Comparative population genetic structure of two Ixodidae ticks (*Ixodes ovatus* and *Haemaphysalis flava*) in Niigata Prefecture, Japan

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Abstract

Background

Ixodid tick species such as *Ixodes ovatus* and *Haemaphysalis flava* function as important vectors of tick-borne diseases in Japan. The study of the genetic patterns of tick populations can reveal information regarding the spread of tick-borne disease. We hypothesized that *I. ovatus* and *H. flava* have different population genetic structure because of their host mobility in different tick life stages despite sharing of hosts.

Methods

Samples (*n* = 1 to 77) were collected in 29 (*I. ovatus*) and 17 (*H. flava*) sampling locations across Niigata. In this study, we used genetic structure at two mitochondrial loci (*cox1*, 16S rRNA gene) to infer gene flow patterns of *I. ovatus* and *H. flava* from Niigata Prefecture, Japan.

Results

For *I. ovatus*, pairwise *F*<sub>ST</sub> and analysis of molecular variance (AMOVA) analyses of *cox1* sequences indicated significant among-population differentiation. This was in contrast to *H. flava*, for which there were only two cases of significant pairwise differentiation and no overall structure. A Mantel test revealed isolation by distance and there was positive spatial autocorrelation of haplotypes in *I. ovatus* *cox1* and 16S sequences, but non-significant results were observed in *H. flava* in both markers. We found three genetic groups (China 1, China 2 and Japan) in the *cox1* *I. ovatus* tree. Newly sampled *I. ovatus* grouped together with a published *I. ovatus* sequence from northern Japan and were distinct from two other *I. ovatus* groups that were reported from southern China.
Conclusions

The three genetic groups in our data set suggest the potential for cryptic species within the lineage. While many factors can potentially account for the observed differences in genetic structure, including population persistence and large-scale patterns of range expansion, we propose that differences in the mobility of hosts of tick immature stages (small mammals in *I. ovatus*; birds in *H. flava*) may be driving the observed patterns.

Background

Tick-borne diseases are a public health concern and their control is often challenging due to the fact that it involves a complex transmission chain of vertebrate hosts and ticks interacting in a changing environment [1]. In order to understand the vector-host relationship, studies on the spatial distribution and population genetic structure of ticks are needed. Population genetic studies have helped in understanding the dispersal patterns and potential pathogen transmission in ticks [2]. The direction, distance and gene flow patterns between tick populations can be inferred from population genetic studies. For example, if high gene flow is observed there might be a greater chance of colonizing new areas or of re-colonizing areas following successful vector control programs [3]. Therefore, estimates of genetic structure and gene flow in tick populations may give insights regarding the spread of tick-borne disease.

Host mobility can affect the genetic patterns of tick populations, although its effects are not consistent. Several studies have reported low levels of gene flow in ticks with less mobile hosts (e.g. smaller mammals) and high levels of gene flow in ticks with highly mobile hosts [2]. For example, *Amblyomma americanum* and *A. triste*
(Ixodidae, Acari) exhibited high gene flow across various spatial scales (137, 000 km$^2$ to 2.78 million km$^2$) and this was attributed to their hosts’ dispersal capabilities (large mammals and birds) [4-6], while low gene flow was observed in Amblyomma dissimile populations and this was attributed to the low mobility of its hosts (small mammals, reptiles and salamanders) [7]. In contrast, some studies have reported low gene flow despite the high mobility of the host, for example in Ixodes scapularis [8] and Ornithodoros coriaceus [9] which can be attributed to the host mobility.

