Update on prevalence of *Babesia canis* and *Rickettsia* spp. in adult and juvenile *Dermacentor reticulatus* ticks in the area of Poland (2016–2018)

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Ornate dog tick, *Dermacentor reticulatus* is an important vector of *Babesia canis*, and *Rickettsia* spp. and other pathogens of veterinary and public health interest. The current study is the first to investigate the long-term changes in prevalence of these pathogens in expanding tick populations in Central Europe. Molecular techniques (PCR, sequencing) were applied for the detection of pathogen DNA in adult (n = 2497) and juvenile ticks (1096 larvae and 410 nymphs). DNA of *Rickettsia* spp. was identified in 35% of adults and 12.6% of juvenile ticks. DNA of *B. canis* was detected in 3% of adult ticks and only in ticks from the Eastern region (regional prevalence 6%). As previously, no *B. canis*-positive ticks were found in Western Poland, including ticks from Wrocław area (n = 298). DNA of *B. canis* was identified in 0.33% of juvenile ticks (in 3 pools of larvae and 2 nymphs) from the Eastern region. In the current study we confirmed high occurrence of *R. raoultii* in adults ticks from all four zones and relatively high prevalence of *B. canis* in the Eastern population of *D. reticulatus*, corresponding well with high incidence of canine babesiosis in this area of Poland. Finally, we confirmed *R. raoultii* and *B. canis* infection in all life stages of *D. reticulatus* ticks.

Vector-borne diseases constitute a serious problem of medical and veterinary importance³⁴. Among canine tick-borne diseases, canine babesiosis, caused by apicomplexan parasite *Babesia canis* is of the greatest significance in central and eastern Europe⁵. Ornate dog tick, *Dermacentor reticulatus* is a main if not the only vector of *B. canis*⁴.

Recent study on the distribution of canine babesiosis in Poland in 2018 and occurrence of adult ticks in endemic areas and areas historically free of this tick species (2016–2018) revealed great discrepancy both in distribution of babesiosis cases and tick abundance, with 'hot spot' in Central and Eastern Poland⁵. The present study aimed to investigate the prevalence of *B. canis* in questing adult ticks collected from these regions and to find the link with distribution of babesiosis in Poland. Furthermore, *D. reticulatus* ticks are example of fast-spreading tick species and these dynamic changes in geographical range may be accompanied by changes in prevalence of pathogens in ticks. As we have previously determined prevalence of *B. canis* and *Rickettsia* spp. in ticks collected in years 2012–2014 from different regions of Poland⁶, the present study provides the update on prevalence of these pathogens in previously examined regions/sites and let us compare the long-term changes in infection frequency.

It is worth to underline, that in the previous studies *B. canis* was not detected in adult questing ticks from Western Poland, neither in juvenile ticks collected from rodents⁶–⁸ despite sporadic occurrence of canine babesiosis in this region⁵. In the current study we attempted to collect and examine a significant number of questing *D. reticulatus* ticks from Western Poland, including ticks collected in Wrocław area, in location with confirmed babesiosis cases in dogs.

Juvenile *D. reticulatus* ticks are heavily understudied in comparison to adult ticks, due to their hidden nidicolous style of life¹. One of the aims of our current study was to examine a significant number of larva and nymphs
collected from rodents in the Eastern region of *D. reticulatus* occurrence for the presence of *B. canis* and *Rickettsia* spp. DNA and to compare prevalence of these pathogens between adult and juvenile ticks from the same sites.

Transovarial transmission of *B. canis* in *D. reticulatus* is believed to constitute the main route of transmission in vector population enabling persistence of parasite in certain tick populations. Despite this belief, only few studies have confirmed the occurrence of *B. canis* in larvae and nymphs of ornate dog tick. We have confirmed transovarial transmission of *B. canis* from infected females collected from dogs to eggs and larvae hatched in laboratory conditions in present study we searched for *Babesia* spp. in juvenile ticks obtained from rodents, mainly voles. Detection of *B. canis* DNA in larvae and nymphs feeding on rodents (main reservoir of *B. microti*) would prove the efficiency of vertical transmission of *B. canis*. We hypothesize that an effective transovarial and transstadial transmission of *B. canis* in ticks are the main cause for its persistence in *D. reticulatus* population in *B. canis* hyperendemic areas in Central and Eastern Poland.

