

OPEN ACCESS

EDITED BY Alessia Libera Gazzonis, University of Milan, Italy

REVIEWED BY

Emma Gillingham, UK Health Security Agency (UKHSA), United Kingdom Gerardo Fracasso, University of Antwerp, Belgium

*CORRESPONDENCE Gergő Keve ⊠ keve.gergo@univet.hu

SPECIALTY SECTION This article was submitted to Parasitology, a section of the journal Frontiers in Veterinary Science

RECEIVED 18 January 2023 ACCEPTED 22 February 2023 PUBLISHED 22 March 2023

CITATION

Keve G, Csörgő T, Benke A, Huber A, Mórocz A, Németh Á, Kalocsa B, Tamás EA, Gyurácz J, Kiss O, Kováts D, Sándor AD, Karcza Z and Hornok S (2023) Ornithological and molecular evidence of a reproducing *Hyalomma rufipes* population under continental climate in Europe. *Front. Vet. Sci.* 10:1147186. doi: 10.3389/fvets.2023.1147186

COPYRIGHT

© 2023 Keve, Csörgő, Benke, Huber, Mórocz, Németh, Kalocsa, Tamás, Gyurácz, Kiss, Kováts, Sándor, Karcza and Hornok. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.

Ornithological and molecular evidence of a reproducing *Hyalomma rufipes* population under continental climate in Europe

Gergő Keve^{1,2*}, Tibor Csörgő^{3,4,5}, Anikó Benke^{5,6}, Attila Huber^{5,7}, Attila Mórocz^{5,8}, Ákos Németh^{5,9,10}, Béla Kalocsa⁵, Enikő Anna Tamás^{5,11}, József Gyurácz^{5,12}, Orsolya Kiss^{5,13}, Dávid Kováts^{4,5,14}, Attila D. Sándor^{1,2,15}, Zsolt Karcza⁵ and Sándor Hornok^{1,2}

¹Department of Parasitology and Zoology, University of Veterinary Medicine, Budapest, Hungary, ²ELKH-ÁTE Climate Change: New Blood-Sucking Parasites and Vector-Borne Pathogens Research Group, Budapest, Hungary, ³Department of Anatomy, Cell- and Developmental Biology, Eötvös Loránd University, Budapest, Hungary, ⁴Ócsa Bird Ringing Station, Ócsa, Hungary, ⁵BirdLife Hungary, Budapest, Hungary, ⁶Fenékpuszta Bird Ringing Station, Keszthely, Hungary, ⁷Aggtelek National Park Directorate, Jósvafő, Hungary, ⁸Duna-Dráva National Park Directorate, Pécs, Hungary, ⁹Kiskunság National Park Directorate, Kecskemét, Hungary, ¹⁰Kiskunság Bird Protection Association, Izsák, Hungary, ¹¹Faculty of Water Sciences, University of Public Service, Baja, Hungary, ¹²Department of Biology, Eötvös Loránd University, Savaria, Szombathely, Hungary, ¹³Faculty of Agriculture, Institute of Animal Sciences and Wildlife Management, University of Szeged, Hódmezővásárhely, Hungary, ¹⁴Hungarian Biodiversity Research Society, Budapest, Hungary, ¹⁵Department of Parasitology and Parasitic Diseases, University of Agricultural Sciences and Veterinary Medicine, Cluj-Napoca, Romania

Reports on adult Hyalomma ticks in certain regions of the Carpathian Basin date back to the 19th century. These ticks were thought to emerge from nymphs dropping from birds, then molting to adults. Although the role of migratory birds in carrying ticks of this genus is known from all parts of Europe, in most countries no contemporaneous multiregional surveillance of bird-associated ticks was reported which could allow the recognition of hotspots in this context. From 38 passeriform bird species 956 ixodid ticks were collected at seven locations in Hungary. Tick species were identified as *lxodes ricinus* (n = 598), Ixodes frontalis (n = 18), Ixodes lividus (n = 6), Haemaphysalis concinna (n= 321), and D. reticulatus (n = 1). All 12 Hyalomma sp. ticks (11 engorged nymphs and 1 unengorged larva) were identified as H. rufipes based on three mitochondrial markers. This species was only found in the Transdanubian region and along its southeastern border. The Common Blackbird and the European Robin were the two main hosts of I. ricinus and I. frontalis, whereas H. concinna was almost exclusively collected form long-distance migrants. The predominant hosts of H. rufipes were reed-associated bird species, the Sedge Warbler and the Bearded Reedling, both in their nesting period. This study provides ornithological explanation for the regional, century-long presence of adult Hyalomma ticks under continental climate in the Transdanubian Region of the Carpathian Basin. More importantly, the autochthonous occurrence of a H. rufipes population was revealed for the first time in Europe, based on the following observations: (1) the bird species infested with H. rufipes are not known to migrate during their nesting period; (2) one larva was not yet engorged; (3) the larva and the nymphs must have belonged to different local generations; and

(4) all *H. rufipes* found in the relevant location were identical in their haplotypes based on three maternally inherited mitochondrial markers, probably reflecting founder effect. This study also demonstrated regional and temporal differences in tick species carried by birds.

KEYWORDS

bird migration, Ixodes, Haemaphysalis, Hyalomma, Central Europe, Hungary

Introduction

In the temperate zone of Europe, pathogens transmitted by hard ticks (Acari: Ixodidae) are responsible for the majority of the vector-borne diseases (1). On this continent \sim 55 ixodid species occur (2). From among these, the number of tick species that are regarded as indigenous will likely increase in several countries, in part due to climate change and the emergence of new, thermophilic tick species from the south.

In this scenario, the first prerequisite for the establishment of new tick species in any region is their (repeated) introduction, for which a very important natural route is via bird migration. Migratory birds are long-known carriers of ticks, most importantly Hyalomma species, from the south to temperate regions of Europe (3), even its northernmost parts (4). However, birds usually carry immature ticks, larvae and nymphs of Hyalomma species (5), therefore in case of these thermophilic ticks, another crucial prerequisite prior to establishment is the ability of nymphs detaching from birds to molt to adults. This was already reported for both Hyalomma marginatum and Hyalomma rufipes from several countries north of the Mediterranean Basin, as exemplified by the UK (6), and the Netherlands (7) in western Europe, Sweden in northern Europe (8) or Hungary in Central Europe (9). Consequently, Hyalomma adults might also overwinter (10), increasing the chances for future establishment of permanent, reproductive populations.