In Japan, tick-borne diseases are an increasing public health concern, affecting humans and animals [10]. A total of 8 genera of ticks have been recorded in Japan, composed of 47 species: 43 Ixodidae species and 4 Argasidae species [11]. Out of 47 species, 21 species are known to parasitize humans [12]. Among these 21 species, Ixodes ovatus, the main vector of Lyme disease [13] and Haemaphysalis flava, a vector of severe fever with thrombocytopenia syndrome (SFTS) and Japanese spotted fever (JSF), were reported in Japan [10,14]. A previous study found that the hosts of adult I. ovatus were mainly hares and can also be large mammals (e.g. cows and horses), and the hosts of immature forms were small rodents [15]. H. flava adult host preference are cows, dogs, horses, wild boar, bear, and deer, while immature forms only parasitized birds [15]. Despite the similarity of I. ovatus and H. flava in which blood meal from different vertebrate host is taken in each life stage (larva, nymph and adult) [16], comparative population genetic studies on two Ixodid tick species such as I. ovatus and H. flava from the same sampling location remain nonexistent. Thus, our study is important because there are no previous studies about dispersal movement of two important tick species: I. ovatus and H. flava, we conclude that low gene flow might be observed in I. ovatus due to the lower mobility of hares in the adult stage and small rodents in its immature stage while H. flava
will exhibit high gene flow because of its avian mediated dispersal in its immature stage.

In addition to tick population structure studies, genetic studies may also reveal the presence of cryptic species, where morphologically identified individuals might represent more than one species [17]. Previous studies on *Rhipecephalus appendiculatus* [18], *I. holocyclus* [19] and *I. ovatus* [20] have revealed the presence of cryptic species based on the clustering of haplotypes in a phylogenetic tree, and concluded that morphological criteria for species differentiation alone are equivocal and that genetic analysis is important. A putative *I. ovatus* species complex was identified based on the presence of 3 distinct haplotype clusters in both *cox1* and 16S rRNA genes; 2 groups in China and one in Japan that included haplotypes from North America [20].

Here, we studied the population genetic structure of *I. ovatus* and *H. flava* and also tested for the presence of cryptic species using mitochondrial DNA sequences of *cox1* and the 16S rRNA gene. We hypothesized that *I. ovatus* and *H. flava* may have differences in their population genetic structure despite some overlap in their adult host preference because of the differences in mobility of the immature tick hosts. We predicted that *H. flava* exhibits high gene flow patterns due to avian-mediated dispersal of immature ticks, in contrast to *I. ovatus* which uses small mammals as hosts during its immature stages.

**Methods**

**Study site, collection, sampling and identification**

From April 2016 until November 2017, ticks were collected by the standard flagging methods [21] across Niigata Prefecture in Japan and a total of 29 sampling locations
were surveyed for *H. flava* and *I. ovatus* ticks (Additional File 1. Table S1). Altitude per location ranged from 8 -1402 m/a.s.l. and the geographic distance between sites ranged from 4.28 - 247.65 km. Collected ticks were stored in microcentrifuge tubes with ethanol at 4°C. A total of 2,103 individual tick samples were collected.

Identification of sex, developmental stage and species was performed using a compound microscope and identification keys of [15].

**DNA extraction, PCR amplification and sequencing**

Genomic DNA (*I. ovatus* n=320; *H. flava* n=220) from each identified tick was extracted using the Isogenome DNA extraction kits (NIPPON GENE Co. Ltd., Tokyo, Japan) following the manufacturer’s recommended protocol. Prior to DNA extraction, each tick was washed with alcohol and PBS solution. DNA quality was checked using a NanoDrop™ 2000 Spectrophotometer (Thermo Scientific™). Two mitochondrial genes were amplified by polymerase chain reaction (PCR): Cytochrome Oxidase 1 (*cox1*) (658 base pairs) using the primer pairs LCO-1490 (5'-GGTCAACAAATCATAAAGATATTGG-3') and HCO1-2198 (5'-AAACTTCAGGGTGACCAAAAAATCA-3') [22] and the 16S rRNA gene (16S) (407 base pairs) with the following primer pairs 16S+1 (5'CTGCTCAATGAATTTAAATTGC-3') and 16S – 1 (5' -CGGTCTAAACTCAGATCATGTAGG-3') [23]. All PCR amplifications of both *cox1* and 16S were performed with a final volume of 10 µl with 1µl of genomic DNA. The PCR reaction for both markers were composed of the following: 10x Ex Taq buffer, 25mM MgCl₂, 2.5mM dNTP, 10µm of forward and reverse primers and 5 units/µl of TaKaRa Ex Taq™ (Takara Bio Inc.). The *cox1* PCR amplification were as follows: an initial denaturation of 94°C for 2 minutes, denaturation of 94°C for 30 seconds, annealing of 38°C for 30 seconds, extension of 72°C for 1 minute for 30 cycles and
final extension of 72°C for 10 minutes. While the 16S PCR amplification followed the protocol of [23] with some modifications, initial denaturation of 94°C for 3 minutes, denaturation of 94°C for 30 seconds, annealing of 50°C for 40 seconds, extension of 72°C for 40 seconds for 30 cycles and final extension of 72°C for 5 minutes. PCR products were purified using the QIAquick 96 PCR Purification Kit (Qiagen) in accordance with the manufacturer’s instructions, and sequenced by Eurofin Genomics, Inc., Tokyo, Japan.