**Materials and methods**

**Adult ticks.** In the main study, adult questing ticks were collected in three-year period, 2016–2018, including 864 ticks from 2016, 877 from 2017 and 756 from 2018.

In the main study, 2497 adult ticks (1446 females and 1051 males) were examined for pathogen occurrence: 1264 ticks from the Eastern region of *D. reticulatus* occurrence (1142 from the eastern endemic area and 122 from the eastern expansion zone) and 1233 ticks from the Western region of *D. reticulatus* occurrence (635 from the western endemic area and 598 from the western expansion zone) (zones marked in Fig. 1, following Dwużnik-Szarek et al.).

All procedures of adult *D. reticulatus* collection were described in details in Dwużnik-Szarek et al. Additionally, 298 questing adult *D. reticulatus* (152 females, 146 males) collected in Wrocław area in 2019, located within the city where cases of canine babesiosis were recorded (Kiewra, unpublished), were also examined.

**Larvae and nymphs collected from rodents.** Altogether, 1096 larvae in 150 pools and 410 nymphs processed individually, were examined for *Rickettsia* spp. and *B. canis*. Description of collection of juvenile feeding ticks is provided in Dwużnik-Szarek et al. Briefly, rodents were trapped in habitats preferable for *D. reticulatus* (meadows, fallow lands and wetlands, etc.) in four sites in the Eastern region of *D. reticulatus* occurrence, endemic for *D. reticulatus* (Fig. 1): three sites in Mazovia voivodeship: Stoski + Franciszków (analyzed together as they are situated in close proximity of a few kilometers and represent similar open submerged habitat) and Białobrzegi and one site in Warmia-Mazuria voivodeship—Urwitałt (coordinates in Dwużnik-Szarek et al.). Additionally, data on *Babesia* and *Rickettsia* occurrence in juvenile *D. reticulatus* collected from Białobrzegi and Urwitałt + Talty (these sites were also analyzed together, for their close proximity and habitat similarity [mixed forest]) was included in statistical analyses of prevalence.

**DNA isolation.** Genomic DNA was extracted from ticks (larvae, nymphs, adults) using the Genomic Mini AX Tissue Spin kit (A&A Biotechnology, Gdańsk, Poland) in accordance with the manufacturer’s protocol.

Adult *D. reticulatus* collected from the environment were processed individually. SPEX SamplePrep Freezer/Mill 6875D (Laboratory Equipment for Sample Preparation & Handling, Rickmansworth, Great Britain) was used to prepare the tick tissue homogenate for the subsequent extraction. This equipment enables the complete homogenization of 24 samples in the temperature of liquid nitrogen (~195.6°F) within 2 min. During one cycle, tick was placed in cryogenic vial and frozen in liquid nitrogen. Then it was pulverized into a homogenate using to prepare the tick tissue homogenate for the subsequent extraction. This equipment enables the complete homogenization of 24 samples in the temperature of liquid nitrogen (~195.6°F) within 2 min. During one cycle, tick was placed in cryogenic vial and frozen in liquid nitrogen. Then it was pulverized into a homogenate using a Laboratory Equipment for Sample Preparation & Handling, Rickmansworth, Great Britain) was used to prepare the tick tissue homogenate for the subsequent extraction. This equipment enables the complete homogenization of 24 samples in the temperature of liquid nitrogen (~195.6°F) within 2 min. During one cycle, tick was placed in cryogenic vial and frozen in liquid nitrogen. Then it was pulverized into a homogenate using a homogenizer (Nippon Genetics Europe, Düren, Germany). From each rodent a maximum of 50 larvae (5 pools, up to 10 larvae in one pool) and 5 nymphs (processed individually) were tested.

