

Haemosporidian infections in skylarks (*Alauda arvensis*): a comparative PCR-based and microscopy study on the parasite diversity and prevalence in southern Italy and the Netherlands

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Abstract Changes in agricultural management have been identified as the most probable cause for the decline of Skylark (*Alauda arvensis*) populations in Europe. However, parasitic infections have not been considered as a possible factor influencing this process. Four hundred and thirty-four Skylarks from the Southern Italy and the Netherlands were screened for haemosporidian parasites (Haemosporida) using the microscopy and polymerase chain reaction (PCR)-based methods. The overall prevalence of infection was 19.5%; it

was 41.8% in Italian birds and 8.3% in Dutch birds. The prevalence of *Plasmodium* spp. was 34.1% and 6.5% in Skylarks from Italy and Netherlands, respectively. Approximately 15% of all recorded haemosporidian infections were simultaneous infections both in Italian and Dutch populations. Six different mitochondrial cytochrome *b* (cyt *b*) lineages of *Plasmodium* spp. and three lineages of *Haemoproteus tartakovskyi* were found. The lineage SGS1 of *Plasmodium relictum* was the most prevalent at both study sites; it was recorded in 24.7% of birds in Italy and 5.5% in the Netherlands. The lineages SYAT05 of *Plasmodium vaughani* and GRW11 of *P. relictum* were also identified with a prevalence of <2% at both study sites. Two *Plasmodium* spp. lineages (SW2 and DELURB4) and three *H. tartakovskyi* lineages have been found only in Skylarks from Italy. Mitochondrial cyt *b* lineages SYAT05 are suggested for molecular identification of *P. vaughani*, a cosmopolitan malaria parasite of birds. This study reports the greatest overall prevalence of malaria infection in Skylarks during the last 100 years and shows that both *Plasmodium* and *Haemoproteus* spp. haemosporidian infections are expanding in Skylarks so it might contribute to a decrease of these bird populations in Europe.

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Introduction

Pigment-forming haemosporidians (Haemosporida) are a clearly phylogenetically defined group of obligate hetero-

geneous parasites, which inhabit in birds all over the world except the Antarctic (Greiner et al. 1975; McClure et al. 1978; Atkinson and van Riper 1991; Bishop and Bennett 1992). Over 45% of bird species of the world fauna have been currently investigated with respect to infection with haemosporidians (Valkiūnas 2005). *Haemoproteus* spp. was recorded in approximately 50% and *Plasmodium* and *Leucocytozoon* spp. in approximately 30% of the investigated bird species (Valkiūnas 2005). Although these parasites are widespread geographically, their prevalence varies markedly between different regions, host species and populations (Greiner et al. 1975; Valkiūnas 2005).

The Skylark (*Alauda arvensis*) is largely distributed in the temperate zone of Europe, Asia and Northern Africa, with 13 subspecies described (Cramp 1988; Donald 2004). In Western Europe, the Skylark is undergoing a rapid population decline in recent years, as are many other farmland birds (BirdLife International 2004; Donald et al. 2006). The decline of farmland birds is frequently associated with changed conditions during the breeding season and deterioration of the breeding habitat (Donald 2004; Newton 2004). Although the impact of parasites on bird populations is often overlooked in wildlife ornithology, parasitism certainly is an important factor in conservation biology and should therefore be considered in biodiversity preservation studies (Valkiūnas 2005; Parker et al. 2006). The establishment of parasites in new hosts and geographic areas is often associated with changes in virulence and might lead to devastating outbreaks among resident bird populations, which is particularly well documented in relatively simple island ecosystems in Hawaii islands (Atkinson and van Riper 1991) and recently recorded in Galápagos (Levin et al. 2009). A possible role of blood parasites as a factor influencing the decline of Skylark populations has not been considered because of limited knowledge about haemosporidian infections in this bird species (Bennett et al. 1982; Valkiūnas 2005). The aim of this study was to describe distribution, diversity, prevalence and intensity of haemosporidian parasites in Skylarks from two sites in Europe.

Materials and methods

Study sites and collection of blood samples

One hundred and forty-six Skylarks were caught in the Volturino Plain (41°02' N, 13°55' E) located 40 km north of Naples, Italy between 1 and 30 of October in 2006 and 2007. Two hundred and eighty-eight Skylarks were sampled in the northern part of the Netherlands (52°55' N, 006°18' E) between May 2006 and December 2007: 12 of them were caught during the period of establishment of

breeding territory, 145 adult birds—during the breeding season, 9—during moult, 45—during autumn migration and 2—during winter. Additionally, 71 nestlings have been sampled in nests; they were 5–7 days old. Skylarks were caught using mist nets or traps. The birds were banded to avoid resampling. Blood samples were taken by puncturing the brachial vein. Blood films were air-dried, fixed in absolute methanol in the field and stained with Giemsa solution in the laboratory as described by Valkiūnas et al. (2008b).