**Sequence data analysis**

We assembled forward and reverse reads for each individual using CodonCode Aligner ver 1.2.4 software (https://www.codoncode.com/aligner/). No ambiguous bases were observed and the low quality bases were removed in the start and end of the reads. Multiple alignment of sequences was performed using the MAFFT alignment online program with default settings (https://mafft.cbrc.jp/alignment/server/). To ensure sequence quality and correct species identification, we checked the similarity of the sequences against reference sequences from GenBank using BLASTN (Basic Local Alignment Search Tool - Nucleotide, https://blast.ncbi.nlm.nih.gov/Blast.cgi?PROGRAM=blastn&PAGE_TYPE=BlastSearch&LINK_LOC=blasthome). The quality of the final aligned sequences was checked in Mesquite ver 3.5 [24] wherein the protein coding genes were translated into amino acids to confirm the absence of stop codons.

**Population genetic analysis**

We analyzed the final sequences of both *H. flava* and *I. ovatus* per genetic marker separately using DNASp ver 6.12.03 [25] and Arlequin ver 3.5.2.2 [26] for the
following parameters: number of haplotypes, average number of polymorphic sites, average number of nucleotide differences, gene/haplotype diversity and nucleotide diversity. The population genetic structure among and within sampling locations in Niigata Prefecture was calculated by analysis of molecular variance (AMOVA) performed in Arlequin with 1023 permutations. The degree of molecular genetic differentiation between sampling locations defined as populations was assessed by calculation of the pairwise $F_{ST}$ values in Arlequin.

To determine if the genetic differentiation was influenced by geographical distance or altitudinal differences among populations, we performed Mantel Test in GenAlEx ver 6.51b2 [27]. Two tests per species and marker were conducted. First we compared pairwise genetic (pairwise $F_{ST}$ values) and geographical distances (km) and second we compared genetic distance ($F_{ST}$ values) with altitudinal differences (m/a.s.l.). The geographic distances were obtained from the GPS coordinates (latitude and longitude) recorded during the sampling. All Mantel tests were assessed using 9999 permutations for the significance of the correlation.

The spatial component of genetic variation was further assessed by spatial autocorrelation [28] using GenAlEx [27]. The autocorrelation coefficient ($r$) was computed from the geographic distance between sampling locations and the genetic distance (pairwise $F_{ST}$ values). This measures the genetic similarity between individual pairs within an identified distance class. The size of the distance class influences the estimation of the spatial autocorrelation. We identified the appropriate distance class based on the observed distribution of pairwise geographic distance between the sampling locations (data not shown). The distance class sizes used were the following: 20, 40, 60, 80, 100, 120, 140, 160, 180, 200,
220, 240, 260, 280, 300, and 320 kilometers. Bootstrap estimates of \( r \) and random permutations were set at 9999 each for the test for significance. The upper and lower error bars in the correlogram bound the 95% confidence interval of \( r \) value as identified by bootstrap resampling. The upper and lower confidence limits bound the 95% confidence interval of the null hypothesis of no spatial structure in the data set.