**Pathogen detection.** For detection of *Rickettsia* spp., primers Cs409 (5′-CCTATGGCTATTATGCTTTGC-3′) and Rp1258 (5′-ATTGCAAAAAAGTACAGTGAAACG-3′) were used for amplification of a 750 bp fragment of the citrate synthase (*gltA*) gene as follows: initial denaturation in 95°C for 5 min, 40 cycles of denaturation at a temperature of 95°C for 30 s, 45 s of primer annealing in 59°C, and elongation in 65°C for 1 min.
For molecular screening of *Babesia* spp., primers CRYPTO R (5'-GCT TGA TCC TTC TGC AGG TTC ACC TAC-3') and CRYPTO F (5'-AAC CTG GTT GAT CCT GCC AGT-3') were used to amplify ~1200 bp fragment of 18S rDNA in the first step of nested-PCR reaction. In second reaction primers BabGR2 (5'-CCA AAG ACT TTG ATT TCT CTC-3') and BabGF2 (5'-GYY TTG TAA TTG GAA TGA TGG-3') amplified ~550 bp fragment of 18S rDNA20. Reaction conditions were as described in Tołkacz et al.21. For species-specific detection of *B. canis*, primers BcCOX1nR (5'-GGC CCT GTT CGG TAT TGC AT-3') and BcCOX1nF (5'-CCA TTT TGT TCT TTC AAT TGG TGC -3') were used to amplify ~328 bp fragment of mitochondrial cytochrome c oxidase subunit 1 (cox1) gene22. Reaction conditions were as follows: 94 °C for 5 min, followed by 40 cycles at 94 °C for 20 s, 59 °C for 30 s, 72 °C for 45 s and final elongation at 72 °C for 7 minutes22. DNA of *B. canis* was used as positive control, negative controls were performed with 2 μl of sterile water in the absence of template DNA. PCR products were visualized on 1.5% agarose gel stained with Midori Green Stain (Nippon Genetics Europe, Düren, Germany).

Selected PCR products were sequenced by a private company (Genomed, Warsaw, Poland). Sequence alignments were carried out using BLAST-NCBI. Molecular phylogenetic analyses were performed in Molecular Evolutionary Genetics Analysis (MEGA) X open access software (https://www.megasoftware.net/) using Maximum Likelihood method of tree-construction. The evolutionary model was chosen with accordance to the data (following implemented model test in MEGA X) and bootstrapped over 1000 randomly generated sample trees.

**Figure 1.** Map of endemic areas and expansion zones for adult questing *Dermacentor reticulatus* ticks (Dwużnik-Szarek et al. 2021) and sites of the collection of juvenile ticks. The map was designed using ArcGIS (ESRI) version 10.8.1 software (institutional licence purchased by the University of Warsaw, Warsaw, Poland). Briefly, each georeferenced location of tick collection (listed in Dwużnik-Szarek et al.) was projected as a point type .shp layer and then used as the raw data for spatial analyses. A radius buffer was calculated for each point, allowing to interpolate a range of occurrence of the tick. The base layer consisted of contour map of Poland: country borders and largest administrative units (voivodeships).
Table 1. Prevalence of *B. canis* and *R. raoultii* in *D. reticulatus* ticks in four zones. Nc not calculated.

| Region (no. of ticks) | Babesia canis | 95%CI | Rickettsia raoultii | 95%CI | Zones | Babesia canis | 95%CI | Rickettsia raoultii | 95%CI |
|-----------------------|---------------|-------|---------------------|-------|-------|---------------|-------|---------------------|-------|
| Eastern (1264)        | 5.9% (74/1264) | 4.7–7.3 | 35.9% (454/1246)    | 33.3–38.6 | Eastern endemic zone (1142) | 6.1% (70/1142) | 4.9–7.6 | 35.7% (408/1142)    | 33.0–38.5 |
|                       |               |       |                     |       | Eastern expansion zone (122) | 3.3% (4/122) | 1.1–7.6 | 37.7% (46/122)     | 29.5–46.5 |
| Western (1233)        | 0             | nc    | 33.5% (413/1233)    | 30.9–36.2 | Western endemic zone (635) | 0 | nc | 30.7% (195/635)    | 27.2–34.4 |
|                       |               |       |                     |       | Western expansion zone (598) | 0 | nc | 36.5% (218/598)    | 32.7–40.4 |

**Statistical analysis.** Minimum Infection Rate (MIR) was calculated for pools of larvae; if a sample was positive it was assumed that only one tick specimen in the pool was infected. Analyses regarding larvae were conducted on MIR.