A complementary blood sample (20–50 µl) was collected using heparinized microcapillaries and stored in non-lysis SET buffer (Waldenström et al. 2004) or in 96% ethanol (only Dutch samples). In the field, the samples were stored at ambient temperature (Italy) or on ice (the Netherlands) and later stored at –20°C in the laboratory. The blood samples were analysed by molecular methods between 1 and 24 months after their collection. In total, 434 samples were collected at both study sites (Table 1).

Examination of blood films and parasite morphology

An Olympus BX51 light microscope equipped with an Olympus DP12 digital camera and imaging software DP-SOFT was used to examine blood slides, prepare illustrations and to take measurements. Blood films were examined for 10–15 min at low magnification (×400), and then

Table 1 Prevalence of haemosporidian infections in Skylark *Alauda arvensis* at two study sites in Europe, 2006–2007

Parasite species and mitochondrial cytochrome <i>b</i> lineage	Prevalence	
	Italy (N=146)	The Netherlands (N=288)
<i>Haemoproteus tartakovskiyi</i>		
hALARV1	1 (0.7) ^a	0
hALARV2	1 (0.7)	0
hALARV3	2 (1.4)	0
Total	4 (2.7)	0
<i>Plasmodium (Haemamoeba) relictum</i>		
pSGS1	36 (24.7)	16 (5.5)
pDELURB4	1 (0.68)	0
pGRW11	4 (2.7)	0
pMOTALB1	0	1 (0.3)
Total	41 (28.0)	17 (5.9)
<i>Plasmodium (Novyella) vaughani</i>		
pSYAT5	3 (2.0)	2 (0.6)
<i>Plasmodium (Novyella) sp.</i>		
pSW2	6 (4.1)	0
Unidentified simultaneous infections	8 (5.5)	5 (1.7)
Grant total	62 (42.5)	24 (8.3)

^a Percentage is given in parentheses

at least 100 fields were studied at high magnification ($\times 1,000$), as described by Valkiūnas et al. (2008b). We used the morphometric features (Table 2) and identified parasites according to Valkiūnas (2005). The intensity of infection was estimated as a percentage by actual counting of the number of parasites per 1,000 red blood cells or per 10,000 red blood cells if infections were light (i.e. $<0.1\%$), as recommended by Godfrey et al. (1987).

The morphology of gametocytes of *Haemoproteus tartakovskiyi* from Skylarks was compared with the type and voucher specimens of *H. tartakovskiyi* from its type host the Common crossbill (*Loxia curvirostra*) (accession no. 413.91) at the Collection of the Institute of Ecology, Nature Research Centre, Vilnius. The morphology of *Plasmodium (Novyella) vaughani* from the Skylark was compared with the type and voucher material of the same parasite from its type vertebrate host the American robin (*Turdus migratorius*) and additional vertebrate host the Blackbird (*Turdus merula*) (accession nos. 635, 639, 654 and 655) in the Garnham Collection at the Natural History Museum, London.

Extraction of DNA, PCR, sequencing and analysis of molecular data

The DNA was extracted using a standard ammonium acetate method (Sambrook et al. 2002). Diluted total DNA was used as the template in PCR assays for detection of the parasites, using primers and temperature profiles as in Hellgren et al. (2004). The method consists of a nested PCR assay that amplifies a part of the parasites mitochondrial cytochrome *b* (*cyt b*) gene in two steps, first an initial PCR (primers HaemNFI/HaemNR3; 570 bp excluding primers) that amplifies parasites from all of the three genera followed by a second step that separates *Leucocytozoon* spp. (primer HaemFL/HaemR2L; 478 bp excluding primers) from parasites of the genera *Plasmodium* and *Haemoproteus* spp. (primers HaemF/HaemR2; 480 bp excluding primers). By amplifying the parasite DNA in two PCRs, the sensitivity of the screening is increased (Waldenström et al. 2004; Hellgren et al. 2004). Positive or negative infections were seen as presence or absence of bands on a 2% agarose gel using 1.5 μ l of the final PCR

Table 2 Morphometric parameters of mature blood stages of *Haemoproteus tartakovskiyi* (lineage hALARV1) and *Plasmodium (Novyella) vaughani* (lineage pSYAT05)