The genetic relationships among the sampling locations (i.e., populations) were calculated by unweighted pair group method with the arithmetic mean (UPGMA) cluster method using the APE package [29] and R program [30] to create dendrogram using the genetic distance matrix (pairwise \( F_{ST} \) values) generated from GenAlEx.

**Phylogenetic analysis**

We constructed maximum likelihood (ML) gene trees for \( \text{cox1} \) and 16S haplotype sequences of \( I. ovatus \) and \( H. flava \) using PhyML ver 3.1 [31] under the default settings. We applied HKY and GTR nucleotide substitution models for \( \text{cox1} \) and 16S, respectively, as suggested by jModelTest ver 2 [32]. Additional sequences from China (MH208506, MH208512, MH208514, MH208522, MH208515-19, MH208524, MH208531, MH208574, MH208577, MH208579, MH208681-87, MH208689-93, MH208706, KU664519 [20]), Japan (Hokkaido, Yamanashi and Aomori) (AB231670, AB819241, AB819243, AB819244 [33-34]) and the USA (U95900; [35]) were included. We used \( Ixodes canisuga \) as an outgroup (KY962023, KY962074; [36]).

**Results**

The number of successfully sequenced and analyzed \( H. flava \) ranged from 1 to 36 individuals per study site, with more successfully sequenced for \( \text{cox1} \) (220/223;
98.7%) than for 16S (172/223; 77.1%). The number of *I. ovatus* ranged from 1 to 36 individuals per study site and, as with *H. flava*, more were successfully sequenced for *cox1* (307/320; 95.9%) than for 16S (284/320; 88.8%) (Additional File 1. Table S1). There were 63 and 60 *cox1* haplotypes in *H. flava* and *I. ovatus*, respectively, and 40 and 24 16S haplotypes (Table 1). Haplotype diversity ranged from 0.442 to 0.852 and nucleotide diversity ranged from 0.001 to 0.006 for both markers and species (Table 1).

AMOVA results revealed a high among-population divergence (41.54%) in *I. ovatus* *cox1* sequences, whereas both *cox1* and 16S markers of *H. flava* indicated no significant genetic variation among populations (Table 2). Global $F_{ST}$ values were highest in *I. ovatus* with values 0.415 in *cox1* and 0.074 in 16S sequences. Pairwise $F_{ST}$ values (0.500 to 1.00) of *I. ovatus* based on *cox1* showed significant genetic differences between most pairs of populations such as between locations 18 and 25 and between locations 28 and 25 (Additional File 2 Table S2). No significant differences were observed in 16S (Additional File 3 Table S3). Pairwise $F_{ST}$ values from the *cox1* marker of *H. flava* showed no significant genetic differences except between locations 18 and 15 and 18 and 25 (Additional File 4 Table S4). No significant genetic differentiation was also observed in 16S marker of *H. flava* except in between locations 3 and 20 and 4 and 20 (Additional File 5 Table S5).

The Mantel test of *I. ovatus* showed evidence of isolation by geographic distance in both markers: (*cox1* $r = 0.197, p = 0.008$; 16S $r = 0.293, p = 0.011$) (Figures 1a, b). In contrast, there was no evidence for isolation by altitudinal difference (*cox1* $r = -0.066, p=0.225$; 16S $r = -0.023, p = 0.577$). Both Mantel tests were not significant for *H. flava* (distance: *cox1* $r= -0.145, p= 0.23$; 16S $r = -0.049, p= 0.33$; altitude: $r$
= 0.092, p = 0.30; 16S r = 0.217, p= 0.06). The spatial autocorrelation test in *I. ovatus* revealed positive and significant correlation between genetic and geographic distance from 20 – 40 km, and 60 – 80 km, and 20 – 50 km and 120 – 160 km from *cox1* and 16S, respectively (Figure 1c, d).

