



Research Article

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
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High genetic diversity of *Echinococcus canadensis* G10 in northeastern Asia: is it the region of origin?

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Abstract

Echinococcus canadensis consists of 4 genotypes: G6, G7, G8 and G10. While the first 2 predominantly infect domestic animals, the latter are sylvatic in nature involving mainly wolves and cervids as hosts and can be found in the northern temperate to Arctic latitudes. This circumstance makes the acquisition of sample material difficult, and little information is known about their genetic structure. The majority of specimens analysed to date have been from the European region, comparatively few from northeast Asia and Alaska. In the current study, *Echinococcus* spp. from wolves and intermediate hosts from the Republic of Sakha in eastern Russia were examined. *Echinococcus canadensis* G10 was identified in 15 wolves and 4 cervid intermediate hosts. Complete mitochondrial cytochrome c oxidase subunit 1 (*cox1*) sequences were obtained from 42 worm and cyst specimens from Sakha and, for comparison, from an additional 13 G10 cysts from Finland. For comparative analyses of the genetic diversity of G10 of European and Asian origin, all available *cox1* sequences from GenBank were included, increasing the number of sequences to 99. The diversity found in northeast Asia was by far higher than in Europe, suggesting that the geographic origin of *E. canadensis* (at least of G10) might be northeast Asia.

Introduction

Cystic echinococcosis (CE) is a zoonotic infectious disease caused by several members of the cestode species complex *Echinococcus granulosus* sensu lato (s.l.). CE is considered by the World Health Organization as one of the most important ‘neglected zoonotic diseases’ (WHO, 2020). The life cycle of these parasites is obligate between carnivorous definitive (mostly canids) and herbivorous intermediate hosts (Romig *et al.*, 2017). The adult worms, which are only a few millimetres in length, live in the small intestine of the definitive hosts, and their eggs are released into the environment with the feces. After oral ingestion of the eggs *via* contaminated water or food by intermediate hosts, the larval cyst-like metacestode may develop in various organs, most often in the liver and lungs. The protoscoleces, which develop in large numbers in these cysts, grow into adult worms when ingested by the definitive host during predation or scavenging (Thompson, 2017). If infectious eggs are accidentally ingested by humans, development of metacestodes in internal organs may occur, leading to CE due to damage of organs by the growth of cysts, which may ultimately be fatal (Kern *et al.*, 2017; Thompson, 2017).

The species complex *E. granulosus* s.l. consists of 5 distinct species: *E. granulosus* sensu stricto (s.s.), *Echinococcus equinus*, *Echinococcus ortleppi*, *Echinococcus felidis* and *Echinococcus canadensis*. The latter species is the most diverse, holding distinct genotypes (G6, G7, G8 and G10), and there is debate whether *E. canadensis* should be split into 2 or even 3 species (Lymbery *et al.*, 2015; Yanagida *et al.*, 2017; Laurimäe *et al.*, 2018a). The various genotypes, formerly named ‘strains’, differ with respect to life cycle, geography and genetics. Although the closely related and globally distributed G6 and G7 are mostly associated with a livestock–dog life cycles (often involving camels or pigs), G8 and G10 only occur in the Northern Hemisphere and are sylvatic in nature involving mainly wolves and cervids such as moose and reindeer as hosts (Romig *et al.*, 2017). According to the distribution of their hosts, G8 and G10 can be found in the northern temperate to Arctic latitudes (Romig *et al.*, 2017).

While all variants of G6 and G7 are genetically close (and are often referred to as the G6/7 cluster) (Addy *et al.*, 2017), the relationship of G8 and G10 with each other and with G6/7 is differently resolved depending on whether nuclear or mitochondrial DNA is considered. Analysis of the mitochondrial genome suggests a closer affinity of G10 to G6/7, while some

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nuclear marker genes support a clade with G8 and G10 (Yanagida *et al.*, 2017; Laurimäe *et al.*, 2018a).

Taxonomic decisions on *E. canadensis* have been postponed so far due to a lack of sufficient data, especially on the northern genotypes G8 and G10. In contrast to G6/7, whose genetic structure is well known, only very few specimens of G8 and G10 have ever been characterized and showed little intra-strain variability. However, the great majority came from the European region, and the study of the nuclear genome was performed exclusively with sample material from Europe (Laurimäe *et al.*, 2018a). The northeastern part of Asia is particularly data-deficient, although this is the only known region where all genotypes of *E. canadensis* have been recorded, leading to speculation on the geographic origin of the species (Konyaev *et al.*, 2013; Ito *et al.*, 2014; Zhang *et al.*, 2014; Yang *et al.*, 2015; Wu *et al.*, 2018; Hua *et al.*, 2019).