Feature	Measurements (μm) ^a	
	<i>Haemoproteus tartakovskiyi</i>	<i>Plasmodium vaughani</i>
Meront		
Length		3.7–7.5 (5.4 \pm 0.8)
Width		1.6–3.3 (2.3 \pm 0.5)
Area		6.2–12.7 (9.7 \pm 1.5)
Area of globule		0.5–1.2 (0.8 \pm 0.2)
No. of pigment granules		1–2 (1.2 \pm 0.4)
No. of merozoites		4–8 (5.6 \pm 1.3)
Macrogametocyte		
Length	10.6–12.3 (11.6 \pm 0.5)	9.6–12.2 (11.3 \pm 0.8)
Width	3.1–4.8 (4.1 \pm 0.4)	2.5–3.3 (2.8 \pm 0.2)
Area	39.3–53.9 (49.4 \pm 4.1)	23.2–39.2 (28.6 \pm 4.5)
Gametocyte nucleus		
Length	2.1–2.9 (2.5 \pm 0.2)	2.2–4.2 (2.9 \pm 0.5)
Width	1.4–2.6 (1.9 \pm 0.4)	1.3–2.8 (2.1 \pm 0.4)
Area	2.3–5.6 (3.7 \pm 0.8)	2.9–6.3 (4.2 \pm 1.1)
Number of pigment granules	13–22 (16.1 \pm 2.4)	3–8 (5.3 \pm 1.3)
Microgametocyte		
Length	10.7–12.7 (11.5 \pm 0.5)	10.5–12.8 (12.1 \pm 0.7)
Width	3.9–5.3 (4.4 \pm 0.4)	2.8–3.7 (3.2 \pm 0.3)
Area	40–57.9 (50.6 \pm 3.8)	29.8–43.6 (35.4 \pm 3.9)
Gametocyte nucleus		
Length	6.1–9.2 (7.6 \pm 1)	5.8–7.9 (6.8 \pm 0.8)
Width	3–4.6 (4 \pm 0.4)	2.5–3.2 (2.9 \pm 0.3)
Area	18.7–35.3 (27.7 \pm 4.1)	11.7–22 (16.8 \pm 2.8)
Number of pigment granules	9–20 (14.8 \pm 2.6)	4–9 (5.8 \pm 1.2)

^aAll measurements ($n=31$) are given in micrometers. Minimum and maximum values are provided, followed in parentheses by the arithmetic mean and standard deviation

product. Samples which showed positive amplification were sequenced using the procedures described by Bensch et al. (2000). Amplified fragments were sequenced from the 5' end with the primer HaemF. We used dye terminator cycling sequencing (big dye) kit and the samples were loaded on an ABI PRISM™ 3100 sequencing robot (Applied Biosystems, Florida, USA).

The obtained sequences were edited and aligned using the BioEdit programme (Hall 1999). The appearance of double peaks in the sequence was considered as mix infection. All unique lineages, i.e. sequences differing by one or more nucleotide base pair, were sequenced in the reversed direction with the complement primer HaemR2. The 38 taxa (total 474 nucleotides) in the final alignment were used for Bayesian analysis. Bayesian phylogeny was constructed using mrBayes version 3.1.2 (Ronquist and Huelsenbeck 2003). We used the General Time Reversible model including invariable sites and variation among sites (GTR+I+G) as suggested by the software mrModeltest 2.2 (Nylander 2004, software available from <<http://www.ebc.uu.se/systzoo/staff/nylander.html>>). Two simultaneous runs were conducted with a sample frequency of every 100th generation over three million generations. Convergence in phylogeny estimation for each analysis was assessed using the programme Tracer (Rambaut, A. & Drummond, A., available at <http://evolve.zoo.ox.ac.uk>) and used to indicate the appropriate “burn-in” period. The 25% of the trees were discarded as burn-in period. The remaining trees were used to construct a majority rule consensus tree. The phylogenies were visualised using Tree View 1.6.6. (software available from <<http://evolution.genetics.washington.edu/phyliip/software.html>>). We used sequences of haemosporidian parasites, which species were positively identified (for linkage of parasite lineages with their morphospecies, see Križanauskienė et al. 2006; Palinauskas et al. 2007; Krone et al. 2008; Martinsen et al. 2008; Valkiūnas et al. 2008a, b; Zehindjiev et al. 2008; Bensch et al. 2009; Križanauskienė et al. 2010). GenBank accession numbers and MalAvi reference names (see Bensch et al. 2009) of all lineages mentioned in this article are given in Fig. 1

The sequence divergence between the different lineages was calculated with the use of a Jukes–Cantor model of substitution, with all substitution weighted equally, implemented in the programme MEGA version 4 (Tamura et al. 2007) where the pairwise deletion was selected.

Student's *t* test for independent samples was used to determine statistical significance between mean linear parameters. Prevalences were compared by Yates corrected Chi-square test. A *P* value of 0.05 or less was considered significant.