The UPGMA cluster dendrogram from the pairwise *F*$_{ST}$ values from the *cox1*, *I. ovatus* (Figure 2) revealed 2 genetic clusters across the 29 sampling locations, where the cluster 2 included populations from the more western sites including the island (location 24 and 25) while cluster 1 mostly included populations in the northern and southern part of Niigata Prefecture (Figure 3). No evident genetic clustering on the dendrogram was observed from the *I. ovatus* 16S sequences and in both *cox1* and 16S *H. flava* sequences (Additional File 6 Figure S1; Additional File 7 Figure S2; Additional File 8 Figure S3).

The *cox1* gene tree of 60 *I. ovatus* haplotypes indicated three groups: group 1 of the published sequences from western China, one large group 2 containing the 58 haplotypes and 2 divergent haplotypes (Hap 60 and Hap 59), and group 3 from western China (Figure 4). The 2 divergent haplotypes were singletons. One was found in location 29 and the other in location 7. The published haplotype from Hokkaido [33] occurred within our large group and the published sequences from western China did not. The *I. ovatus* 16S gene tree (24 newly sequenced haplotypes) (Additional File 9 Figure S4) also recovered published haplotypes from Japan (Yamanashi and Aomori; [34]) as well as from North America [35] within the same cluster as our new sequences. The *cox1* haplotype sequences of *H. flava* were similar to reference sequences from China (Additional File 10 Figure S5) [unpublished manuscript] while the 16S haplotype sequences (Additional File 11 Figure S6) showed similarities with Japan (Kagoshima, Fukui Aomori, Fukui,
Yamanashi, Kagawa and Ehime [34]) and China [unpublished manuscript] reference sequences from Genbank.

Discussion

**Contrasting population genetic structures between *H. flava* and *I. ovatus***

Our results supported our hypothesis that *H. flava* and *I. ovatus* display contrasting population genetic structure. The noticeable high gene flow in *H. flava* may be a result of the higher level of mobility of its hosts such as birds in tick’s immature stage and the more pronounced genetic structure and lower gene flow in *I. ovatus* may be due to the restricted ability of its small mammalian hosts at the immature stage to disperse further. 28 species of birds were previously reported as hosts of *H. flava*, mostly from the order Passeriformes known as migratory birds [37]. The host for adults of *I. ovatus* included large mammals such as hares while immature forms are found on small mammals such as small rodents and rats [15]. The movement of small mammals is limited because of environmental and ecological factors such as difficulty in crossing roads and higher possibility of road kill due to vehicle collision and or traffic [38], which may explain the genetic structuring in *I. ovatus*. The host preference of the adults *H. flava* are large mammals (e.g. cows, horses, wild boar, deer, bear) and hares, dogs, rodents while the immature forms parasitize birds [15]. Mammals and birds may have wide habitats ranges that may allow maintaining the gene flow of *H. flava* between the locations in Niigata, as previously observed in *Amblyomma americanum* populations [5,6,39], and *Ixodes ricinus* [40]. Birds present the greatest potential for farther dispersal of *H. flava* that could reduce its genetic structuring [41] that supports the results of our study. Several alternative factors such as tick behavior, biology and ecology can also
affect the tick dispersal movement. For example, long movement patterns, resistance against unfavorable weather conditions causing high survival in their hosts and low host specificity [4] are possible reasons for the low genetic structuring in \textit{H. flava} populations. It could also be influenced by ecological factors such as changes in land use and cover, wildlife and forestry management as observed in the previous study of [42] in the \textit{Ixodes ricinus} tick populations from Europe. Previous study on two tick species: \textit{Hyalomma rufipes} and \textit{Amblyomma hebraum} also displayed contrasting genetic pattern despite the overlap sharing of highly mobiles hosts such as birds [43] due to the capacity of the immature stages of ticks to survive off the hosts in various habitat conditions [44-46]. Population structure can also be influenced by assortative mating (e.g. \textit{Ixodes ricinus}) wherein genetically similar individuals tend to mate, rather than random mating resulting in increased genetic divergence [47]. Our study does not have supporting data to test these alternative factors thus it is suggested in future studies to further analyze these factors that could play a role in the tick’s dispersal movement.