Recently, the emergence of Hyalomma marginatum was reported in a previously non-endemic region of the Mediterranean Basin in southern France, but it was stated that even in such newly invaded areas this tick species probably remains exclusively Mediterranean and cannot expand outside this climatic range (11). On the other hand, north of the Mediterranean region, in the Carpathian Basin [geographically including both Hungary and the Transylvanian Basin: (12)], adult ticks from the genus Hyalomma are long-known for their autochthonous occurrence under continental climate. This was already reported in the 19th century (13), and later confirmed (14, 15). At the same time, in the absence of detailed morphological description, the species in the Carpathian Basin remained uncertain, because some hints were more relevant to H. rufipes [e.g., the name Hyalomma aegyptium: (15)], while others to H. marginatum [as implied in the predominance of the species referred to from Hungary in the Mediterranean Basin: (14)]. More recently, H. rufipes adults were found on cattle on two occasions in Hungary (9), and one adult on the same host species 10 years later by citizen science method (16).

Interestingly, these century-long reports on the presence of adult *Hyalomma* ticks in the Carpathian Basin attest that the chance for their occurrence is more likely in certain endemic areas of the country. However, this hypothesis was not yet tested from the point of view of bird migration, despite the long-known import of *Hyalomma* nymphs by birds into this geographical region (17). In light of the above, the aim of this study was to perform a pilot survey focusing on the comparison of tick species carried or imported by birds at various locations in Hungary. These locations were meant to represent most of the Carpathian Basin where important stopover sites can be found along the Adriatic Flyway of bird migration.

Materials and methods

Sample collection

In this study, birds mist-netted at seven ringing stations (Figure 1) by standard ornithological mist-nets (mesh size 16 mm) were examined for the presence of ticks, between March and November, 2022. The main characteristics of ringing stations are as follows:

- Tömörd Bird Ringing Station (coordinates: 47°21'N, 16°39'E): situated in northwestern Hungary, next to a small lake. It is surrounded by cultivated lands, bushes and deciduous forest, predominantly oak trees.
- (2) Ócsa Bird Ringing Station (47°19'N, 19°13'E): situated in north central Hungary, on the edge of a wetland. It is surrounded by arable fields, poplar plantations with several interspersed open-pit gravel mines (18).
- (3) Bódva Valley Bird Ringing Station (coordinates: 48°27'N, 20°42'E): situated in northeastern Hungary, and located in the valley of Bódva River. The river is surrounded by the mosaics of gallery forests, wet meadows, *Prunus* scrubs and arable lands. The adjacent hillsides are covered mostly by oak forests.
- (4) Fenékpuszta Bird Ringing Station (46°44'N, 17°14'E): situated in southwestern Hungary, next to the largest lake in Central Europe, Lake Balaton. The vegetation type is predominantly reed.
- (5) Izsák, Lake Kolon Bird Ringing Station (coordinates: 46°46'N, 19°19'E): situated in central Hungary, across Lake Kolon in the reedbed. The vegetation type is typical for

Abbreviations: *cox*1, cytochrome *c* oxidase subunit I.

marshes, heavily covered with reeds. It is surrounded by cultivated lands and planted forests.

- (6) Dávod, Lake Földvár Bird Ringing Station (coordinates: 46°0'N, 18°51'E): situated in south central Hungary on the shores of Lake Földvár which is an oxbow lake, that was formed naturally from river Danube. The lake is surrounded by reedbed. Sand Martins (*Riparia riparia*) were ringed in the sand mines of Baja (coordinates: 46°12N, 18°58'E) which are close to this area.
- (7) Lake Fehér Ornithology Camp (coordinates: 46°20'N, 20°6'E). The camp is located near to Lake Fehér, which is large fishpond system the greatest saline lake. The vegetation type is reedbed with sparse shrubs and trees.

Ticks were removed from the skin of birds with fine tweezers and stored in 96% ethanol. Data of collection (date, location, avian host species, ring number) were recorded. *Ixodes* and *Haemaphysalis* species were identified morphologically (2), whereas *Hyalomma* species molecularly as outlined below.

For data comparison and presentation, ornithological traits were assigned to bird species according to Csörgő et al. (19). English bird species names are capitalized in accordance with the international recommendations (https://bou.org.uk/british-list/bird-names/).

DNA extraction

Ticks of the genus *Hyalomma* were disinfected on their surface with sequential washing for 15 s in 10% NaClO, tap water and distilled water. For the DNA extraction, the larva was used without incision, whereas nymphs were cut dorsally on the idiosoma. DNA was extracted with the QIAamp DNA Mini Kit (QIAGEN, Hilden, Germany) according to the manufacturer's instruction, including an overnight digestion in tissue lysis buffer and Proteinase-K at 56 °C. Extraction controls (tissue lysis buffer) were also processed with the tick samples to monitor cross-contamination.

Molecular identification of *Hyalomma* species

The cytochrome oxidase subunit I (*cox*1) gene was chosen as the first target for molecular analysis. The PCR was modified from Folmer et al. (20) and amplifies an ~710-bp-long fragment of the gene. The primers HCO2198 (5'-TAA ACT TCA GGG TGA CCA AAA AAT CA-3') and LCO1490 (5'-GGT CAA CAA ATC ATA AAG ATA TTG G-3') were used in a reaction volume of 25 μ l, containing 1 U (stock 5 U/ μ l) HotStarTaq Plus DNA Polymerase, 2.5 μ l 10 × CoralLoad Reaction buffer (including 15 mM MgCl₂), 0.5 μ l PCR nucleotide Mix (stock 10 mM), 0.5 μ l of each primer (stock 50 μ M), 15.8 μ l ddH2O and 5 μ l template DNA. For amplification, an initial denaturation step at 95°C for 5 min was followed by 40 cycles of denaturation at 94°C for 40 s, annealing at 48°C for 1 min and extension at 72°C for 1 min. Final extension was performed at 72°C for 10 min.

Another PCR was used to amplify an ~460-bp-fragment of the 16S rDNA gene of Ixodidae (21), with the primers 16S+1 (5'-CTG CTC AAT GAT TTT TTA AAT TGC TGT GG-3') and 16S-1 (5'-CCG GTC TGA ACT CAG ATC AAG T-3'). Other reaction components, as well as cycling conditions were the same as above, except for annealing at 51°C. In addition, a conventional PCR reaction was used with the primer pairs T1B (5'-AAA CTA GGA TTA GAT ACC CT-3') and T2A (5'-AAT GAG AGC GAC GGG CGA TGT-3') to amplify an ~360-bp-long fragment from the 12S rRNA gene from all DNA extracts (22, 23). The PCR was modified with the following conditions. An initial denaturation step at 95°C for 5 min was followed by 5 cycles of denaturation at $94^\circ C$ for 30 s, annealing at 50° C for 30 s and extension at 72° C for 30 s and 30 cycles of denaturation at 94°C for 30 s, annealing at 53°C for 30 s and extension at 72°C for 30 s. Final extension was performed at 72°C for 7 min (23, 24).