For the analysis of prevalence (% PCR-positive ticks), we applied maximum likelihood techniques based on log linear analysis of contingency tables in the IBM SPSS Statistics: PS IMAGO PRO Academic v.7 (institutional license purchased by the University of Warsaw, Warsaw, Poland).

For adult ticks factors such as tick sex (two levels: male, female), region of *D. reticulatus* occurrence (two levels: the Western region, the Eastern region), zones (four levels: western endemic zone, western expansion zone, eastern endemic zone, western expansion zone), season (two levels: spring, autumn), year (three levels: 2016, 2017, 2018) were used in models with the presence or absence of pathogen DNA (*B. canis*, *Rickettsia* spp.) considered as a binary factor (0, 1). In case of juvenile *D. reticulatus* ticks, sites (three levels: Białobrzegi, Franciszków + Stoski, Urwitalt + Talty) and tick stage (larva or nymph) were used in models with the presence or absence of pathogen DNA (*B. canis*, *Rickettsia* spp.) considered as a binary factor (0, 1).

For each level of analysis in turn, beginning with the most complex model, involving all possible main effects and interactions, those combinations that did not contribute significantly to explaining variation in the data were eliminated in a stepwise fashion beginning with the highest level interaction (backward selection procedure). A minimum sufficient model was then obtained, for which the likelihood ratio of $\chi^2$ was not significant, indicating that the model was sufficient in explaining the data. Values of $P < 0.05$ were considered as significant.

**Ethics approval.** All of the procedures (trapping and handling of free-living rodents) were conducted with the approval of the First Warsaw Local Ethics Committee for Animal Experimentation in Poland (ethical license number: 706/2015), according to the principles governing experimental conditions and care of laboratory animals required by the European Union and the Polish Law on Animal Protection. Following collection of ticks, animals were released at the point of capture.

**Results**

**Adult questing Dermacentor reticulatus.** In total, 36.1% of examined ticks were infected with at least one pathogen. In the Western region of *D. reticulatus* occurrence (western endemic area + western expansion zone) 33.3% of ticks were infected, in the Eastern region (eastern endemic area + eastern expansion zone) 38.7% were infected (pathogen presence/absence × region of *D. reticulatus* occurrence: $\chi^2_1 = 7.30, P = 0.007$).

*Babesia canis* was detected in adult ticks with total prevalence of 3.0% (74/2497) and as we suspected, DNA of *B. canis* was detected only in ticks from the Eastern region of *D. reticulatus* occurrence (Table 1, NS). No *B. canis*-positive ticks were found among 298 ticks collected from Wrocław area.

Slight difference in prevalence of *B. canis* was also observed between two eastern zones with about 6% of infected ticks in the eastern endemic area in comparison to about 3% in eastern expansion zone (not significant; NS) (Table 1).

Additionally, higher percentage of *B. canis*-positive ticks was noted in spring than in autumn (6.8% [95 CI]: 5.1–8.8%) vs 4.7% [95 CI]: 3.2–6.7%), respectively) (*Babesia* presence/absence × season: $\chi^2_1 = 23.60, P < 0.001$). The highest prevalence of *B. canis* was noted in 2018 (8.5% [5.9–11.9%]), followed by 2017 (5.6% [3.9–7.9%]) and 2016 (4.0% [2.4–6.2%]) (*Babesia* presence/absence × year of tick collection: $\chi^2_2 = 6.99, P = 0.03$). Tick sex had no effect on prevalence of *B. canis* (NS).