The representative blood slides were deposited in the Nature Research Centre, Vilnius, Lithuania (accession nos. 47733–47736 NS). Sequences of new parasite lineages

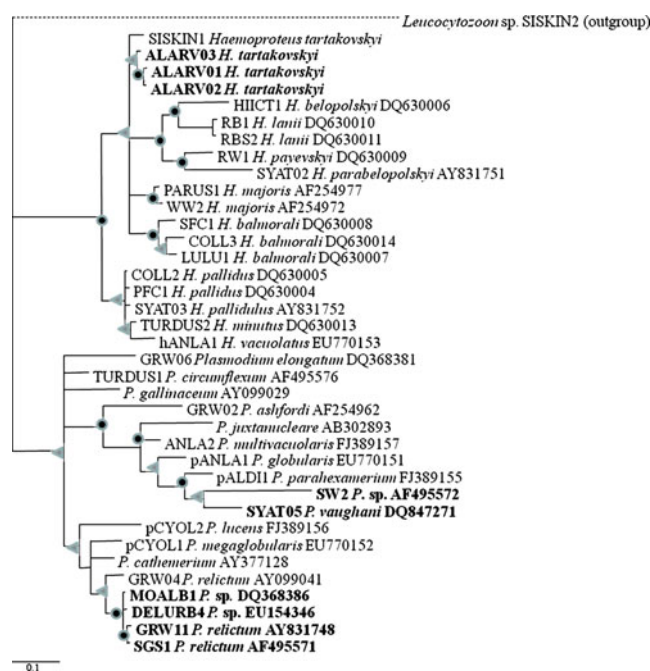


Fig. 1 Bayesian phylogeny of cytochrome *b* gene lineages of positively identified species of avian pigment-forming haemosporidian parasites. Lineages recorded in the Skylark *A. arvensis* are given in bold. Names of the lineages (when available) are given before the species names of parasites; GenBank accession numbers of the lineages are provided after the parasite species names. Nodal support values (circles—80–100%; triangles—60–70%) indicate posterior clade probabilities

were deposited in GenBank (nos. GU289671, GU289672 and GU289673).

Results

Molecular analysis of blood samples

Only pigment-forming haemosporidian parasites of the genera *Haemoproteus* and *Plasmodium* were found using molecular techniques; that is in accord to microscopy data (see below) (Table 1). The overall prevalence of haemosporidian infection was 19.5%; it was 41.8% in Italian and 8.3% in Dutch Skylarks ($P < 0.001$). We detected three lineages of *Haemoproteus* spp. and six lineages of *Plasmodium* spp. in 85 infected Skylarks (Fig. 1). *Plasmodium* parasites were found in 34.1% of Italian and 6.5% of Dutch Skylarks ($P < 0.001$). *Haemoproteus* were present in 2% of the Italian birds and absent from the Dutch Skylarks. In Dutch Skylarks, the prevalence of *Plasmodium* spp. was 4.6% for adults and 9.9% in nestlings ($P < 0.05$, not significant), indicating the active malaria transmission at the study site. According to the PCR diagnostics, 5.5% and 1.7% simultaneous infections have been detected in Italian and Dutch populations, respec-

tively. All recorded malarial infections were simultaneous in nestlings.

At both study sites, the lineage SGS1 of *Plasmodium relictum* was most prevalent; it was recorded approximately in 24.7% of birds in Italy and 5.5% in the Netherlands (Table 1). This lineage together with the lineages pGRW11, pMOALB1, pDELURB4 and pGRW4 form well-supported clade of *P. relictum* morphospecies (Fig. 1). Lineages pGRW11, pDELURB4 and pSW2 has been recorded only in Italian birds and lineage pMOALB1 was recorded only in Dutch Skylarks; these lineages were rare (prevalence < 5%). *Plasmodium* lineage pSYAT05 was found in both studied populations (Table 1).

Three *Haemoproteus* lineages (hALARV1, hALARV2 and hALARV3) were found only in Skylarks from Italy. These lineages cluster together with lineage hSISKIN1 of *H. tartakovskiyi* (the *p* distances between these lineages ranged from 0.2% to 2.3% with a total mean distance of 1.25%) and form well-supported clade with the latter parasite (Fig. 1).

Microscopic investigation

All samples were examined microscopically, and the PCR-based diagnostics (both positive and negative results) was confirmed by microscopic observations. Microscopic examination revealed undetected by PCR simultaneous haemsporidian infections, which were present approximately in 15% of infected birds in both Italian and Dutch populations. Over 60% of all recorded infections were light (<0.001%), so it could be regarded as chronic. For some of the detected *cyt b* lineages (pSW2, pMOTALB1 and pDELURB4), we were unable to identify species due to low intensity of parasitemia and absence of all necessary blood stages on the slides.

