PROTOZOOLOGY - SHORT COMMUNICATION



Hepatozoon in Eurasian red squirrels Sciurus vulgaris, its taxonomic identity, and phylogenetic placement

David Modrý 1,2,3,4 · Lada Hofmannová 4 · Petr Papežík 4 · Karolina Majerová 1,5 · Jan Votýpka 1,5 · Václav Hönig 1,6 · Daniel Růžek 1,6 · Kristýna Hrazdilová 7,8

Received: 24 October 2020 / Accepted: 18 June 2021
© The Author(s), under exclusive licence to Springer-Verlag GmbH Germany, part of Springer Nature 2021

Abstract

Adeleorid apicomplexan parasites of the genus *Hepatozoon* Miller, 1908 are broadly distributed among the rodents. Broader molecular data on *Hepatozoon* from Palaearctic squirrels are necessary for evaluation of diversity and origin of *Hepatozoon* in Eurasian red squirrel *Sciurus vulgaris* populations, considering ongoing invasion by Gray squirrel *S. carolinensis*. Our report brings a set of molecular data from a population of *S. vulgaris* in the Czech Republic, non-invaded by any invasive squirrel species. Cadavers of 41 Eurasian red squirrels were examined using nested PCR targeting 18S rRNA gene; 30 animals tested positive for the presence of *Hepatozoon* spp. DNA in at least one tissue. Phylogenetic analysis of obtained sequence types revealed relatedness to sequences of *Hepatozoon* sp. from *S. vulgaris* from Spain and the Netherlands, forming a sister clade to *Hepatozoon* isolates from other European rodents. The fact that all available 18S rRNA gene sequences form a monophyletic clade is interpreted as a presence of a single *Hepatozoon* species in *S. vulgaris* in continental Europe, most probably *Hepatozoon sciuri*. The presented molecular data on the *Hepatozoon* from European squirrels provides a basis for future studies on possible exchange of *Hepatozoon* species between Eurasian red and gray squirrels.

Keywords Hepatozoon · Squirrel · Sciurus · 18S rRNA gene · Europe · Hemoparasites

Section Editor: Leonhard Schnittger

- ☑ David Modrý david.modry@paru.cas.cz
- ¹ Institute of Parasitology, Biology Centre, Czech Academy of Sciences, Branišovská 31, Ceske Budejovice, Czech Republic
- Department of Botany and Zoology, Faculty of Science, Masaryk University, Brno, Czech Republic
- Department of Veterinary Sciences, Faculty of Agrobiology, Food and Natural Resources/CINeZ, Czech University of Life Sciences, Prague, Czech Republic
- Department of Pathology and Parasitology, Faculty of Veterinary Medicine, University of Veterinary Sciences, Brno, Czech Republic
- Department of Parasitology, Faculty of Science, Charles University, Prague, Czech Republic
- Veterinary Research Institute, Brno, Hudcova 70, 62100 Brno, Czech Republic
- CEITEC, University of Veterinary Sciences, Brno, Czech Republic

Published online: 31 July 2021

Faculty of Medicine in Pilsen, Biomedical Center, Charles University, Plzeň, Czech Republic

Introduction

Adeleorid apicomplexan parasites of the genus Hepatozoon Miller, 1908 are broadly distributed among the terrestrial vertebrates, representing the most diversified and complex assemblage of haemogregarines (Smith 1996). Ubiquitous presence and unique appearance of gametocytes within white blood cells of mammalian hosts make Hepatozoon commonly reported and discussed vector-borne pathogen since the beginning of twentieth century. European squirrel Hepatozoon was originally described as Haemogregarina sciuri Coles, 1914 in mononuclear leucocytes of captive Eurasian red squirrels Sciurus vulgaris Linnaeus, 1758 in England (Coles 1914); later, the same name was used for a haemogregarine reported from Italian S. vulgaris (Franchini 1932). At the North American continent, Hepatozoon griseisciuri Clark, 1958 was described, including its life cycle, from gray squirrels S. carolinensis Gmelin, 1788 from Maryland (Clark 1958). The fact that the original description by Coles (1914) from England mentioned that the examined squirrels were caged with captive gray squirrels S. carolinensis opens a question of possible introduction