Sequencing

In all PCRs non-template reaction mixture served as negative control. Extraction controls and negative controls remained PCR negative in all tests. Purification and sequencing of the PCR products were done by Biomi Ltd. (Gödöllő, Hungary). Quality control and trimming of sequences were performed with the BioEdit program, then alignment with GenBank sequences by the nucleotide BLASTN program (https://blast.ncbi.nlm.nih.gov). New sequences were submitted to GenBank (*cox*1: OQ108291-OQ108294, 16S rRNA: OQ103402-OQ103405, 12S rRNA: OQ103398-OQ103401).

Statistical analyses

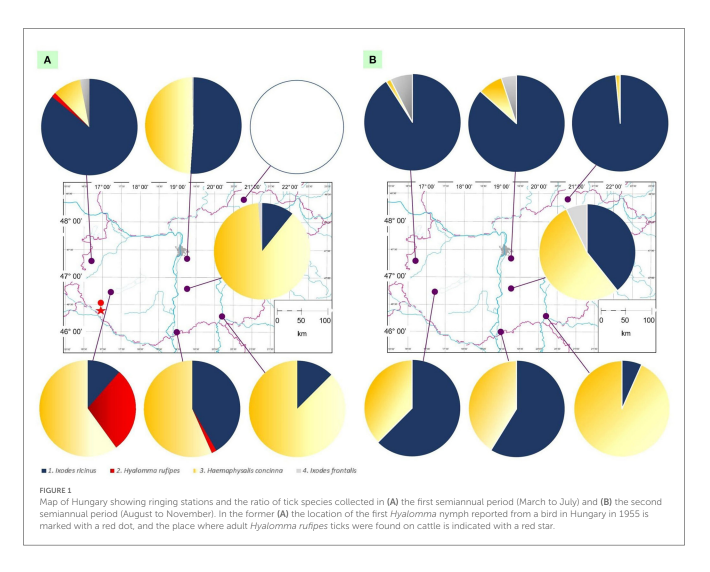
Fisher exact test (https://www.langsrud.com/fisher.htm) was used to compare prevalence rates between tick species and differences were regarded significant if P < 0.05.

Results

Species and developmental stages of ticks infesting birds

During 2022, 539 individuals of 38 passeriform bird species were found to be tick-infested (~39'000 birds were caught during the study period), from which altogether 956 ixodid ticks were collected (505 from March to July, and 451 from August to November: Table 1). The majority of developmental stages were nymphs (n = 588), but 352 larvae and 16 females were also present. On most birds (n = 381) only a single tick was found. The maximum number of ticks removed from a single bird was 30, and the mean intensity of tick-infestation was 1.78 tick/tick-infested bird in the whole study period.

Based on morphological characteristics, the ticks belonged to the following species: *Ixodes ricinus* (n = 598), *Ixodes frontalis* (n = 1000), *Ixodes frontalis* (n = 10000), *Ixodes frontalis* (n = 100000), *Ixodes frontalis* (n = 1000000), *Ixodes frontalis* (n = 10000000), *Ixodes frontalis* (n = 10



18), *Ixodes lividus* (n = 6), *Haemaphysalis concinna* (n = 321), and *D. reticulatus* (n = 1) (Supplementary Table 1). Morphologically, the twelve *Hyalomma* sp. ticks could only be identified on the genus level and so molecular identification was necessary. All *Hyalomma* nymphs were in a similar, advanced state of engorgement, but the single larva was flattened, appeared unengorged.

Based on the 16S rRNA gene, Hyalomma nymphs belonged to three haplotypes (Table 2). One of these collected in south Hungary (OQ103402) had 100% (383/383 bp) sequence identity to H. rufipes previously collected from a bird in north-central Hungary (Ócsa: KU170517) and another in Egypt (MK737650). The second haplotype (collected in northwest Hungary: OQ103403) differed in two, and the third haplotype (all other specimens: OQ103404-OQ103405) in one position of their 16S rRNA sequence, meaning 99.5% and 99.7% sequence identities to the above two reference sequences, respectively (Table 2). One haplotype (OQ108291) differed in one position, whereas all other H. rufipes specimens were 100% (645/645 bp) identical in the sequenced part of their cox1 gene (OQ108292-OQ108294) to a tick collected from a Eurasian Reed Warbler (Acrocephalus scirpaceus) in the Netherlands (MT757612) and another reported from Malta (OL339477). Interestingly, these cox1 sequences were more different (in two bps) from H. rufipes collected from a bird in a previous study in north-central Hungary (Ócsa: KU170491). In addition, all *H. rufipes* nymphs and the larva had identical 12S rRNA sequences (OQ103398-OQ103401), with 100% (341/341 bp) sequence identity to ticks collected from birds in Malta (OL352890) and in Italy (MW175439). Thus, the genus *Hyalomma* was exclusively represented by *H. rufipes* (n = 12).

Host-associations of tick species and the migratory habits of their avian hosts

Associations of ticks collected in this study with different bird species are summarized in Supplementary Table 2. The Common Blackbird (*Turdus merula*) ($n_{birds} = 58$, with 22.4% of total *I. ricinus* ticks) and the European Robin (*Erithacus rubecula*) ($n_{birds} = 105$, with 32.4% of total *I. ricinus* ticks) were the two main hosts of *I. ricinus* in both the spring and the autumn tick collection periods. The preferred hosts of *I. frontalis* were also these two bird species ($n_{birds} = 4$, with 22.2% of total *I. frontalis* ticks; and 5, with 38.9% of all *I. frontalis* ticks, respectively). *Haemaphysalis concinna* most often infested the Sedge Warbler (*Acrocephalus schoenobaenus*) ($n_{birds} = 49$, with 17.1% of total *H. concinna* ticks) and Savi's

Spring migration and summer nesting period (March–July):										
Tick species	Male	Female	Nymph	Larva	Total					
Ixodes ricinus	0	0	190	28	218					
Haemaphysalis concinna	0	0 179		85	264					
Hyalomma rufipes	0	0 11		1	12					
Ixodes frontalis	des frontalis 0		1	1	4					
Ixodes lividus	lividus 0		0	0	6					
Dermacentor reticulatus 0		1 0		0	1					
Autumn migration period (August–November):										
Ixodes ricinus	0	0	167	213	380					
Haemaphysalis concinna	0	0	37	20	57					
Hyalomma rufipes	0	0	0	0	0					
Ixodes frontalis	0	7	3	4	14					

TABLE 1 Total number of ticks collected in the first and second semiannual periods, according to their species and developmental state.