H. tartakovskiyi (Figs. 1 and 2e–h) (lineages hALARV1, hALARV2 and hALARV3), *P. (Novyella) vaughani* (Figs. 1 and 2i–l) (lineage pSYAT05) and *P. relictum* (lineage pSGS1) were identified using morphological features of blood stages of the parasites. Lineages pMOALB1 and pDELURB4 of *Plasmodium* spp. are closely related to the lineages pSGS1, pGRW11 and pGRW4 of *P. (Haemamoeba) relictum* (Palinauskas et al. 2007; Knowles et al. 2010) with genetic difference among them between 0.2% and 2.3% (Fig. 1); these lineages probably belong to this morphospecies. However, our material is incomplete for these parasites' unequivocal identification using morphological characters because the recorded infections were too light. The unidentified *Plasmodium* lineage (pSW2) is genetically distant from other *Plasmodium* spp. lineages (from 6.6% to 12.3%); based on available morphological features, it belongs to subgenus *Novyella*. Additional material is needed for identification of this parasite species.

The parasites of the lineages hALARV1, hALARV2 and hALARV3 (Figs. 1 and 2e–h) are indistinguishable morphologically among each other and from *H. tartakovskiyi*; they are also genetically similar to the lineage pSISKIN1 of *H. tartakovskiyi* (Figs. 1 and 2a–d) from its type host, the Common crossbill (*L. curvirostra*) and additional host the Siskin (*Spinus spinus*). Genetic divergence between these lineages varies between 0.2% and 2.1%. We consider all these lineages as intraspecies genetic variation of the same morphospecies, i. e. *H. tartakovskiyi*.

The intensity of parasitemia was light in all infected birds; it was <0.05% in the great majority of our samples. The highest intensity of malaria infection (0.77%) was recorded in one bird infected with the *P. relictum* lineage SGS1 in Italy on 27 October 2006. All *Haemoproteus* infections were of low intensity (between 0.015% and 0.018%). Because (1) *H. tartakovskiyi* has been recorded in Skylarks for the first time and (2) lineages for molecular identification of *P. vaughani*, a widespread agent of avian malaria have not been determined, we describe recorded in this bird parasites briefly and link these morphospecies with their *cyt b* lineages.

Description of parasites

P. (Novyella) vaughani Novy and MacNeal, 1904 (Fig. 2i–l, Table 2)

Avian host: Skylark (*A. arvensis*) (Passeriformes, Alaudidae)

DNA sequences: Mitochondrial *cyt b* gene lineage pSYAT05 (479 bp), GenBank accession no. DQ847271

Prevalence: five of 434 (1.2%) (Table 1)

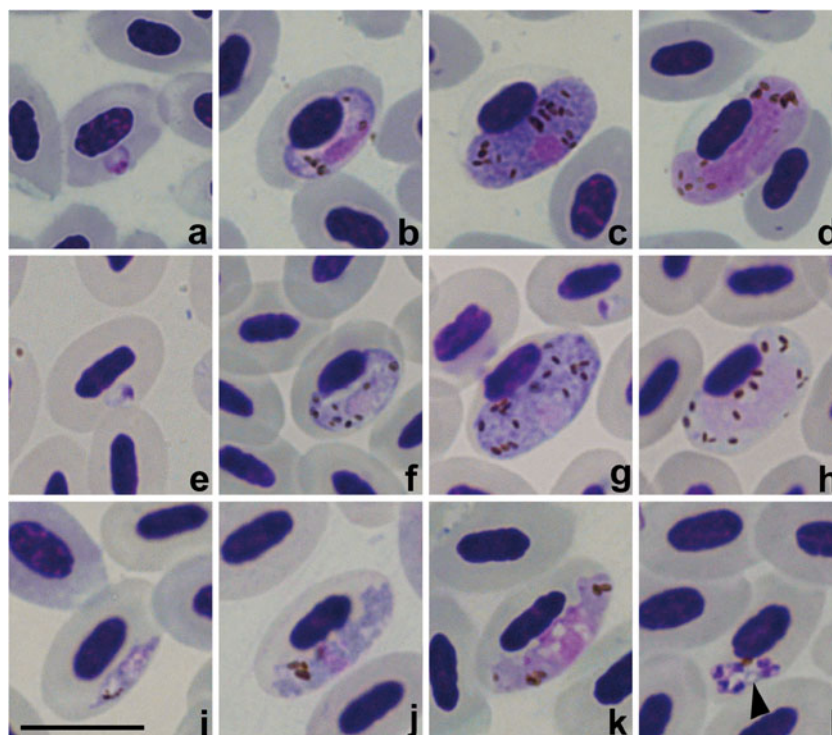
Additional hosts: The lineage pSYAT05 of *P. vaughani* has been recorded in seven bird species: American robin (*T. migratorius*), Blackbird (*T. merula*), Blackcap (*Sylvia atricapilla*), Sardinian warbler (*Sylvia melanocephala*) and Tomtit (*Petroica macrocephala*) (Bensch et al. 2009; Bentz et al. 2007; Hellgren et al. 2007; Martinsen et al. 2008).