of squirrel *Hepatozoon* to British Isles by invasive *S. carolinensis*. Under such scenario, the pathogenicity of *Hepatozoon* observed in the original study as well as in recent accounts from British Isles (Simpson et al. 2006, 2013) can relate to its origin from other host species, similar to fatal infections in European squirrels by squirrel pox virus spread by invasive *S. carolinensis* (Sainsbury et al. 2000). Broader molecular data on *Hepatozoon* from Palaearctic squirrels (both from areas invaded and not invaded by gray squirrels) are necessary for evaluation of diversity and origin of *Hepatozoon* in Eurasian red squirrel populations. Our report brings a set of molecular data from a population of *S. vulgaris* in the Czech Republic, non-invaded by any invasive squirrel species.

Material and methods

Cadavers of 41 Eurasian red squirrels Sciurus vulgaris, which were accidentally killed (mostly road kills) or had died in rescue centers, were examined, originating from more than 30 localities throughout the Czech Republic (Table \$1, Supplementary material). At necropsy, tissues were removed under sterile conditions, and a piece $(5 \times 5 \text{ mm})$ of each tissue was placed into a 2-ml Eppendorf tube, where it was cut into 1-mm pieces; the liver, spleen, and blood were the only tissues used in this study. Blood was obtained from heart of the cadaver with sterile Pasteur pipette, and 50–100 µl of the blood was placed into a 2-ml Eppendorf tube. Blood and organ samples were stored at -20 °C until further analyses. The blood samples did not require any mechanical processing; however, liver and spleen samples were homogenized in RLT buffer (Qiagen) with addition of beta-mercaptoethanol using sterile stainless steel beads (Qiagen) and Tissue Lyzer II as 30% (w/v) suspensions. After addition of 20 µl of proteinase K, the samples were incubated 30 min at 57 °C. The lysate was cleared by centrifugation, and the supernatant collected to a clean sterile microtube. The blood samples were only treated with 20 µl of proteinase K for 30 min at 57 °C. DNA was isolated from 180 µl of the lysate using DNeasy Blood and Tissue kit (Qiagen) according to the manufacturer's instructions. The elution volume was 200 µl. All samples were screened using sensitive nested PCR assay targeting nuclear 18S rRNA gene. For the first round of PCR amplifying ~ 1800 bp, reactions were performed in 15 µl reaction volume, consisting of 1 µl template DNA, 400 nM of each primer (HAM 1F: 5'-GCCAGT AGTCATATGCTTGTC-3′, HPF 2R: 5′- GACTTCTCCTTC GTCTAAG-3' (Criado et al. 2006)), and 7.5 µl of 2×PCRBIO Taq Mix Red (PCR Biosystems, UK). The second round of nested PCR was done in 25 µl using 1 µl of PCR product from the first round as template, 400 nM of each primer (EF-M: 5'-AAAACTGCAAATGGCTCATT-3', ER-M: 5'-CTYGCG CTTACTAGGCATTC-3', modified for this study (Kvičerová et al. 2008), and 12.5 µl of 2×PCRBIO Taq Mix Red (PCR Biosystems, UK). Cycling conditions were identical for both reactions following manufacturer's recommendations with annealing temperature 52 °C and elongation time of 45 s. PCR products were visualized on 1% agarose gel stained by MIDORIGreen Advance (Nippon Genetics Europe, Germany).