DNA of *Rickettsia* spp. was detected in 34.7% of total ticks. Prevalence of *Rickettsia* spp. was similar in the Eastern and Western regions of *D. reticulatus* occurrence (Table 1, NS). There were some minor differences in percentage of PCR-positive ticks between four zones. Higher prevalence of *Rickettsia* was detected in ticks collected in the both expansion zones in comparison to the endemic regions (Table 1, NS). Season of tick collection and tick sex had no effect on prevalence of *Rickettsia* (NS). Interestingly, we detected significant differences in prevalence of this pathogen between years of tick collection. The highest prevalence, 39.4% [36.1–42.6%] was noticed in 2016 followed by 35.8% [32.7–39.0%] in 2017 and 28.2% [25.1–31.5%] in 2018 (*Rickettsia* presence/absence × year: $\chi^2_1 = 23.26, P < 0.001$).

Co-infections of *Babesia* and *Rickettsia* could be recorded only in ticks collected in the Eastern region of the *D. reticulatus* occurrence, and only 3.2% (40/1264) of ticks from this region carried two pathogens (co-infection × region of *D. reticulatus* occurrence: $\chi^2_1 = 59.15, P < 0.0001$).
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Table 2. Comparison of B. canis prevalence between juvenile and questing adult ticks D. reticulatus in endemic sites. *Stoski + Franciszków together for juvenile tick stages. **Urwitałt + Talty together for juvenile tick stages.

| Site                  | Larvae   | 95% CI | Nymphs  | 95% CI | Adults | 95% CI |
|-----------------------|----------|--------|---------|--------|--------|--------|
| Białobrzegi           | 0.35% (3/859) | 0.1–0.9 | 0.81% (1/124) | 0.09–3.7 | 1.85% (1/54) | 0.2–8.3 |
| Stoski*               | 0% (0/52) | 0      | 0% (0/65) | 0      | 3.28% (8/244) | 1.6–6.1 |
| Urwitałt**            | 0% (0/185) | 0      | 0.45% (1/221) | 0.05–2.1 | 5.56% (8/144) | 2.7–10.2 |
| Total                 | 0.27% (3/1096) | 0.08–0.7 | 0.49% (2/410) | 0.10–0.53 | 3.85% (17/442) | 2.3–5.6 |

Juvenile Dermacentor reticulatus. Total prevalence of pathogens in juvenile D. reticulatus, including MIR, was 12.6% [11.1–14.3%]. Among instars, 8.7% [7.1–10.3%] of larvae and 23.4% [19.5–27.6%] of nymphs were positive for at least one pathogen (pathogen presence/absence × tick stage: $\chi^2_1 = 53.04$, P < 0.001).

Total prevalence by site (L + N) was the highest in Urwitałt + Talty, Masuria—19.2% [15.6–23.3%], followed by Stoski + Franciszków in Mazovia—16.2% [10.4–23.7%], with the lowest value in Białobrzegi (Mazovia)—9.6% [7.8–11.5%] (pathogen presence/absence × site: $\chi^2_1 = 35.49$, P < 0.001).

Rickettsia spp.. 12.6% [11.0–14.3%] of examined juvenile D. reticulatus ticks were Rickettsia-positive. Prevalence of Rickettsia was more than twice higher in nymphs compared to larvae (22.9% [19.1–27.2%] vs 8.7% [MIR] [7.1–10.4%], respectively) (Rickettsia present/absence × tick stage: $\chi^2_1 = 50.1$, P < 0.001).

Among sites, the highest prevalence of this pathogen was detected in juvenile ticks in Urwitałt + Talty, 19.0% [15.4–23.0%], followed by Stoski + Franciszków, 16.2% [10.4–23.2%] and Białobrzegi, 9.5% [7.8–11.4%] (Rickettsia presence/absence × site: $\chi^2_1 = 24.38$, P < 0.001).