Geographical distribution: the lineage pSYAT05 was recorded in the USA, Europe and New Zealand (Bensch et al. 2009; Bentz et al. 2007; Hellgren et al. 2007; Martinsen et al. 2008), so it seems to be cosmopolitan in distribution.

Representative blood films: Voucher specimen (accession number 47736 NS *A. arvensis*, 4 October 2007, collected by P. Zehndjiev) is deposited in the Nature Research Centre, Vilnius, Lithuania. Simultaneous infection of *P. relictum* is present in the voucher slide 47736 NS.

Erythrocytic meronts (Fig. 2l): Develop in mature erythrocytes; they were seen anywhere in the host cells. Fully grown meronts are variable in form, most frequently are roundish, oval or irregular; mature meronts contain between four and eight merozoites (Fig. 2l, Table 2); one clearly defined round refractive colourless globule is

Fig. 2 *H. tartakovskyi* from the blood of its type vertebrate host, the crossbill *L. curvirostra* (a–d, lineage hSISKIN1) and the Skylark *A. arvensis* (e–h, lineage hALARV03), and *P. (Novyella) vaughani* (i–l, lineage pSYAT05) from the Skylark: a–b, i, e young gametocytes; c, g, j—macrogametocytes; d, h, k—microgametocytes; l—erythrocytic meront. Arrow—a refractive colourless globule. Giemsa-stained thin blood films. Bar=10 μ m



present in each meront (Fig. 2l, Table 2); it is frequently located close to a clump of pigment granules. The influence of meronts on infected erythrocytes is not pronounced.

Macrogametocytes (Fig. 2j) develop in mature erythrocytes and are elongated in form. The cytoplasm is homogeneous in appearance, sometimes contains large vacuoles. Parasite nucleus is prominent (Table 2), variable in shape and usually median in position (Fig. 2j). Pigment granules are few (Table 2), elongated or sometimes roundish, of medium size (0.5–1.0 μ m), randomly scattered throughout the cytoplasm or clumped in small groups. The influence of gametocytes on infected erythrocytes is not pronounced.

Microgametocytes (Fig. 2i, k): The general configuration is as for macrogametocytes with the usual haemosporidian sexual dimorphic characters. The parasite nucleus is diffuse, and its size is markedly variable in different gametocytes (Fig. 2i, k).

H. tartakovskyi Valkiūnas 1986 (Fig. 2e–h, Table 2)

Avian host: Skylark (*A. arvensis*) (Passeriformes, Alaudidae)

DNA sequences: Mitochondrial *cyt b* gene lineages hALARV1 (479 bp, GenBank accession no. GU289671), hALARV2 (479 bp, GenBank accession no. GU289672) and hALARV3 (479 bp, GenBank accession no. GU289673).

Prevalence: four of 434 (1.0%) (Table 1)

Additional hosts: The lineages hALARV1, hALARV2 and hALARV3 have been recorded only in Skylarks so far.

Closely related lineage hSISKIN1 is prevalent in crossbills and siskins in Europe (Fig. 1).

Geographical distribution: The lineages hALARV1, hALARV2 and hALARV3 have been recorded only in Italy.

Representative blood films: Voucher specimens (accession numbers 47733 NS, 47734NS and 47735 NS, *A. arvensis*, 23–26 October 2006, collected by P. Zehtindjiev) are deposited in the Institute of Ecology, Nature Research Centre, Vilnius, Lithuania.

Young gametocytes (Fig. 2e, f): The earliest forms (Fig. 2e) can be seen anywhere in the infected erythrocytes; they are roundish or oval, each possesses a large nucleus and prominent cytoplasm. As parasite develops, gametocytes adhere to the erythrocyte nuclei and extend longitudinally along the nuclei markedly displacing them laterally (Fig. 2f).

Macrogametocytes (Fig. 2g): The cytoplasm is homogeneous in appearance, sometimes contains small vacuoles. Gametocytes markedly displace the nucleus of erythrocytes laterally, frequently to the envelope of the host cells (Fig. 2c, g). Parasite nucleus is oval or roundish usually more or less median in position (Fig. 2g); pigment granules are numerous (Table 2), oval and roundish, of medium size (0.5–1.0 μ m), usually randomly scattered throughout the cytoplasm.