In the present study, we collected samples of *Echinococcus* spp. from a variety of wild and semi-domestic definitive and intermediate hosts in the Republic of Sakha (Russian Federation), and examined them for species, genotypes and haplotypic diversity to obtain new insights into the structure of the *E. canadensis* cluster.

Materials and methods

Parasite material

The parasite material originated from wild and semi-domestic animals from the Republic of Sakha, located in the Russian Far East in northeast Asia (Fig. 1).

Echinococcus worms from definitive hosts and cyst material from intermediate hosts were available for the present study. Adult worms were isolated from small intestines of 94 legally hunted wolves (*Canis lupus*) from Sakha in the period 2011–2015 and preserved in a fixative solution. One cyst each from moose (*Alces alces*), elk (*Cervus canadensis*) and roe deer (*Capreolus pygargus*) and 2 from reindeer (*Rangifer tarandus*) could be opportunistically collected from hunted animals in Sakha during meat inspection and preserved in 70% ethanol (EtOH).

In addition, 13 cyst samples of *E. canadensis* G10 stored in 70% EtOH from 3 reindeer and 10 moose from Finland were also available for molecular analyses (Fig. 1).

Sample preparation

The content of the sample tubes with the adult worm material was examined under an inverted microscope. Individual worms or worm fragments were aspirated with a pipette and each transferred to a 200 μ L polymerase chain reaction (PCR) tube containing 20 μ L of 0.02 M sodium hydroxide (NaOH). Of the cyst samples, a small tissue piece of $\sim 1 \times 1$ mm² in size was cut with a scalpel and each transferred to separate tubes containing 30 μ L of 0.02 M NaOH. The isolated single worms and cyst material were lysed at 95°C for 15 min (Nakao *et al.*, 2003). The lysate was used directly as a template for the subsequent molecular analyses or stored at –20°C until further use. For the cyst samples that failed to yield a positive PCR result, DNA was obtained by proteinase K digestion followed by phenol–chloroform extraction and EtOH precipitation as described previously (Dinkel *et al.*, 1998).

DNA amplification and sequencing

Species identification was done by nested PCR and sequencing. The target gene was the ~ 1600 bp long mitochondrial cytochrome c oxidase subunit 1 (*cox1*) gene. To obtain the complete *cox1* gene,

different primer combinations were used targeting the front part or the back part of the gene. If the PCR remained negative, an attempt was made to amplify a *cox1* fragment of ~ 200 bp in length to identify the species. If this attempt also remained negative, additional fragments of other mitochondrial genes, the cytochrome b (*cob*, ~ 170 bp) or NADH dehydrogenase subunit 1 gene (*nad1*, ~ 190 bp), were targeted. All primers used are listed in Table 1.

The amplification was done *via* nested PCR. For the first PCR, a reaction volume of 25 μ L was prepared. The mixture contained 10 mM Tris-hydrochloric acid (Tris-HCl) (pH 8.3), 50 mM potassium chloride (KCl), 2 mM magnesium chloride (MgCl₂), 200 μ M of each deoxynucleotide triphosphates (dNTPs), 6.25 pmol of each first PCR primer and 0.625 U of Taq polymerase (Applied Biosystems, Carlsbad, CA, USA). One microlitre of the cyst or worm lysate, or extracted DNA was added to the PCR mixture as a template. The volume of the nested PCR reaction mixture was 50 μ L and consisted of 10 mM Tris-HCl (pH 8.3), 50 mM KCl, 2 mM MgCl₂, 200 μ M of each dNTPs, 12.5 pmol of each nested PCR primer and 1.25 U of Taq polymerase and 2 μ L of the first PCR product as template DNA.