Based on previous studies [48-49], isolation by distance suggests that the populations are in equilibrium of migration and genetic drift at the spatial scale of Niigata Prefecture. Our results were congruent with a previous study of \textit{Amblyomma ovale} that showed positive relationship of geographic and genetic distance at the spatial scale of the whole of Brazil whose main hosts for the immature stage are small rodents while tapir and jaguar in its adult stage [50]. In contrast, \textit{H. flava} populations showed an absence of isolation by distance. In circumstances when geographic distance can’t explain the genetic structure of tick populations, other factor such as ecological variables (e.g. land use) maybe affecting the genetic structuring within tick populations [7, 40, 51].
A significant and positive spatial autocorrelation of *I. ovatus* in both *cox1* and *16S* sequences was observed within the geographic range of 20 – 40 km and 60- 80 km in *cox1* sequences and 20 – 50 km and 120 – 160 km range in the *16S* sequences. We can infer that the positive *r* values at the maximum geographic rages of 60-80 km in *cox1* and 120 - 160 km in *16S* *I. ovatus* sequences indicate that gene flow is occurring in the wide geographic range. The lack of significance at 40-60 km in *cox1* may suggest patch structure indicating genetic similarity or variation [28] as observed on limpet species, *Siphonaria raoulensis* from New Zealand that also displayed recovery of positive *r* values at greater distances [52]. However, no autocorrelation was observed in *H. flava* at any distance class, suggesting that gene flow occurring has no specific pattern favoring certain distances [53-55]. Since the sampling geographic range of our study was within Niigata Prefecture and distances >220 km were not included in our sampling, we suggest that future studies collect samples at a wider geographic scale to more precisely determine the geographic distance at which autocorrelation exists.

*16S* *I. ovatus* sequences showed genetic differentiation (pairwise *F*<sub>ST</sub> = 0.5-0.6) in few pairs of sampling locations as compared to the *cox1* sequences which may be due to lower variability of the marker thus providing inadequate genetic variation for population genetic analysis, as previously observed in *Amblyomma ovale* [50] and *Rhipicephalus microplus* [56-57]. The low variability of the *16S* in *I. ovatus* may not be sufficient to infer intraspecific relationship and is further supported by its low (*nd* = 0.001) nucleotide diversity as compared in *cox1* *I. ovatus* (*nd* = 0.004). Thus, it may also explain the grouping observed in the *cox1* *I. ovatus* tree which is absent in the *16S* *I. ovatus* sequences. Despite of these, we used *16S* mitochondrial gene because this gene mutates at a rate that is informative for species-level
phylogenetic and broad biogeographic inferences [2]

The negative pairwise $F_{ST}$ values obtained in *H. flava* sequences do not have any biological meaning. These results from the unbiased estimator allowing a negative $F_{ST}$ when the sample size is small. [58-60]. It can also be noted that we found significant pairwise $F_{ST}$ values in 2 sampling location in *H. flava* *cox1* sequences probably due to the fact that both locations have only one tick sample. Although, some sampling sites in this study have small sample size ($n < 10$), the *cox1* and 16S sequences provided valuable insight into the genetic structure and gene flow patterns of the present-day tick populations. Thus, for future research, we recommend to increase the sample size per sampling site.

**Species complex formation in *I. ovatus* *cox1* sequences**

Our *cox1* *I. ovatus* gene tree showed Japanese individuals to form a group that was distinct from haplotypes from southern China [20]. Despite the low gene flow we found in *cox1* *I. ovatus*, haplotypes found from Niigata were closely related to the published sequence from Hokkaido, Japan which may indicate that these ticks originated from a diverse set of geographical locations in Japan which might be transported by its hosts or is undergoing recent population expansion from northern Japan (Hokkaido) to south (Niigata) or *vice versa*. We found three groups (China 1, Japan and China 2) and 2 slightly divergent *cox1* haplotypes (Hap 60 and 59) in the Japan group of *I. ovatus* in Niigata. Considering two or more cryptic species can be concealed in one morphologically described species [59], the occurrence of the 3 groups and the divergent haplotypes suggests that *I. ovatus* may be a species complex. Previous studies have also observed species complexes in *Ixodes* and *Rhipicephalus* [19-20,57;62] suggesting that morphological criteria for species
differentiation alone are equivocal and that genetic analysis is important.