Microgametocytes (Fig. 2h): The general configuration is as for macrogametocytes with the usual haemosporidian sexual dimorphic characters. The parasite nucleus is

diffuse, and its size is variable in different gametocytes (Fig. 2h).

Discussion

Molecular and microscopy approaches were combined in investigations of haemosporidian parasites of Skylarks for the first time during this study. The PCR-based methods were less sensitive in determining simultaneous infections than microscopic examination of blood films. That should be taken in consideration in field studies of blood parasites using general primers, as previously discussed by Valkiūnas et al. (2006) and Martínez et al. (2009).

P. relictum (lineages pSGS1) was the most prevalent haemosporidian parasite in Skylarks. That was expected because this lineage is widespread and actively transmitted in the Old World (Palinauskas et al. 2007; Bensch et al. 2009). Unidentified to species level, *Plasmodium* lineages pDELURB4 and pMOALB1 are closely related to pSGS1 and pGRW11 (Fig. 1), so probably belong to the same morphospecies *P. relictum*, but further morphological studies are needed to prove that.

Linkage between DNA sequences and identifications based on traditional morphological species can provide important information about life history strategies for parasitologists and evolutionary biologists studying phylogenetic relationships of these organisms; it can also be used for molecular identification of parasites (Križanauskienė et al. 2010). Unfortunately, the number of incorrectly identified species is increasing in GenBank (Valkiūnas et al. 2008a). To ensure parasites' species identification, we used museum type specimens in our identifications and also provided brief description of reported parasites in this study.

P. vaughani is second only to *P. relictum* in frequency of occurrence in birds. According to the previous studies, this avian malaria parasite has been reported in numerous bird species belonging to many host families and even orders, but is particularly common in passerines (Garnham 1966; Valkiūnas 2005), so the record of *P. vaughani* in Skylarks was not unexpected. In spite of worldwide distribution, molecular identification of *P. vaughani* has not been developed yet. That is important to do because the majority of natural malarial infections are light, so frequently are difficult to identify to species level in single blood films. We suggest using the lineage pSYAT05 for molecular identification of *P. vaughani*. Morphological features and measurements of blood stages of parasites of the lineage pSYAT05 are indistinguishable from *P. vaughani* from its type vertebrate hosts, the American robin. Gametocytes (Fig. 2i–k) and erythrocytic meronts (Fig. 2l), which are typical for *P. vaughani*, predominate among parasites of the pSYAT05 lineage in our material. Importantly, the same

lineage was found in the USA in the American robin, the type host of *P. vaughani* (Martinsen et al. 2008); it also present in the black bird, which is a common host of this parasite in Europe (Valkiūnas 2005; Hellgren et al. 2007). These data are in accord with former microscopic investigations, which showed cosmopolitan distribution and broad avian host range of *P. vaughani* (Garnham 1966; Corradetti and Scanga 1973; Bennett et al. 1982; Valkiūnas 2005). Formerly, *P. vaughani* was found in Skylarks only in Kazakhstan (prevalence is 3%, see Yakunin and Zhazylytaev 1977). This parasite has been reported in European Skylarks for the first time during this study.

We found the lineage SW2 of *Plasmodium* (*Novyella*) sp. only in Italian birds. This lineage is common in sedge warblers (*Acrocephalus schoenobaenus*) in Africa (Waldenström et al. 2002); it was found in Skylarks for the first time during this study. We were unable to identify this parasite to species level because intensity of infection was light.

The gametocytes of the lineages hALARV1, hALARV2 and hALARV3 are indistinguishable from each other in all their main qualitative and morphometric parameters. Comparison of blood stages of these Skylark parasites with the type specimens of *H. tartakovskiyi* (lineage hSISKIN1, Fig. 2a–d) from its type vertebrate host, the Common crossbill (Valkiūnas 1986) showed that all these haemoproteids are indistinguishable. We consider all these lineages as intraspecific variation of *H. tartakovskiyi* and attribute them this species. That is in accord to our phylogenetic analysis and is similar to the level of intraspecific variation reported, for instance in *Haemoproteus balmoralis* (Fig. 1). Lineages of *H. tartakovskiyi* are prevalent in Common crossbills, Hawfinches and Siskins in Europe (Bensch et al. 2009). This haemoproteid has been reported in Skylarks for the first time during this study. Because *H. tartakovskiyi* normally is prevalent in fringillid birds (Valkiūnas 1986), it might be that our report of this parasite is a case of new emerging haemosporidian infection in European Skylarks. *H. tartakovskiyi* is transmitted by biting midge *Culicoides impunctatus* (Valkiūnas and Liutkevičius 2002); its development in avian host and virulence remains unknown.