The conditions during amplification for the PCRs were as follows: initial denaturation at 95°C for 5 min followed by 35 cycles with denaturation at 95°C for 30 s, annealing at 50°C for 30 s, elongation at 72°C for 60 s (for front and back parts of *cox1* gene) or 30 s (for small fragments of *cox1*, *cob* and *nad1* genes) and a final elongation at 72°C for 5 min. Amplification results were obtained on a 1.5% agarose gel stained with GelRed™. PCR products were purified with a High Pure PCR Product Purification Kit (Roche, Mannheim, Germany) and sent to Microsynth Seqlab GmbH (Göttingen, Germany) for sequencing. The sequences were viewed, edited and assembled using the GENTle V1.9.4 program (Manske M., University of Cologne, Germany) and compared with GenBank entries using the NCBI basic local alignment search tool (BLAST) for species identification.

Analyses of haplotype networks and diversity indices

The haplotypes and the genetic diversity of the obtained complete *cox1* sequences were analysed. Haplotype identification and creation of haplotype networks were calculated with TCS 1.2 software (Clement *et al.*, 2000) using statistical parsimony (Templeton *et al.*, 1992). Networks were visualized using the online program tcsBU (Santos *et al.*, 2016). The haplotype and nucleotide diversity indices were analysed by DnaSP 6.12 (Rozas *et al.*, 2017) and analysis of molecular variance, degree of genetic differentiation (pairwise fixation index, *F*_{st}) and neutrality indices of Tajima's *D* and Fu's *F*_s were calculated using Arlequin v 3.5.2.2 software (Excoffier *et al.*, 2005).

In addition to 13 samples from Finland examined here, all *E. canadensis* G10 sequences of the complete *cox1* gene available in GenBank were included in the analyses for comparison. The samples behind the sequences originated from eastern Russia (Sakha), Mongolia, China (Tibet), the USA (Alaska) and Europe (Estonia, Sweden, Finland and Russia) and are listed in Table 2. For the comparison of the genetic diversity and neutrality indices of the sequences found in northeast Asia and Europe, the frequencies of the haplotypes discovered in different studies were extracted from the corresponding literature (Table 2).

Results

Parasite species identification

Adult worms or worm fragments could be isolated from the 94 wolf samples. Depending on the sample, between 1 and 36



Figure 1. Map of Russia with the Republic of Sakha (red), blue ellipse shows area of sample collection in Sakha; green ellipse shows Finland (source: www.wikipedia.com; CC BY-SA 4.0).

worms or worm fragments, in total 713 were isolated individually and examined. Amplification of at least one of the small 170–200 bp gene fragments was successful only in 191 worms from 35 samples. *Echinococcus canadensis* G10 was detected in 15 wolves (1–14 worms per wolf) and *Echinococcus multilocularis* in 2 wolves (1 and 2 worms). Of the *E. multilocularis* specimens only the small *nad1* fragment could be amplified resulting in ~100 bp long sequences. The comparison with GenBank did not allow an assignment to known haplotypes. Identical sequences were detected in isolates from e.g. Europe, China or

St. Lawrence Island. Twenty-one wolves showed infection with juvenile *Taenia* sp., of which a ~200 bp *nad1* fragment had 95.5% similarity with *Taenia multiceps* (NC012894; Jia *et al.*, 2010); 2 of these wolves were co-infected with *E. canadensis* G10. In 38 *E. canadensis* G10 worms from 10 wolves, amplification and sequencing of the complete *cox1* gene was successful. Analysis of the intermediate hosts also revealed infection with *E. canadensis* G10 in 2 reindeer, 1 moose and 1 elk; the cyst from the roe deer could not be amplified. All of the 13 cyst samples from Finland (3 reindeer and 10 moose) belonged to *E. canadensis* G10. From all these cyst samples the complete *cox1* gene could be sequenced. A total of 42 *cox1* sequences from Sakha and 13 sequences from Finland were thus available for the further haplotype analysis (Table 2). The sequences were deposited at NCBI GenBank under the accession numbers OR420689–OR420703.