TABLE 2 Data of Hyalomma ticks collected in 2022 from birds at various ringing stations in Hungary.

lsolate code	Bird species	Date	Region of <i>Hyalomma</i> sp. Hungary (number, stage (location)		GenBank accession numbers according to the three genetic markers		
					16S rRNA	Cox1	12S rRNA
BA2	SYL COM	May 14	South (Dávod)	H. rufipes (1xN)	OQ103402	OQ108291	OQ103398
GJ10	FIC HYP	April 23	Northwest (Tömörd)	H. rufipes (1xN)	OQ103403	OQ108292	OQ103399
BE02	ACR SCH	June 26	Southwest (Fenékpuszta)	H. rufipes (1xN)	OQ103404	OQ108293	OQ103400
BE03	ACR SCH	June 26	Southwest (Fenékpuszta)	H. rufipes (1xN)	OQ103404	OQ108293	OQ103400
BE04	ACR SCH	June 26	Southwest (Fenékpuszta)	H. rufipes (1xN)	OQ103404	OQ108293	OQ103400
BEO5	PAN BIA	June 26	Southwest (Fenékpuszta)	H. rufipes (1xN)	OQ103405	OQ108294	OQ103401
BE06	PAN BIA	June 26	Southwest (Fenékpuszta)	H. rufipes (1xL, 5xN)	OQ103405	OQ108294	OQ103401

Identical background color in cells of the same column of a genetic marker indicates identical sequences

ACR SCH, Acrocephalus schoenobanenus; FIC HYP, Ficedula hypoleuca; PAN BIA, Panurus biarmicus; SYL COM, Sylvia communis.

Warbler (*Locustella luscinioides*) ($n_{birds} = 65$, with 55.1% of all *H. concinna* ticks) (Supplementary Table 3). *Hyalomma rufipes* was only collected on repeated occasions from Sedge Warblers (*A. schoenobaenus*) and Bearded Reedling (*Panurus biarmicus*), and once from a Common Whitethroat (*Curruca communis*) and from a European Pied Flycatcher (*Ficedula hypoleuca*). *Ixodes lividus* was only found once, on its specific host, the Sand Martin (*Riparia riparia*). Importantly, with the exception of the accidental finding of a single *D. reticulatus* female on a Common Blackbird, all other females (n = 16) belonged to the two ornithophilic tick species *I. frontalis* (n = 9) and *I. lividus* (n = 6).

During the spring, at the ringing station in north-central Hungary (Ócsa) where the highest number of tick-infested birds were caught and which contributed the most balanced ratio of birds with different migratory habits to the study, there was a highly significant (P < 0.0001) difference between the host associations of *I. ricinus* and *H. concinna*, since the former predominated on resident and short-distance migrant bird species, but *H. concinna* on long-distance migrants. In the autumn, taking into account all ringing stations, the difference between these two tick species in the

same comparison, and the association of *I. frontalis* with resident and short-distance migrant bird species was also highly significant (P < 0.0001).

Spatiotemporal occurrence of tick species

Ixodes ricinus and *H. concinna* were found to infest birds in both the spring and autumn collection periods (Figure 1), whereas the presence of *H. rufipes* was restricted to the first half of the year (Figure 1 and Tables 1, 2), and *I. frontalis* predominated in the autumn period (Figure 1). Importantly, *H. rufipes* was collected from long-distance migrant birds in south and northwest Hungary in May and April, respectively (Table 2). However, all other specimens of this species were removed from birds in the middle of summer (late June) at one ringing station in the southwestern part of the country (Fenékpuszta).

During the spring period, *I. ricinus* was the predominant tick species in the north, whereas *H. concinna* in central and south Hungary (Figure 1 and Supplementary Table 1). However, in the

autumn, *I. ricinus* represented the highest number of ticks from birds in north as well as in southwestern parts of the country, and *H. concinna* at two ringing stations, in central and southeast Hungary (Figure 1). *Hyalomma rufipes* was only found in the Transdanubian region and in one case along the southern reach of the Danube. On the other hand, *I. frontalis* could only be collected in northern and central locations during both spring and autumn and was absent from birds in southern parts of the country (Figure 1).

Taken together, *I. ricinus* and *H. concinna* occurred on birds at all sampling sites, but their ratio was different according to these sites and semiannual periods. At the same time, the spatiotemporal distribution was limited in case of *H. rufipes* and *I. frontalis*.

Discussion

In Hungary, studies on tick-infestations of birds date back to more than half a century (17), and have been extensively performed on annual or tri-annual bases focusing on the same ringing station in the north-central part of the country [Ócsa: (25-27)]. Similar reports on ticks from avian hosts are available from numerous European countries, as exemplified by Sweden (28), The Netherlands and Belgium (29), Germany (30), or Italy (31). Relevant studies have also been reviewed recently (32, 33). However, leaving aside opportunistic and sporadic collections of ticks from birds, the present study is the first "horizontal tick survey" from birds in the Carpathian Basin and probably also in a broader geographical context. This implies that ticks were removed and their species identified at several ringing stations simultaneously in the course of 1 year, allowing not only the regional comparison of tick burdens carried by birds, but also assessing the significance and need of similar studies on a larger, continental scale.

In this study, six species of ixodid ticks (three prostriate and three metastriate) were collected from birds. The most significant finding related to tick species diversity was the *H. rufipes*-infestation of three long-distance migrant and a resident bird species. *Hyalomma rufipes* is the vector of Crimean– Congo haemorrhagic fever virus, *Babesia occultans, Rickettsia aeshlimannii, Ehrlichia* spp., *Coxiella burnetii, Rickettsia conorii* and *Anaplasma marginale* (2).