Plasmodium (*Novyella*) sp. (lineage pSW2) infection has been recorded in African migrating Sage Warbler (*Acrocephalus schoenobaenus*) (Waldenström et al. 2002). According to our study, this malaria parasite has been reported for the first time in Skylarks, so it should be considered as a possible new pathogen in this bird species, so it might contribute to decrease of Skylarks' population in Europe. It is interesting that the lineage pSW2 have been reported in non-migrant Tawny owl (*Strix aluco*) in Germany (Krone et al. 2008), so transmission certainly takes place in Europe.

Only two haemosporidian parasite species have been identified in Skylarks in Western Europe in the

twentieth century: *Haemoproteus alaudae* and *Plasmodium* (*Haemamoeba*) *supraecox* (Bennett et al. 1982; Valkiūnas 2005). Malaria parasites have been recorded incidentally in this bird, and the overall prevalence of haemosporidians was reported to be <10% before massive decline of Skylarks populations in Western Europe (Peirce 1981). Interestingly, these two haemosporidian parasites were not recorded during this study. However, we found at least four additional species (three *Plasmodium* and one *Haemoproteus*) and nine different mitochondrial DNA lineages of haemosporidians, which have not been reported in Skylarks in Western Europe before. It is worth noting that *P. relictum* has been reported in one of five examined Skylarks in Georgia (Burtikashvili 1978) and *P. vaughani* was seen in two of 65 examined Skylarks in Kazakhstan (Yakunin and Zhazylytaev 1977), so these parasites have broad range of distribution in Skylarks. Recently, *Plasmodium* spp. were found in 20% of Skylarks in France (Chavatte et al. 2009). However, during this study, we report highest prevalences and genetic diversity of malaria parasites and haemosporidians than ever been reported in this bird before. Additionally, *H. tartakovskiyi* seems to be an emerging parasite in Italian Skylarks because it has been reported only in birds belonging to the Fringillidae so far; it is prevalent in Siskins, Common crossbills and Hawfinch in Europe (Valkiūnas et al. 2003; Valkiūnas 2005). These data indicate possible ongoing colonization of Skylarks by blood parasites in changing environment conditions. That warrants further investigation, particularly due to recent reports on mortality of birds caused by emerging *Haemoproteus* spp. infections in Europe (Olias et al. 2011). It worth noting that transmission of the same lineages of haemosporidians between passerines of different families certainly occurs in Europe (Križanauskienė et al. 2006, 2010), but epidemiological significance of this phenomenon remains insufficiently understood.

It is important to note that significantly greater prevalence of pigment-forming haemosporidian infections was recorded in Skylarks sampled in Italy than in the Netherlands (Table 1). The birds sampled in Italy were caught during their seasonal migration; they probably came from Northeastern Europe to the study site and were on their way to wintering quarters in the Mediterranean.

Importantly, malarial parasites were prevalent in 5–7 days old nestlings during this study. Furthermore, all recorded *Plasmodium* spp. infections were simultaneous in nestlings; such infections have been reported to be particularly virulent in birds (Zehtindjiev et al. 2008; Palinauskas et al. 2009). Because *Plasmodium* spp. are particularly virulent and even lethal in juveniles (Garnham 1966; Valkiūnas 2005), it seems probable that malaria might be a factor contributing to mortality in European Skylarks due

to direct or indirect (via predators) elimination of nestlings and/or fledglings.

The Skylarks are “species of Conservation Concern in Europe”, owing to the populations’ decline of over 50%, particularly in West Europe since 1960s and mainly since 1980s (BirdLife International 2004). Although there are examples of dramatic impact of malaria parasites and other haemosporidians on wildlife, particularly in ecosystems where malaria has been emerging (Garnham 1966; Atkinson et al. 2001; Valkiūnas 2005; Levin et al. 2009), blood parasites have not yet been considered in understanding the decline of Skylark populations. This study does not provide final answer if blood parasites contribute to the decline of Skylarks populations; additional field studies comparing distribution and virulence of certain species of haemosporidians between areas where Skylarks are decreasing and where populations are stable are needed to answer this questions satisfactorily. However, our data indicate that haemosporidian parasites are likely emerging in Skylarks, so should be considered in conservation programmes. That is particularly true due to recent reports about mortality in birds caused by emerging haemosporidian infections in Europe (Olias et al. 2011; Valkiūnas 2011). Thus, the recent decline of Skylark populations in Western Europe is accompanied by increased prevalence and diversity of haemosporidian parasites, which should be considered as possible factors contributing to the population decline process and need further research.

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