Babesia canis. DNA of B. canis was identified in 0.33% [0.13–0.73%] of instars, with 0.27% [0.08–0.73%] in larvae (3 positive pools collected from Microtus oeconomus, Myodes glareolus and Apodemus agrarius in Białobrzegi) and in 0.49% [0.10–0.53%] of nymphs (one positive nymph collected from M. oeconomus in Białobrzegi, one nymph removed from M. glareolus, Urwitałt) (NS). DNA of B. canis was detected in juvenile ticks from Białobrzegi and Urwitałt + Talty but not in much lower number of instars originated from Stoski and Franciszków (NS) (Table 2).

Comparison of B. canis prevalence between juvenile ticks and questing adult ticks collected from endemic areas. 442 adult ticks from three endemic sites, Urwitałt, Białobrzegi and Stoski were examined for B. canis presence. DNA of B. canis was detected in all tick stages (in larvae, nymph and adult ticks) in Białobrzegi site (Table 2). Although there were some differences in prevalence between sites, they were insignificant (NS; Table 2).

Molecular identification of pathogen species/genotypes. Rickettsia spp.. 42 of 867 Rickettsia-positive PCR products were sequenced. Among these, 38 were obtained from adult questing ticks (14 originated from eastern endemic area, four from eastern expansion zone, nine from western endemic area and 11 from western expansion zone). Four sequences were derived from juvenile D. reticulatus ticks: three sequences were obtained from larvae (Stoski) and one from a nymph (Białobrzegi). All obtained sequences displayed the highest identity (99.83–100%) to R. raoultii (GenBank accession numbers: MN388798 and MN550896). The phylogenetic tree, incorporating 25 sequences obtained in this study and 22 reference sequences from GenBank, is presented in Fig. 2. The tree topology showed that sequences obtained from examined ticks clustered on the one separate branch, as expected from BLAST analysis, constituting the R. raoultii clade.

Babesia spp. Among 74 PCR products from adult D. reticulatus ticks, 31 Babesia-positive samples were sequenced. Sequences representing cox1 gene (n = 26) showed high similarity (in range 99.7–100%) to the sequence of B. canis derived from red fox from Poland (MN147867) and B. canis derived from a dog, USA (KC207822). Three sequences were derived from ticks collected in the eastern expansion zone and 22 from the eastern endemic region of D. reticulatus occurrence, Masovian voivodeship and one from Urwitałt, Warmia-Mazuria voivodeship. A representative tree for cox1 sequences (ten sequences derived from this study and 11 reference sequences from GenBank), obtained using the Maximum Likelihood method and Hasegawa-Kishino-Yano model is presented in Fig. 3. Our sequences (GenBank accession numbers OL549270- OL549279) clustered on one separate branch with the other two B. canis cox1 gene sequences deposited in GenBank.

Five PCR products of Babesia detected in adult ticks, representing 550 bp-fragment of 18S rDNA, were sequenced successfully. Representative sequence was deposited in GenBank under accession number MZ363934 and showed 99.8% identity (488/489) to B. canis derived from red fox (Poland), domestic dog (China) and D. reticulatus tick (Kazakhstan) (GenBank accession numbers MN134074, MK571831 and MK070118, respectively).

Additionally, two of three PCR products of Babesia-positive larvae (cox1 fragment) were sequenced successfully. Both samples were essentially identical (identity above 99%) to the sequence of B. canis, MN147867, from red fox, Poland.
**Figure 2.** Molecular phylogenetic analysis of a 750 bp fragment of the citrate synthase (gltA) gene of *Rickettsia raoultii*. The evolutionary history was inferred by using the Maximum Likelihood method and Hasegawa-Kishino-Yano model. The tree with the highest log likelihood (−1924.77) is shown. The percentage of trees in which the associated taxa clustered together is shown next to the branches. Initial tree(s) for the heuristic search were obtained automatically by applying Neighbor-Join and BioNJ algorithms to a matrix of pairwise distances estimated using the Maximum Composite Likelihood (MCL) approach, and then selecting the topology with superior log likelihood value. A discrete Gamma distribution was used to model evolutionary rate differences among sites (5 categories (+G, parameter = 0.3308)). This analysis involved 58 nucleotide sequences. There were a total of 703 positions in the final dataset.