All PCR products of expected size were purified from the gel using the Gel/PCR DNA Fragment Extraction Kit (Geneaid Biotech, Taiwan) and directly sequenced by Macrogen capillary sequencing services (Macrogen Europe, the Netherlands) using the amplification primers in both directions. All obtained sequences were carefully edited using Geneious 11.0.2 (Kearse et al. 2012) and compared with those available in GenBank by the BLASTn algorithm (http://blast.ncbi.nlm.nih.gov/Blast.cgi). Unique acquired sequences were deposited in GenBank (Acc. No. MN104636-40). Phylogenetic analysis was conducted on dataset of all 18S rRNA gene sequences available in public databases (available by the end of 2018) for the proposed Bartazoon clade (Karadjian et al. 2015; Maia et al. 2016). In order to reduce the dataset, only sequences longer than 1000 nt and differing > 5 bp from each other were included in the final analyses. All (three) available sequences from Sciurus spp. host were included in analyses. In total, 69 sequences were aligned using MAFFT v7 with the Q-INS-I iterative method applying parameters by default (Katoh et al. 2019); poorly aligned regions were eliminated using the least stringent Gblocks options (Talavera and Castresana 2007) resulting in an alignment of 1570 nt in length. Phylogenetic trees were inferred by maximum likelihood method using IQ-TREE v. 1.6.beta4 (Nguyen et al. 2015). A best-fit evolution model (TVM + F + R3, transversion model AG = CT, using empirical base frequencies and relaxing the assumption of Gamma-distributed rates using 3 categories) was chosen based on the Bayesian information criterion (BIC) computed by ModelFinder (Kalyaanamoorthy et al. 2017). Branch supports were assessed by the ultrafast bootstrap (UFBoot) approximation (Minh et al. 2013) and by the SHlike approximate likelihood ratio test (SH-aLRT) (Guindon and Gascuel 2003). Trees were visualized and edited using FigTree v1.4.1 (http://tree.bio.ed.ac.uk/software/figtree/); final graphic outputs were processed by InkScape v.092.1 (http://www.inkscape.org/). Differences in presence of Hepatozoon spp. DNA in examined tissues were computed using Fisher Exact test in the GraphPad Prism 8 software (GraphPad Software, La Jolla California, USA).

Results and discussion

Of the 30 individuals positive for the presence of *Hepatozoon* spp. DNA, 56 high-quality sequences of 18S rRNA gene were acquired from different tissues (Table S1), making



the infection prevalence 73%; the prevalence was higher in males (80%) than in females (66%). To reveal the most suitable tissue for Hepatozoon spp. detection, only animals with all three tissues available (n=34) were considered for statistical analysis. The prevalence (at least one tissue positive) in this subset was 79.4% (27/34). Blood and spleen tissues (67.6 and 64.7%, respectively) were significantly more positive (p=0.015 and p=0.028, respectively) than liver (35.3%, 12/34).

High-quality sequence of 18S rRNA gene was acquired for 56 samples (Supplementary Table 1); no double peaks suggesting the mix infections were observed in chromatograms. Comparing results of sequencing with GenBank database confirmed 98.59–98.86% (17–20 nucleotide differences) identity with *Hepatozoon* sp. from a squirrel from Spain (Acc. No. EF222259). Five sequence types (with 99.71–99.93% identity) were represented in our dataset. In majority of samples, two dominant sequence types were identified: SV12-7 in 45 samples and SV146-6 in 8 samples differing in one nucleotide. Additionally, three other sequence types (SV52-4, SV174-7, SV194-7) were represented by a single sequence only. These sporadic sequences

differed by a single nucleotide from the two main sequence types as well as from sequences acquired from other tissue of the same individual and were not included in phylogenetic analyses. The biological relevance of this variability needs to be confirmed. Phylogenetic analysis of the two major sequence types from our study confirmed their relatedness to already published sequences of *Hepatozoon* sp. from *Sciurus vulgaris* from Spain and the Netherlands, forming a sister clade to *Hepatozoon* isolates from other European rodents (Fig. 1). A single available sequence from *Sciurus carolinensis* from the USA (Acc. No. JF491240) remained well-separated from all *S. vulgaris* sequences.

PCR detection followed by Sanger sequencing revealed a common presence of the *Hepatozoon* throughout the population of Eurasian red squirrels in the Czech Republic. Similarly, using histopathological examination and detection in blood smears, Simpson et al. (2006) confirmed *H. sciuri* in 37% of the examined red squirrels in the UK, reporting a single partial 18S rRNA gene sequence obtained from a squirrel from Isle of Wight, with 97.9% identity to *H. erhardovae*. Sequence identity within the squirrel and rodent subclades ranged between 98.1 and 99.9%, while between these