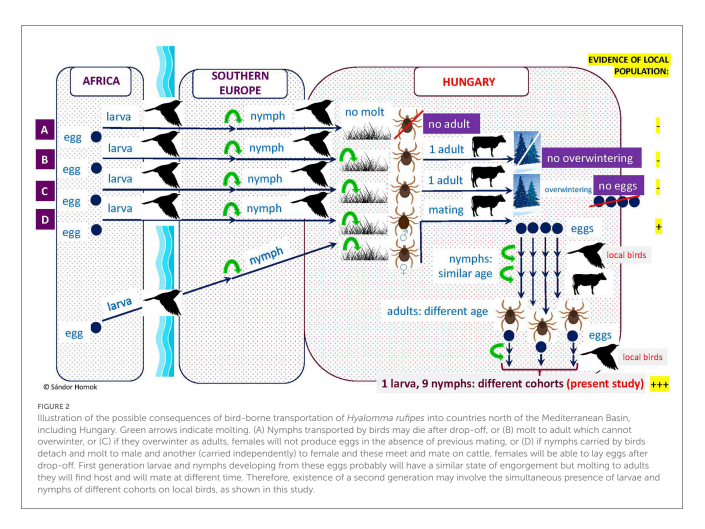
Importantly, *H. rufipes* was collected in south and northwestern Hungary during late spring in 2022, as in a previous study (27). However, in this study all remaining 10 specimens were removed from birds in the middle of summer (late June) at one ringing station in the southwestern part of the country (Fenékpuszta), i.e., in the same county (Zala) where *Hyalomma*-infestation of a bird was diagnosed for the first time in Hungary in 1955 (17). In the same region, *Hyalomma* sp. ticks were reported to occur (34) and *H. rufipes* adults were identified on two occasions from cattle (9) (Figure 1A).

It is utterly unlikely that all five individuals of the two avian host species of these 9 fully engorged nymphs and one unengorged larva of *H. rufipes* (sampled on June 26) carried these ticks into Hungary from abroad. *Hyalomma rufipes* has a two-host life cycle, and engorged nymphs drop off from the host after 21– 29 days of infestation (35). One of the avian hosts shown to harbor nymphs of *H. rufipes* in this study, the Sedge Warbler (*A. schoenobaenus*) typically arrives in Hungary from the wintering grounds in Africa between April and early May (19), and late June (when its *Hyalomma*-infestation was diagnosed) is in the middle of its nesting period, without migration. On the other hand, the other repetitive host of *H. rufipes* in this study, the Bearded Reedling (*P. biarmicus*) is an *a priori* resident bird species, with rarely documented limited movements, but according to ringing data (19) these "vagrancies" never occur in its summer nesting period.

Regarding the results of molecular analyses, it is not surprising that all H. rufipes individuals collected in 2022 from birds in Hungary (n = 12) had identical 12S rRNA haplotypes, because this genetic marker was shown to be identical in case of a much larger set of *H. rufipes* ticks (n = 48) collected from birds with probably different geographical origin (36). However, in this study the sequenced part of the cox1 gene was also identical between all H. rufipes (n = 11) collected in the Transdanubian part of Hungary, in particular in case of those 10 ticks which were removed from birds at the same ringing station in southwest Hungary (Fenékpuszta). Hyalomma rufipes was shown to differ remarkably in its cox1 haplotype in case of ticks carried by birds with different geographical origin (36). Moreover, the ratio and presence or absence of certain Hyalomma cox1 haplotypes were demonstrated to be site- and population-specific, usually with multiple haplotypes even within the same population (37). Therefore, finding of exclusively one cox1 haplotype among 10 H. rufipes ticks collected in one location (Fenékpuszta) raises the possibility that these ticks represent the same population. Their genetic similarity is probably a consequence of founder effect. Taken together, all three studied mitochondrial, maternally inherited genetic markers were identical only between H. rufipes individuals collected in the latter place, also supporting the common maternal aborigine of these ticks.

In addition, the apparently unengorged state of the *H. rufipes* larva on one of these birds also argues against the foreign origin of its tick-infestation. Note that in a previous study only molting (i.e., advanced stage) *H. marginatum* larvae were found on birds in Hungary, and all other stages were nymphs (26, 27). Importantly, hitherto molecularly verified *H. rufipes* larvae were only reported from birds in south European countries [reviewed by Keve et al. (33)], and typically only nymphs of this tick species arrive on birds in countries north of the Mediterranean Basin if these originate from Africa [Figure 2; (33)].

Probably not all ticks carried by migratory birds in the spring were imported by them from southern countries, as they can become infested with ticks of the local fauna, particularly when avian hosts arrive from their wintering grounds during the activity peak of local tick populations. Similarly to previous bird tick studies in the Carpathian Basin (27) and most countries north of the Mediterranean Basin [e.g., (28)], I. ricinus was the tick species most commonly collected from birds in 2022 in Hungary. Haemaphysalis concinna was the second most abundant tick species on birds, which, however, seems to be unique to the Carpathian Basin and its region (33). Both of these tick species (I. ricinus, H. concinna) indigenous to Hungary tend to infest birds which arrive in their main activity periods (38), therefore I. ricinus (peak activity: April) is mainly found on residents and short-distance migrants (typically arriving early spring), whereas H. concinna (peak activity: May) on long-distance migrants (usually arriving late spring) (19).



New tick-host associations revealed in this study include the presence of *H. rufipes* on the Bearded Reedling (*P. biarmicus*), and infestation of Moustached Warbler (*Acrocephalus melanopogon*) with *H. concinna*. Although *D. reticulatus* seldom occurs on birds (33), its immature developmental stages were reported from avian hosts (including the Common Blackbird, *T. merula*) (39). The collection of its adult on a Blackbird during this study is probably an accidental finding.

Considering the regional occurrence of tick species on birds in the Carpathian Basin, *H. concinna* is a thermophilic tick species (40), and this is in accordance with its predominance on birds in central-south Hungary during the spring, and central-southeastern Hungary in the autumn (i.e., the warmest regions of the country: Supplementary Figure 1). On the other hand, the reason for the absence of *I. frontalis* from birds in the southern part of Hungary maybe twofold. First, the relevant sampling locations are near water surfaces where the predominant bird species (e.g., Savi's Warbler, *L. luscinioides*) are not known to be hosts or (e.g., the Sedge Warbler, *A. schoenobaenus*) are exceptional hosts of this tick species (33). Second, in these places bird mist-netting (i.e., tick collection) was terminated sooner than the late autumn peak activity of *I. frontalis* in the relevant region (41).

In this study, *H. rufipes* was only found in the Transdanubian region and once along the southern Danube, in line with the reported 130-year-long endemicity of *Hyalomma* species in the country (13–15). While *Hyalomma*-infestation was previously

reported on non-water-associated bird species (*E. rubecula*, *C. communis*) in the springtime in north-central Hungary (Ócsa) (26, 27), this is the first occasion when ticks of this genus were observed on reed-dwelling birds in another region of Hungary, in a different season (during summer). This also raises the question on what the differences between the relevant two habitats in terms of landscape, vegetation and avian hosts are.

