difference of *P. matutinum* (pLINN1) and P. tejerai (pSPMAG01) from the widespread lineages pSGS1, pGRW4, pGRW11 of P. relictum ranges between 5.7 and 6.1% and between 5.9 and 6.8%, respectively. That suggests possible independent evolution of P. matutinum (pLINN1) and P. tejerai (pSPMAG01) in regard to *P. relictum* and probably other *Haemamoeba* parasites.

It is interesting to mention that P. matutinum and P. tejerai possess large circular vacuoles in their growing erythrocytic meronts (Fig. 2G). That is not a case in P. relictum, P. cathemerium and P. gallinaceum (Garnham, 1966; Valkiūnas, 2005). Because P. matutinum and P. tejerai are closely phylogenetically related (Fig. 1), it might be that such vacuolization is a morphological character, which indicates evolutionary relationships between malaria parasites of subgenus Haemamoeba. Large vacuoles are also present in erythrocytic meronts of P. giovannolai and P. griffithsi. Molecular characterization of the latter two parasites is needed to test this assumption. It is worth mentioning that similar large vacuoles have been described in blood stages of several Haemoproteus parasites, which is a sister group to Plasmodium (Valkiūnas, 2005; Bensch et al. 2016), but the origin and metabolic function of such vacuoles in blood stages of haemosporidian parasites remains unclear. It is noteworthy that sporogonic stages of avian haemosporidian parasites, particularly often zygotes, ookinetes and early oocysts, contain cytoplasmic inclusions, which are the gatherings of amorphous dense material called a crystalloid material (Valkiūnas, 2005; Atkinson et al. 2008). The crystalloid material is washed out during fixation with alcohols and looks like an empty space, which is usually described as a 'vacuole' in stained preparations under a light microscope. The crystalloid likely performs energy functions and takes part in the metabolism of lipids (Desser and Bennett, 1993; Valkiūnas, 2005; Atkinson et al. 2008). Vacuoles in the cytoplasm of P. matutinum and P. tejerai are similar in appearance to the vacuoles reported in sporogonic stages of haemosporidians, and may be homologous to them. Histochemical and electron microscopy studies of growing erythrocytic meronts of P. matutinum and related parasites are needed to answer this question.

Former experimental studies showed that virulence of different P. matutinum strains varies markedly. Italian strain caused lethal malaria in avian hosts, but American strain was less virulent, with no mortality reported in experimentally exposed birds (Corradetti et al. 1960; Garnham, 1966; Valkiūnas, 2005). Field observations indicate that this infection is associated with diseases and even mortality in native and endemic birds in New Zealand (Castro et al. 2011; Howe et al. 2012; Sijbranda et al. 2016). This study shows that the lineage pLINN1 of P. matutinum was relatively benign in canaries during first 4 months after inoculation of infected blood. However, the sample size was limited only to two individual birds, and this observation should be treated with a caution. It is important to note that the genetically similar parasite has been recently found in dead blackbirds Turdus merula in Austria (Dinhopl et al. 2015), suggesting that some virulent lineages of P. matutinum might be present in Europe, as has been formerly reported by Corradetti et al. (1960) in Italy. Further studies of genetic diversity of P. matutinum are needed for better understanding the virulence of this infection and its role in natural bird populations.

Former studies showed that P. matutinum completed sporogony in Culex pipiens, Culex tarsalis and Culex stigmatosoma mosquitoes, but susceptibility of different isolates was markedly different in same mosquito species. For example, the American strain failed to develop or developed only in few individual C. pipiens mosquitoes, while the Italian strain readily developed and completed sporogony in this mosquito species (Corradetti et al. 1962; Garnham, 1966). Vectors of the lineage pLINN1 of P. matutinum remain unknown. PCR-based reports of this parasite in C. pipiens in the USA and Italy (Table 1) are in accordance with former experimental parasitology studies, suggesting possible involvement of this ornithophilic mosquito in transmission of P. matutinum (pLINN1). Detection of infective sporozoites in mosquitoes is needed to prove this observation.

ACKNOWLEDGEMENTS

This paper benefited from comments of two anonymous reviewers. We thank V. Jusys and V. Eigirdas (Ventės Ragas Ornithological Station, Lithuania) for support during field work.

FINANCIAL SUPPORT

This study was funded by the Research Council of Lithuania (no. MIP-045/2015).

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