**Discussion**

Our current study allowed to monitor long-term dynamic of prevalence of two main pathogens vectored by *D. reticulatus* in endemic areas and zones of expansion in Central Europe. The main finding is stable great difference in prevalence of *B. canis* between the Western and Eastern populations of *D. reticulatus*. Higher prevalence of *B. canis* was also recorded in both endemic areas in comparison to both expansion zones. Additionally, there was the association between the occurrence of *B. canis* in adult questing ticks and juvenile stages in the eastern
cases (n = 1532) and incidence of babesiosis (53/1000 dogs), noted in that region in 2018, and accompanied by experience in the capital city Warsaw over the period of six years is also reflected in high number of canine babesiosis cases in this locality5 and is not confirmed by other studies on results require more investigations, as this high prevalence is not reflected in the number/incidence of babesiosis cases/1000 dogs in 2018, thus confirming maintenance of hyperendemic region for canine babesiosis5,13,23. It shows, that the prevalence is relatively high fatality rate of 2.5%5. Furthermore, in some localities in this region, incidence reached up to 250 stable high prevalence of Babesia canis, 8% prevalence in the Masovian endemic zone and almost 5% prevalence in the eastern expansion zone, west of the Vistula River. This finding supports successful maintenance/circulation of Babesia canis in the Eastern D. reticulatus population. Stable high prevalence of Babesia canis observed in central and eastern Poland (including capital city Warsaw) over the period of six years is also reflected in high number of canine babesiosis cases (n = 1532) and incidence of babesiosis (53/1000 dogs), noted in that region in 2018, and accompanied by relatively high fatality rate of 2.5%5. Furthermore, in some localities in this region, incidence reached up to 250 cases/1000 dogs in 2018, thus confirming maintenance of hyperendemic region for canine babesiosis5,13,23. It is worth to underline, that in the current study the highest prevalence of Babesia canis was observed in Urwiltait, in Warmia-Mazuria voivodeship, one of the oldest area known as endemic for D. reticulatus5,24,25. Babesia canis was identified in numerous recent studies on adult D. reticulatus collected in Eastern and NE Poland with prevalence in range 0.6–7.3%26–29.

In agreement with our previous study, DNA of Babesia canis wasn’t detected in ticks collected from Western Poland, neither in current nor in other studies8,7. We haven’t found Babesia canis DNA in ticks collected from Wroclaw area, despite occurrence of canine babesiosis in this city4. However, the annual number and incidence of canine babesiosis cases in several veterinary clinics from the area of Western Poland were extremely low in 2018- a total 19 cases/year and 0.4/1000 dogs, respectively, in comparison with central and eastern Poland and suggest very low local prevalence in ticks (< 0.1%) or very focal occurrence of infected D. reticulatus ticks. Focal occurrence of Babesia canis-infected ticks was reported earlier in Switzerland30 and the UK31.

Recently, high percentage of Babesia canis-positive Ixodes ricinus ticks was detected near Poznań32. However, these results require more investigations, as this high prevalence is not reflected in the number/incidence of babesiosis cases in this locality4 and is not confirmed by other studies on I. ricinus ticks8.

Rickettsia raoultii was the most common pathogen in both adult and juvenile ticks. Bacteria from this genus are known as intracellular endosymbionts of various invertebrates, including ticks43. Prevalence of R. raoultii in adult ticks in the current study was higher in comparison to data from Austria (4.5%)44, Romania (18%)32, Slovakia (22.3–27%)34 and Ukraine (28%)36. Prevalence was lower than prevalence of Rickettsia observed four years earlier, during our previous study (44.1%)28 and in other studies in Poland (in range 41–44%)26,28. In our previous study significant differences in prevalence of Rickettsia between western and eastern populations were observed4. Prevalence of Rickettsia was by 10% higher in the Western population of D. reticulatus than in the Eastern one. In the present study prevalence was similar across four zones and two regions. However, in previous study (2012–2014) great majority of ticks (n = 1993) originated from the Eastern region of D. reticulatus occurrence and only 592 from the Western one, but in the current study (2016–2018) the number of examined ticks from both regions was similar (1264 vs 1233). Interestingly, although DNA of R. raoultii was identified in larvae and nymphs of D. reticulatus, the pattern was typical for tick-borne pathogens acquired externally (by feeding) and transmitted transstadially, with feeding prevalence from larvae, through nymphs to adults.