Fenékpuszta Bird Ringing Station is situated next to Lake Balaton. Here, the reedbed habitat in the riparian zone narrows to about 150 meters at the site of the mist-nets, where 12 pieces of these stretch across the reedbed completely. Due to uninterrupted reeds, this is an important stopover site for migrating passerines, particularly *Acrocephalus*-species. Based on ringing data, mostly long-distance migrant Sedge Warblers (*A. schoenobaenus*) and Eurasian Reed Warblers (*A. scirpaceus*) stop in this area, but Great Reed Warblers (*A. arundinaceus*) and Savi's Warblers (*L. luscinioides*) are also significant in numbers.

Conversely, in Ócsa Bird Ringing Station the heterogeneous reedbed habitats of the capture locations are interspersed with fast growing shrubs as elderberry (*Sambucus nigra*) and blackberry (*Rubus fruticosus*), with softwood stands (*Salix* spp. and *Populus* spp.) forming most of the vegetation. Thus, the Eurasian Blackcap (*Sylvia atricapilla*) and the European Robin (*Erithacus rubecula*) are two most common short-to-mid-distance migratory species here (18). Regarding the capture rates of the species groups of migrating passerines, there is a significant difference between the

homogeneous reedbed and other habitats (where the reedbed is patchy and alternates with deciduous forests, berry bushes). While *Acrocephalus* spp. account for the largest proportion of birds caught in Fenékpuszta, bush-dwelling warblers present a higher portion in Ócsa.

Based on the above, the existence of at least one indigenous population of *H. rufipes* is evidenced in the western part of Transdanubia, near Lake Balaton, because of the following reasons: (1) most importantly, the recognized avian hosts of *H. rufipes* were extremely unlikely to arrive from abroad shortly prior to their examination, especially not all five of them; (2) one larva was not yet engorged; (3) the larva and the nymphs (in a similar state of engorgement) were probably offspring of two females and must have belonged to different local generations (Figure 2); and (4) all *H. rufipes* found in the relevant location were identical in their haplotypes based on three maternally inherited mitochondrial markers, probably reflecting founder effect.

In addition, adults of H. rufipes are known to occur in the western part of the Carpathian Basin for 130 years, and in the same county (Zala) with its present collections adults of this tick species were found to infest cattle repeatedly (9). Small local populations of H. rufipes were proposed to explain the occasional presence of H. rufipes in Russia (42, 43) and its populations in scattered areas are also known in north Africa (43, 44). However, to our knowledge, this is the first report of a similar phenomenon and its evidence from Europe. One of the most important limiting factor for the survival of this xerophilic tick species under any climate is thought to be the maximum level of precipitation (annual rainfall) which is around 650 mm in southwestern Hungary (Supplementary Figure 1), i.e., similar to what is well-tolerated by H. rufipes in its range within Africa (45, 46). Populations of these ticks probably can survive winter conditions as adults in southwestern Hungary where winter temperatures are among the mildest in the country (Supplementary Figure 1). Nevertheless, H. rufipes is known to have populations in regions with up to 120 days of frost (43). It is also noteworthy here that the likely overwintering of H. rufipes was reported in the Czech Republic (10), north of Hungary. Importantly, the discovered H. rufipes population might act as a "stepping-stone" for this tick species during its northward transportation by birds which use the relevant habitat near Lake Balaton in southwestern Hungary as a stopover site (see above).

On the other hand, no evidence was gained for any further *Hyalomma* populations indigenous in other regions of Hungary, as also indicated by the overall absence of *Hyalomma* ticks from birds in the autumn migration period. Thus, also taking into account the over-century-long presence of adult *Hyalomma* ticks, up to now there was no evidence for their emergence in the Carpathian Basin, but here evidence is reported for the emergence of a local population for the first time.

Similarly relevant to a broader, international context, the most important aim of the present study was also fulfilled, i.e., it was successful to demonstrate differences between sampling sites, indicating that in the above context single-site surveys may be biased (not informative) on the actual risk posed by birds in transporting ticks in a geographical region or country. Therefore, to assess the emergence or increasing presence of a *Hyalomma* species, ticks should be collected (larvae and nymphs from birds, and/or adults from reproductive hosts) extensively and annually in different regions of suspected endemic areas, preferentially by unbiased professionals who should stick to a standard methodology (sampling protocol).

Data availability statement

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article/Supplementary material.

Ethics statement

The Animal Welfare Committee of the University of Veterinary Medicine, Budapest, confirmed that the research does not constitute animal experimentation in accordance with Act No. 28 of 1998 and Government Decree No. 40 of 2013 on animal experiments. Therefore, notification or permission from the Animal Protection Authority was not required. The University Committee raised no objections to the tests either from an ethical or animal welfare point of view. All songbirds were handled and released by experienced ringers of BirdLife Hungary. License for bird ringing was issued by the Pest County Government Authority (https://www.mme.hu/sites/default/files/pe_ktf_97_13_ 2017_vvt.pdf).

Author contributions

GK: conceptualization, study design, sample collection, tick species identification, and manuscript writing. TC, AB, AH, AM, ÁN, BK, ET, JG, and OK: study design, sample collection, and data curation. DK: ornithological categorization and manuscript writing. AS: study design and supervision. ZK: study organization and data availability. SH: conceptualization, study design, DNA extraction, molecular analyses, manuscript writing, and preparation of figures. All authors contributed to the article and approved the submitted version.

Funding

This study was funded by the Eötvös Loránd Research Network (ELKH), Hungary (Project No. 1500107). The open access publication of this study was funded by Project no. TKP2020-NKA-01 implemented with the support provided from the National Research, Development and Innovation Fund of Hungary, financed under the Tématerületi Kiválósági Program 2020 (2020-4.1.1-TKP2020) funding scheme.

Acknowledgments

The authors are grateful to Ms. Nóra Takács for performing PCRs, to Dr. Jenő Kontschán for his advice in the terminology and definition of populations and to Németh Anna for her participation in field trips. The authors also thank all staff members and ringing personnel who contributed to the sample collection.

Conflict of interest

Authors TC, AB, AH, AM, ÁN, BK, ET, JG, OK, DK, and ZK were employed by BirdLife Hungary.

The remaining authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Publisher's note

All claims expressed in this article are solely those of the authors and do not necessarily represent those of their affiliated organizations, or those of the publisher, the editors and the reviewers. Any product that may be evaluated in this article, or claim that may be made by its manufacturer, is not guaranteed or endorsed by the publisher.

References

1. Rochlin I, Toledo A. Emerging tick-borne pathogens of public health importance: a mini-review. *J Med Microbiol.* (2020) 69:781–91. doi: 10.1099/jmm.0.001206

2. Estrada-Peña A, Mihalca AD, Petney TN. Ticks of Europe and North Africa: A Guide to Species Identification. Cham: Springer (2017).