Table 1. Primer pairs used for PCR

Target gene	Primers for the 1st PCR (5'–3')	Primers for the 2nd PCR (5'–3')
<i>cox1</i> front part	F: TTA CTG CTA ATA ATT TTG TGT CAT ^a R: ATA CCA GTA ACA CCT CCA AAC G	F: GTG TCA TTT AGG TTT GAC TTT CTC R: AAC ATA TAC AAC CAA GTA AAC ACC ^a
<i>cox1</i> back part	F: TTT GCT ATG TTT TCT ATA GTG R: CAA AAA CAT ACT TTA AAA AAC TCC	F: CAT CAT ATG TTT ACT GTT GGA TTG G R: GCA TGA TGC AAA AGG CAA ATA AAC
<i>cox1</i> (<200 bp) ^a	F: ACT GTT GGG TTG GAT GTT AAG ACG R: CAT AAC ATA ATG AAA ATG AGC CA	F: TAG TTC TGT TAC TAT GAT TAT AGG R: GTA TCA TGT AAA ACA TTA TCC AAC
<i>cob</i> (<200 bp) ^a	F: TTA TGC TAT ACT TCG GTG TAT TA R: ATA AGG ATA CTC CGG ATG ACA AC	F: TCG GTG TAT TAA TTC GAA GAT TG R: GAT GAC AAC CAC CCA AAT AAG TC
<i>nad1</i> (<200 bp) ^a	F: TGT TTT TGA GAT CAG TTC GGT GTG R: CAT AAT CAA ACG GAG TAC GAT TAG	F: CAG TTC GGT GTG CTT TTG GGT CTG R: GAG TAC GAT TAG TCT CAC ACA GCA

^aHüttner *et al.* (2008).

F, forward primer; R, reverse primer.

Number of haplotypes and parsimony network

A total of 99 complete *cox1* gene sequences of *E. canadensis* G10, including the present sequences and GenBank entries, were available for the calculation of number of haplotypes and network.

Analysis of the sequences of the 42 samples from Sakha revealed 15 haplotypes of *E. canadensis* G10, while only 1 haplotype was detected in the Finnish samples of 13 cysts (Table 2). When all 99 sequences were analysed according to their geographic origin, 3 haplotypes could be detected in 40 sequences from Europe, 17 haplotypes in the 47 eastern Russian sequences (42 from the present study and 5 additional from GenBank) and 19 haplotypes in northeast Asia (including the sequences from eastern Russia, and GenBank entries from Mongolia and Tibet). The 5 sequences with Alaskan origin resulted in 3 haplotypes.

In addition to the 99 *E. canadensis* G10 sequences, 1 sequence each of genotypes 6, 7 and 8 of *E. canadensis* was included in the haplotype network (Fig. 2, Table 2). The network

Table 2. Geographic origin, host, haplotype, accession number and reference of samples and sequences used for the analyses

Geographic region	Sample origin	Haplotype	Host	Accession no.	Reference
Northeast Asia	Sakha	H01	Wolf_2	OR420689	This study
	Sakha	H01	Wolf_3	OR420689	This study
	Sakha	H01	Wolf_3	OR420689	This study
	Sakha	H01	Wolf_4	OR420689	This study
	Sakha	H01	Wolf_4	OR420689	This study
	Sakha	H01	Wolf_4	OR420689	This study
	Sakha	H01	Wolf_4	OR420689	This study
	Sakha	H01	Wolf_4	OR420689	This study
	Sakha	H01	Wolf_4	OR420689	This study
	Sakha	H01	Wolf_2	OR420689	This study
	Sakha	H01	Wolf_5	OR420689	This study
	Sakha	H01	Wolf_6	OR420689	This study
	Sakha	H01	Wolf_6	OR420689	This study
	Sakha	H01	Wolf_8	OR420689	This study
	Sakha	H01	Wolf_9	OR420689	This study
	Sakha	H01	Wolf_9	OR420689	This study
	Sakha	H01	Wolf_10	OR420689	This study
	Sakha	H01	Wolf_10	OR420689	This study
	Sakha	H01	Reindeer	OR420689	This study
	Sakha	H01	Elk	OR420689	This study
	Sakha	H02	Wolf_1	OR420690	This study
	Sakha	H02	Wolf_5	OR420690	This study
	Sakha	H02	Wolf_5	OR420690	This study
	Sakha	H02	Wolf_7	OR420690	This study
	Sakha	H02	Wolf_8	OR420690	This study
	Sakha	H02	Wolf_9	OR420690	This study
	Sakha	H02	Wolf_10	OR420690	This study
	Sakha	H03	Wolf_5	OR420691	This study
	Sakha	H04	Wolf_2	OR420692	This study
	Sakha	H04	Wolf_10	OR420692	This study
	Sakha	H06	Wolf_2	OR420693	This study
	Sakha	H07	Wolf_7	OR420694	This study
	Sakha	H08	Wolf_7	OR420695	This study
	Sakha	H09	Wolf_7	OR420696	This study
	Sakha	H11	Wolf_2	OR420697	This study
	Sakha	H12	Wolf_8	OR420698	This study
	Sakha	H13	Wolf_7	OR420699	This study
	Sakha	H14	Wolf_8	OR420700	This study
	Sakha	H15	Wolf_10	OR420701	This study
	Sakha	H16	Wolf_8	OR420702	This study
	Sakha	H19	Moose	OR420703	This study
	Sakha	H19	Reindeer	OR420703	This study
	Yakutia (Sakha) ^a	H05	Moose	AB777911	Nakao <i>et al.</i> (2013)
	Yakutia (Sakha) ^a	H10	Wolf	AB777912	Nakao <i>et al.</i> (2013)
	Yakutia (Sakha) ^a	H01	Reindeer	AB777913	Nakao <i>et al.</i> (2013)
	Yakutia (Sakha) ^a	H03	Human	AB777914	Nakao <i>et al.</i> (2013)