Figure 3. Molecular phylogenetic analysis of cox1 of Babesia spp. (328 bp). The evolutionary history was inferred by using the Maximum Likelihood method and Hasegawa-Kishino-Yano model. The tree with the highest log likelihood (−1413.72) is shown. The percentage of trees in which the associated taxa clustered together is shown next to the branches. Initial tree(s) for the heuristic search were obtained automatically by applying Neighbor-Join and BioNJ algorithms to a matrix of pairwise distances estimated using the Maximum Composite Likelihood (MCL) approach, and then selecting the topology with superior log likelihood value. The tree is drawn to scale, with branch lengths measured in the number of substitutions per site. This analysis involved 21 nucleotide sequences. There were a total of 236 positions in the final dataset.
Despite high prevalence of these bacteria in *D. reticulatus* population, the impact of *R. raoultii* on tick fitness or feeding process has yet to be elucidated\(^1\)\(^2\). For humans many *Rickettsia* species, including *R. raoultii*, are considered as pathogenic\(^3\). Prevalence of rickettsioses determined by indirect immunofluorescent assay (IFA) in Poland in years 2006–2012 reached 2.7% \(^4\). In North-Eastern Poland, where both *I. ricinus* and *D. reticulatus* ticks are abundant, presence of anti-*Rickettsia* IgG antibodies was confirmed in 51% of foresters and 27% of farmers\(^5\). Although registration of rickettsioses cases is obligatory in Poland, less than half of detected cases are likely reported\(^6\).

In current study, we made the effort to sample both adult and juvenile ticks from the sites endemic for *D. reticulatus* and *B. canis*. *Babesia canis* was identified in juvenile ticks, three larval pools and two nymphs, from one of this endemic sites with prevalence below 1%, while prevalence in adult ticks ranged from 2 to 5.6%. In Białobrzegi and Stoski sites only recently classified as endemic\(^5\), prevalence was much lower than in old endemic sites. But in Białobrzegi, *B. canis* was detected in every tick stage. In agreement with our results, Dunaj et al.\(^8\) reported *B. canis* infection in nine nymphs of *D. reticulatus* from the eastern endemic region. Detection of DNA of *B. canis* in partially engorged small specimens might have been negatively affected by the presence of other babesiae (i.e. *B. microti*) or other parasites (i.e. *Hepatozoon*) in blood meal taken by instars\(^9\). Identification of *B. canis* in larvae and nymphs collected from rodents supports the predicted routes of circulation of this piroplasm in endemic *D. reticulatus* population through transovarial and transstadial transmission. The main animal reservoir of *B. canis* in Poland constitute domestic dogs, as these parasites were rarely recorded in free-living canids, red foxes and grey wolves\(^10\)\(^11\)\(^12\)\(^13\)\(^14\)\(^15\).

**Conclusions**

In the present study we determined the prevalence of *B. canis* and *Rickettsia* spp. in questing adult *D. reticulatus* ticks collected from different regions in Poland. Furthermore, we confirmed high occurrence of *R. raoultii* in adult ticks from all four zones and relatively high prevalence of *B. canis* in the Eastern population of *D. reticulatus*, corresponding well with high incidence of canine babesiosis in this area of Poland. Interestingly, no *B. canis*-positive ticks were found again in Western Poland, including Wrocław area. Finally, we confirmed *R. raoultii* and *B. canis* infection in all life stages of *D. reticulatus* ticks.

Received: 21 December 2021; Accepted: 23 March 2022

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Acknowledgements

The study was funded by the National Science Centre (NCN) Sonata Bis grant no. 2014/14/E/NZ7/00153 (AB).

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Competing interests

The authors declare no competing interests.

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