Conclusion

Two Ixodid ticks, *H. flava* and *I. ovatus* exhibited contrasting patterns of genetic structure in Niigata Prefecture, Japan. We suggest that these differences are due to differences in host mobility of the immature life stages of the ticks. The more pronounced population genetic structure of *I. ovatus* might be influenced by the restricted movement of its small mammalian host in its immature stage, whereas the lack of structure in *H. flava* may be due to its avian-mediated movement in its immature stage. Even though *I. ovatus* populations were genetically structured within Niigata, a published haplotype from Hokkaido was also found, indicating that widespread dispersal is possible. The occurrence of three groups and the divergent cox1 haplotypes in *I. ovatus* emphasizes the need of additional methods in determining the existence of species complex in *I. ovatus* populations in Japan.

List of Abbreviations

*flava* - *Haemaphysalis flava*

*ovatus* - *Ixodes ovatus*

AMOVA - Analysis of molecular variance

UPGMA - unweighted pair group method with arithmetic mean

ML - maximum likelihood

$F_{ST}$ - fixation index

Declarations

**Ethics approval and consent to participate**
Not applicable

Consent for publication

Not applicable

Availability of data and materials

All data generated or analyzed during the study are included in this published article and its supplementary additional files. All the newly generated sequences are available in the Genbank database under the accession numbers _______.

Competing interests

The authors declare that they have no competing interests.

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Authors’ contributions

MAFR, MG, MM, KW conceptualized and the designed the experiment. MS, TT, RA, SI and MOS designed the sampling collection, collected and identified the tick samples of this study. MD and KT conducted the molecular analyses. MAFR, MG and KW performed the data analysis and wrote the manuscript. All authors read and approve the manuscript.

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Tables

Table 1. Summary of *cox1* and *16S* haplotype distribution, variability, genetic diversity of *Ixodes ovatus* and *Haemaphysalis flava* populations in Niigata Prefecture

| Marker | Species     | n  | h  | s  | k     | hd±SD       | nd±SD       |
|--------|-------------|----|----|----|-------|-------------|-------------|
|        | *I. ovatus* | 307| 60 | 65 | 2.728 | 0.852 ± 0.012 | 0.004 ± 0.000 |
|        | *H. flava*  | 220| 63 | 60 | 1.472 | 0.789 ± 0.026 | 0.002 ± 0.000 |
| *16S*  | *I. ovatus* | 284| 24 | 22 | 0.699 | 0.442 ± 0.037 | 0.001 ± 0.000 |
|        | *H. flava*  | 172| 40 | 49 | 2.447 | 0.835 ± 0.024 | 0.006 ± 0.000 |

Abbreviations: *n* sample size; *h* number of haplotypes; *s* number of polymorphic sites; *k* mean number of nucleotide differences; *hd±SD* haplotype diversity ± Standard Deviation; *nd±SD* nucleotide diversity ± Standard Deviation

Table 2. Analysis of molecular variance (AMOVA) using *cox1* and *16S* of *Ixodes ovatus* and *Haemaphysalis flava* populations
| Marker | Species     | df   | ss     | vc     | pv     | $F_{ST}$ |
|--------|-------------|------|--------|--------|--------|----------|
| cox1   | I. ovatus   | Among| 28     | 190.38 | 0.5803 Va| 41.54    | 0.4154** |
|        |             | Within| 278    | 227.04 | 0.8167 Vb| 58.46    |          |
|        | H. flava    | Among| 16     | 12.42  | 0.0039 Va| 0.53     | 0.0053   |
|        |             | Within| 203    | 148.73 | 0.7327 Vb| 99.47    |          |
| 16S    | I. ovatus   | Among| 22     | 14.04  | 0.0260 Va| 7.39     | 0.0739** |
|        |             | Within| 261    | 84.87  | 0.3252 Vb| 92.61    |          |
|        | H. flava    | Among| 12     | 23.6   | 0.0642 Va| 5.21     | 0.0521** |
|        |             | Within| 159    | 185.58 | 1.1672 Vb| 94.79    |          |