3. Hoogstraal H, Kaiser MN, Traylor MA, Gaber S, Guindy E. Ticks (Ixodoidea) on birds migrating from Africa to Europe and Asia. *Bull World Health Organ.* (1961) 24:197.

4. Brinck P, Svedmyr A, von Zeipel G. Migrating birds at Ottenby Sweden as carriers of ticks and possible transmitters of tick-borne encephalitis virus. *Oikos*. (1965) 16:88–99. doi: 10.2307/3564868

5. Capek M, Literak I, Kocianova E, Sychra O, Najer T, Trnka A, et al. Ticks of the *Hyalomma marginatum* complex transported by migratory birds into Central Europe. *Ticks Tick-Borne Dis.* (2014) 5:489–93. doi: 10.1016/j.ttbdis.2014.03.002

6. Hansford KM, Carter D, Gillingham EL, Hernandez-Triana LM, Chamberlain J, Cull B, et al. *Hyalomma rufipes* on an untraveled horse: Is this the first evidence of *Hyalomma* nymphs successfully moulting in the United Kingdom? *Ticks Tick-Borne Dis.* (2019) 10:704–8. doi: 10.1016/j.ttbdis.2019.03.003

7. Uiterwijk M, Ibanez-Justicia A, van de Vossenberg B, Jacobs F, Overgaauw P, Nijsse R, et al. Imported *Hyalomma* ticks in the Netherlands 2018-2020. *Parasit Vectors*. (2021) 14:244. doi: 10.1186/s13071-021-04738-x

 Grandi G, Chitimia-Dobler L, Choklikitumnuey P, Strube C, Springer A, Albihn A, et al. First records of adult Hyalomma marginatum and H. rufipes ticks (Acari: Ixodidae) in Sweden. Ticks Tick-Borne Dis. (2020) 11:101403. doi: 10.1016/j.ttbdis.2020.101403

9. Hornok S, Horváth G. First report of adult *Hyalomma marginatum rufipes* (vector of Crimean-Congo haemorrhagic fever virus) on cattle under a continental climate in Hungary. *Parasit Vectors*. (2012) 5:170. doi: 10.1186/1756-3305-5-170

10. Rudolf I, Kejikova R, Vojtisek J, Mendel J, Penazziova K, Hubalek Z, et al. Probable overwintering of adult *Hyalomma rufipes* in Central Europe. *Ticks Tick-Borne Dis.* (2021) 12:101718. doi: 10.1016/j.ttbdis.2021.101718

11. Bah MT, Grosbois V, Stachurski F, Muñoz F, Duhayon M, Rakotoarivony I, et al. The Crimean-Congo haemorrhagic fever tick vector *Hyalomma marginatum* in the south of France: Modelling its distribution and determination of factors influencing its establishment in a newly invaded area. *Transbound Emerg Dis.* (2022) 69:e2351–65. doi: 10.1111/tbed.14578

12. Krézsek C, Bally AW. The Transylvanian Basin (Romania) and its relation to the Carpathian fold and thrust belt: insights in gravitational salt tectonics. *Mar Pet Geol.* (2006) 23:405–42. doi: 10.1016/j.marpetgeo.2006.03.003

13. Karpelles L. Adalékok Magyarország atka-faunájához [Contributions to the mite-fauna of Hungary]. Math És Természettudományi Közlemények. (1893) 25:439–41.

14. Kotlán S. Adatok a hazai kullancs-fauna ismeretéhez [Data on the Hungarian tick fauna]. Állattani Közlöny. (1919) 18:33–6.

Supplementary material

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fvets.2023. 1147186/full#supplementary-material

SUPPLEMENTARY FIGURE 1

The average annual precipitation (https://www.met.hu/) and temperature (https://www.mozaweb.com/search?search_=_középhomérséklet) in January in Hungary, based on data from the Hungarian Meteorological Service (OMSZ). The site of the discovered *Hyalomma rufipes* population is marked with a star.

SUPPLEMENTARY TABLE 1

Number of ticks for each bird ringing station, shown according to tick species, developmental stages, and collection periods.

SUPPLEMENTARY TABLE 2

Avian host species that were found tick-infested in this study, shown according to collection site and spring or autumn migration intervals (the former including nesting period).

SUPPLEMENTARY TABLE 3

List of all ticks found in this study, sorted by date, collection site, and host species.

15. Kotlán S. A hazai kullancsok rendszere [The classification of the ticks of Hungary]. Állattani Közlöny. (1921) 20:43–50.

16. Földvári G, Szabó É, Tóth GE, Lanszki Z, Zana B, Varga Z, Kemenesi G. Emergence of *Hyalomma marginatum* and *Hyalomma rufipes* adults revealed by citizen science tick monitoring in Hungary. *Transbound Emerg Dis.* (2022). doi: 10.22541/au.164572450.0192 3223/v1

17. Janisch M. Kullancsgazda madarak különféle betegségek közvetítoi [Tick carrier birds spreading disease agents]. *Aquila*. (1960) 67–68:191–194.

18. Csörgő T, Harnos A, Rózsa L, Karcza Z, Fehérvári P. Detailed description of the Ócsa Bird Ringing Station, Hungary. Ornis Hung. (2016) 24:91-108. doi: 10.1515/orhu-2016-0018

19. Csörgő T, Karcza Z, Halmos G, Magyar G, Gyurácz J, Szép T, et al. *Magyar madárvonulási atlasz [Atlas of bird migration in Hungary]*. Budapest: Kossuth Kiadó (2009). p. 672.

20. Folmer O, Black M, Hoeh W, Lutz R, Vrijenhoek R. DNA primers for amplification of mitochondrial cytochrome c oxidase subunit I from diverse metazoan invertebrates. *Mol Mar Biol Biotechnol.* (1994) 3:294–9.

21. Black 4th WC, Piesman J. Phylogeny of hard-and soft-tick taxa (Acari: Ixodida) based on mitochondrial 16S rDNA sequences. *Proc Natl Acad Sci.* (1994) 91:10034–8. doi: 10.1073/pnas.91.21.10034

22. Beati L, Keirans JE. Analysis of the systematic relationships among ticks of the genera *Rhipicephalus* and *Boophilus* (Acari: Ixodidae) based on mitochondrial 12S ribosomal DNA gene sequences and morphological characters. *J Parasitol.* (2001) 87:32–48. doi: 10.1645/0022-3395(2001)0870032:AOTSRA2.0.CO;2

23. Bitencourth K, Voloch CM, Serra-Freire NM, Machado-Ferreira E, Amorim M, Gazêta GS. Analysis of *Amblyomma sculptum* haplotypes in an area endemic for Brazilian spotted fever. *Med Vet Entomol.* (2016) 30:342–50. doi: 10.1111/mve.12174

24. Burkman EJ. Genetic Structure of Amblyomma cajennense (Acari: Ixodidae) Populations Based on Mitochondrial Gene Sequences. Statesboro, GA: Georgia Southern University (2009).