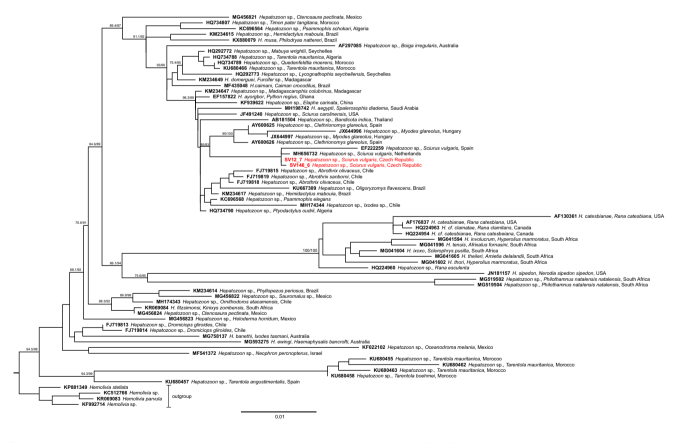


Fig. 1 Phylogenetic analysis assessed by maximum likelihood method based on > 1,000 nt long and differing > 5 bp from each 18S rRNA gene sequences representing genus *Bartazoon* as proposed by Karadjian et al. (2015). Node support values (SH-aLRT/UFB) above

the threshold 80%/95% are displayed. Sequences are marked by their accession numbers, species, host, and country of origin as retrieved from GenBank; newly described sequences from this study are highlighted in red



two clades reached 95.1–98.1%. In our analyses, obtained sequences clustered with other sequences from red squirrel Hepatozoon, forming a clearly monophyletic group within the subclade of European rodent Hepatozoon spp. Interpretation of observed variability is complicated by virtual absence of a species concept in haemogregarine taxonomy. Observed sequence variability fails within variability range of 18S rRNA gene sequences of so far accepted taxa as closely related Hepatozoon erhardovae or more distant Hepatozoon canis. Considering these facts, we believe that observed monophyletic clade of squirrel *Hepatozoon* can be tentatively interpreted as a presence of a single but variable Hepatozoon species in S. vulgaris in continental Europe. Sequencing of more molecular markers including mitochondrial genes as a cytochrome b (Léveillé et al. 2020), morphological data and knowledge of vectors involved as definitive hosts in the life cycle are the only ultimate approach to solve the taxonomy of Eurasian red squirrel Hepatozoon genetic lineages.

At least in two European regions, populations of native Eurasian red squirrels *S. vulgaris* are threatened by invasive Gray squirrel *S. carolinensis*. Latter species is replacing the native squirrel in large part of British Isles from the end of nineteenth century (Lawton et al. 2010) and, more recently, also in northern Italy (Bertolino et al. 2008). In past decades, competition between native and invasive squirrels in the UK involved a squirrel poxvirus, creating a situation classified as disease-mediated invasion (Strauss et al. 2012). The presented molecular data on *Hepatozoon* spp. from European squirrels provides a basis for future studies on possible exchange of *Hepatozoon* species between Eurasian red and gray squirrel in areas of their recent co-occurrence and for understanding of possible impact on native red squirrel populations.

Supplementary Information The online version contains supplementary material available at https://doi.org/10.1007/s00436-021-07229-1.

Acknowledgements We are indebted to all volunteers and staff of rescue centers for enabling access to dead animals.

Funding This research was funded by the Czech Science Foundation (grant number 17-16009S) and by the Ministry of Education, Youth and Sports of the Czech Republic under the project CEITEC 2020 (LQ1601) and by a project No. CZ.02.1.01/0.0/0.0/16_019/0000787.

Declarations

Conflict of interest The authors declare no competing interests.

References

Bertolino S, Lurz PWW, Sanderson R, Rushton SP (2008) Predicting the spread of the American grey squirrel (*Sciurus carolinensis*) in