Abbreviations: $df$ degrees of freedom; $ss$ sum of squares; $Va$, $Vb$ and $Vc$ are associate covariance components; $pv$ percentage variation; *$P<0.05$ **$P<0.01$

Additional Files Description

**Additional File 1.xls Table S1.** Summary of *Haemaphysalis flava* and *Ixodes ovatus* collected from the different locations of Niigata Prefecture and its corresponding sample number

**Additional File 2.xls Table S2.** Pairwise comparison of genetic differentiation ($F_{ST}$) calculated for all *I. ovatus* using the *cox1* gene

**Additional File 3.xls Table S3.** Pairwise comparison of genetic differentiation ($F_{ST}$) calculated for all *I. ovatus* using the *16S* gene
**Additional File 4.xls Table S4.** Pairwise comparison of genetic differentiation ($F_{ST}$) calculated for all *H. flava* using the *cox1* gene

**Additional File 5.xls Table S5.** Pairwise comparison of genetic differentiation ($F_{ST}$) calculated for all *H. flava* using the 16S gene

**Additional File 6.docx Figure S1.** An unweighted pair group method with the arithmetic mean (UPGMA) dendrogram of *I. ovatus* based on the pairwise genetic distance ($F_{ST}$) of 16S among 23 sampling locations across Niigata Prefecture, Japan.

**Additional File 7.docx Figure S2** An unweighted pair group method with the arithmetic mean (UPGMA) dendrogram of *H. flava* based on the pairwise genetic distance ($F_{ST}$) of *cox1* among 17 sampling locations across Niigata Prefecture, Japan.

**Additional File 8.docx Figure S3** An unweighted pair group method with the arithmetic mean (UPGMA) dendrogram of *H. flava* based on the pairwise genetic distance ($F_{ST}$) of 16S among 17 sampling locations across Niigata Prefecture, Japan.

**Additional File 9.docx Figure S4** Phylogenetic tree of 16s haplotype sequences of *I. ovatus* inferred by maximum likelihood analysis with evolutionary model GTR with sequences from China (MH208506, MH208512, MH208514 – MH208519, MH208522, MH208524, MH208531, MH208574, MH208577, MH208579, KU664519), Japan (AB819241, AB819242, AB819243, AB819244) and US (U95900). An outgroup (*Ixodes canisuga*) was also included.

**Additional File 10.docx Figure S5** Phylogenetic tree of *cox1* haplotype sequences of *H. flava* inferred by maximum likelihood analysis with evolutionary model HKY with sequences from China (KY021800- KY021807; KY021810-KY021819, KY003181, JQ62588 - JQ625889; JQ737097; JF758632). An outgroup (*Ixodes canisuga*) was also included.
Additional File 11.docx Figure S6  Phylogenetic tree of 16S haplotype sequences of *H. flava* inferred by maximum likelihood analysis with evolutionary model GTR with sequences from China (KC844858-KC844867; KC844880-KC844882, M696720, KP324926, KX450279) and Japan (AB19177-AB819192). An outgroup (*Ixodes canisuga*) was also included.

Figures

![Figure 1](image)

Figure 1

Plots (a) and (b) show the relationship between the pairwise FST values and the g
Figure 2

An unweighted pair group method with the arithmetic mean (UPGMA) dendrogram
Figure 3

The distribution of the two genetic clusters as observed in the UPGMA cluster den
Figure 4

Maximum likelihood gene tree of cox1 haplotype sequences of I. ovatus with sequ
Supplementary Files

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