25. Hornok S, Karcza Z, Csörgő T. Birds as disseminators of ixodid ticks and tickborne pathogens: note on the relevance to migratory routes. *Ornis Hung.* (2012) 20:86–9. doi: 10.2478/orhu-2013-0010

26. Hornok S, Csörgő T, de la Fuente J, Gyuranecz M, Privigyei C, Meli ML, et al. Synanthropic birds associated with high prevalence of tick-borne rickettsiae and with the first detection of Rickettsia aeschlimannii in Hungary. *Vector Borne Zoonotic Dis Larchmt N.* (2013) 13:77–83. doi: 10.1089/vbz.2012.1032

27. Hornok S, Flaisz B, Takács N, Kontschán J, Csörgő T, Csipak Á, et al. Bird ticks in Hungary reflect western, southern, eastern flyway connections and two genetic lineages of *Ixodes frontalis* and *Haemaphysalis concinna. Parasit Vectors*. (2016) 9:101. doi: 10.1186/s13071-016-1365-0

28. Wilhelmsson P, Jaenson TGT, Olsen B, Waldenstrom J, Lindgren P-E. Migratory birds as disseminators of ticks and the tick-borne pathogens Borrelia bacteria and tick-borne encephalitis (TBE) virus: a seasonal study at Ottenby Bird Observatory in Southeastern Sweden. *Parasit Vectors.* (2020) 13:607. doi: 10.1186/s13071-020-04493-5

29. Heylen D, Fonville M, Docters van Leeuwen A, Stroo A, Duisterwinkel M, van Wieren S, et al. Pathogen communities of songbird-derived ticks in Europe's low countries. *Parasit Vectors*. (2017) 10:497. doi: 10.1186/s13071-017-2423-y

30. Klaus C, Gethmann J, Hoffmann B, Ziegler U, Heller M, Beer M. Tick infestation in birds and prevalence of pathogens in ticks collected from different places in Germany. *Parasitol Res.* (2016) 115:2729–40. doi: 10.1007/s00436-016-5022-5

31. Toma L, Mancuso Ed'Alessio SG, Menegon M, Spina F, Pascucci I, Monaco F, et al. Tick species from Africa by migratory birds: a 3-year study in Italy. *Exp Appl Acarol.* (2021) 83:147–64. doi: 10.1007/s10493-020-00573-4

32. Buczek AM, Buczek W, Buczek A, Bartosik K. The potential role of migratory birds in the rapid spread of ticks and tick-borne pathogens in the changing climatic and environmental conditions in Europe. *Int J Environ Res Public Health*. (2020) 17:2117. doi: 10.3390/ijerph17062117

33. Keve G, Sandor AD, Hornok S. Hard ticks (Acari: Ixodidae) associated with birds in Europe: review of literature data. *Front Vet Sci.* (2022) 9:928756. doi: 10.3389/fvets.2022.928756

34. Janisch M. A hazai kullancsfauna feltérképezése [Geographical distribution of tick species in Hungary]. Állattani Közlöny. (1959) 47:103–10.

35. Magano SR, Els DA, Chown SL. Feeding patterns of immature stages of *Hyalomma truncatum* and *Hyalomma marginatum rufipes* on different hosts. *Exp Appl Acarol.* (2000) 24:301–13. doi: 10.1023/a:1006457325097

36. Hornok S, Cutajar B, Takács N, Galea N, Attard D, Coleiro C, et al. On the way between Africa and Europe: molecular taxonomy of ticks collected from birds in Malta. *Ticks Tick-Borne Dis.* (2022) 13:102001. doi: 10.1016/j.ttbdis.2022.102001

37. Márquez FJ, Caruz A. Phylogeography of *Hyalomma (Euhyalomma) lusitanicum* (Acarina, Parasitiformes, Ixodidae) in Andalusia based on mitochondrial cytochrome oxidase I gene. *Exp Appl Acarol.* (2021) 85:49–61. doi: 10.1007/s10493-021-00652-0

38. Hornok S. Allochronic seasonal peak activities of *Dermacentor* and *Haemaphysalis* spp. under continental climate in Hungary. *Vet Parasitol.* (2009) 163:366–9. doi: 10.1016/j.vetpar.2009. 03.048

39. Akimov IA, Nebogatkin IV. Distribution of Ticks from the Genus Dermacentor (Acari, Ixodidae) in Ukraine. BecmHuk 300π osuu. (2011) 45:e-1. doi: 10.2478/v10058-011-0001-x

40. Hubálek Z, Halouzka J, Juricova Z. Host-seeking activity of ixodid ticks in relation to weather variables. *J Vector Ecol.* (2003) 28:159–65.

41. Reynolds C, Kontschán J, Takács N, Solymosi N, Sándor AD, Keve G, et al. Shift in the seasonality of ixodid ticks after a warm winter in an urban habitat with notes on morphotypes of *Ixodes ricinus* and data in support of cryptic species within *Ixodes frontalis. Exp Appl Acarol.* (2022) 88:127–38. doi: 10.1007/s10493-022-0 0756-1

42. Pomerantsev BI. *Ixodid Ticks (Ixodidae)*. Herndon: VA: American Institute of Biological Sciences (1959). p. 199.

43. Hoogstraal H. African Ixodoidea. VoI. I. Ticks of the Sudan (with special reference to Equatoria Province and with Preliminary Reviews of the Genera *Boophilus*, *Margaropus*, and *Hyalomma*.) US Navy. (1956) 1:479–90.

44. Wallmenius K, Barboutis C, Fransson T, Jaenson TGT, Lindgren P-E, Nystrom F, et al. Spotted fever rickettsia species in hyalomma and ixodes ticks infesting migratory birds in the European Mediterranean area. *Parasit Vectors*. (2014) 7:318. doi: 10.1186/1756-330 5-7-318

45. Kiros S, Awol N, Tsegaye Y, Hadush B. Hard ticks of camel in southern zone of Tigray, Northern Ethiopia. *J Parasitol Vector Biol.* (2014) 6:151–5. doi: 10.5897/JPVB2014.0162

46. Kerario II, Muleya W, Chenyambuga S, Koski M, Hwang S-G, Simuunza M. Abundance and distribution of Ixodid tick species infesting cattle reared under traditional farming systems in Tanzania. *Afr J Agric Res.* (2017) 12:286–99. doi: 10.5897/AJAR2016.12028