- Europe: a call for a co-ordinated European approach. Biol Conserv 141:2564–2575. https://doi.org/10.1016/j.biocon.2008.07.
- Clark GM (1958) *Hepatozoon griseisciuri* n. sp.; a new species of *Hepatozoon* from the grey squirrel (*Sciurus carolinensis* Gmelin, 1788), with studies on the life cycle. J Parasitol 44:52–63. https://doi.org/10.2307/3274829
- Coles AC (1914) Blood parasites found in mammals, birds and fishes in England. Parasitology 7:17–61. https://doi.org/10.1017/S0031 182000006284
- Criado A, Martinez J, Buling A et al (2006) New data on epizootiology and genetics of piroplasms based on sequences of small ribosomal subunit and cytochrome b genes. Vet Parasitol 142:238–247. https://doi.org/10.1016/j.vetpar.2006.07.004
- Franchini G (1932) Su di una leucocito-gregarina dello scoiattolo (*Sciurus vulgaris*). Arch Ital Sci Med Colon 13:385–389
- Guindon S, Gascuel O (2003) A simple, fast, and accurate algorithm to estimate large phylogenies by maximum likelihood. Syst Biol 52:696–704. https://doi.org/10.1080/10635150390235520
- Kalyaanamoorthy S, Minh BQ, Wong TKF et al (2017) ModelFinder: fast model selection for accurate phylogenetic estimates. Nat Methods 14:587–589. https://doi.org/10.1038/nmeth.4285
- Karadjian G, Chavatte JM, Landau I (2015) Systematic revision of the adeleid haemogregarines, with creation of *Bartazoon* n. g., reassignment of *Hepatozoon argantis* Garnham, 1954 to *Hemolivia*, and molecular data on *Hemolivia stellata*. Parasite 22. https://doi.org/10.1051/parasite/2015031
- Katoh K, Rozewicki J, Yamada KD (2019) MAFFT online service: multiple sequence alignment, interactive sequence choice and visualization. Brief Bioinform 20:1160–1166. https://doi.org/10. 1093/bib/bbx108
- Kearse M, Moir R, Wilson A et al (2012) Geneious basic: an integrated and extendable desktop software platform for the organization and analysis of sequence data. Bioinformatics 28:1647–1649
- Kvičerová J, Pakandl M, Hypša V (2008) Phylogenetic relationships among *Eimeria* spp. (Apicomplexa, Eimeriidae) infecting rabbits: evolutionary significance of biological and morphological features. Parasitology 135:443–452. https://doi.org/10.1017/S0031 182007004106
- Lawton C, Cowan P, Bertolino S, et al (2010) The consequences of introducing non-indigenous species: two case studies, the grey squirrel in Europe and the brushtail possum in New Zealand. OIE Rev Sci Tech 29:287–298. https://doi.org/10.20506/rst.29.2.1983
- Léveillé AN, El Skhawy N, Barta JR (2020) Multilocus sequencing of *Hepatozoon* cf. *griseisciuri* infections in Ontario eastern gray squirrels (*Sciurus carolinensis*) uncovers two genotypically distinct sympatric parasite species. Parasitol Res 119:713–724. https://doi.org/10.1007/s00436-019-06583-5
- Maia JP, Carranza S, Harris DJ (2016) Comments on the systematic revision of adeleid haemogregarines: are more data needed? J Parasitol 102:549–552. https://doi.org/10.1645/15-930
- Minh BQ, Nguyen MAT, Von Haeseler A (2013) Ultrafast approximation for phylogenetic bootstrap. Mol Biol Evol 30:1188–1195. https://doi.org/10.1093/molbev/mst024
- Nguyen LT, Schmidt HA, Von Haeseler A, Minh BQ (2015) IQ-TREE: a fast and effective stochastic algorithm for estimating maximum-likelihood phylogenies. Mol Biol Evol 32:268–274. https://doi.org/10.1093/molbev/msu300
- Sainsbury AW, Nettleton P, Gilray J, Gurnell J (2000) Grey squirrels have high seroprevalence to a parapoxvirus associated with deaths in red squirrels. Anim Conserv 3:229–233. https://doi.org/ 10.1111/j.1469-1795.2000.tb00107.x
- Simpson VR, Birtles RJ, Bown KJ et al (2006) *Hepatozoon* species infection in wild red squirrels (*Sciurus vulgaris*) on the Isle of Wight. Vet Rec 159:202–205. https://doi.org/10.1136/vr.159.7. 202



- Simpson VR, Hargreaves J, Butler HM et al (2013) Causes of mortality and pathological lesions observed post-mortem in red squirrels (*Sciurus vulgaris*) in Great Britain. BMC Vet Res 9:229
- Smith TG (1996) The genus *Hepatozoon* (Apicomplexa: Adeleina). J Parasitol 82:565–585. https://doi.org/10.2307/3283781
- Strauss A, White A, Boots M (2012) Invading with biological weapons: the importance of disease-mediated invasions. Funct Ecol 26:1249–1261. https://doi.org/10.1111/1365-2435.12011
- Talavera G, Castresana J (2007) Improvement of phylogenies after removing divergent and ambiguously aligned blocks from protein sequence alignments. Syst Biol 56:564–577. https://doi.org/10.1080/10635150701472164

Publisher's note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