(Continued)

Table 2. (Continued.)

Geographic region	Sample origin	Haplotype	Host	Accession no.	Reference
	Yakutia (Sakha)	H03	Human	AB777914	Konyaev <i>et al.</i> (2013)
	Mongolia	H17	Wolf	AB813184	Ito <i>et al.</i> (2013)
	Mongolia	H17	Wolf	AB813184	Ito <i>et al.</i> (2013)
	Mongolia	H02	Wolf	AB813185	Ito <i>et al.</i> (2013)
	Mongolia	H02	Wolf	AB813185	Ito <i>et al.</i> (2013)
	Mongolia	H17	Human	AB893264	Ito <i>et al.</i> (2014)
	Mongolia	H17	human	AB893264	Ito <i>et al.</i> (2014)
	Tibetan Plateau	H18	Yak	MG597240	Wu <i>et al.</i> (2018)
Europe	Finland	H19	Reindeer	OR420703	This study
	Finland	H19	Moose	OR420703	This study
	Finland	H19	Reindeer	OR420703	This study
	Finland	H19	Moose	OR420703	This study
	Finland	H19	Moose	OR420703	This study
	Finland	H19	Reindeer	OR420703	This study
	Finland	H19	Moose	OR420703	This study
	Finland	H19	Moose	OR420703	This study
	Finland	H19	Moose	OR420703	This study
	Finland	H19	Moose	OR420703	This study
	Finland	H19	Moose	OR420703	This study
	Finland	H19	Moose	OR420703	This study
	Finland	H19	Moose	OR420703	This study
	Finland	H19	Moose	OR420703	This study
	European Russia	H20	Moose	LC184605	Yanagida <i>et al.</i> (2017)
	European Russia	H19	Moose	AB745463	Yanagida <i>et al.</i> (2017)
	European Russia	H19	Moose	AB745463	Yanagida <i>et al.</i> (2017)
	European Russia	H19	Moose	AB745463	Yanagida <i>et al.</i> (2017)
	European Russia	H19	Moose	AB745463	Yanagida <i>et al.</i> (2017)
	European Russia	H19	Moose	AB745463	Nakao <i>et al.</i> (2013)
	European Russia	H19	Moose	AB745463	Nakao <i>et al.</i> (2013)
	European Russia	H19	Moose	OQ161118	Laurimäe <i>et al.</i> (2023)
	European Russia	H19	Moose	OQ161119	Laurimäe <i>et al.</i> (2023)
	European Russia	H19	Moose	OQ161120	Laurimäe <i>et al.</i> (2023)
	European Russia	H19	Moose	OQ161121	Laurimäe <i>et al.</i> (2023)
	Estonia	H05	Moose	OQ161108	Laurimäe <i>et al.</i> (2023)
	Sweden	H19	Reindeer	OQ161122	Laurimäe <i>et al.</i> (2023)
	Finland	H19	Moose	AB745463	Nakao <i>et al.</i> (2013)
	Finland	H19	Moose	AB745463	Nakao <i>et al.</i> (2013)
	Finland	H19	Moose	AB745463	Nakao <i>et al.</i> (2013)
	Finland	H19	Wolf	AB745463	Yanagida <i>et al.</i> (2017)
	Finland	H19	Wolf	AB745463	Yanagida <i>et al.</i> (2017)
	Finland	H19	Moose	OQ161109	Laurimäe <i>et al.</i> (2023)
	Finland	H19	Moose	OQ161110	Laurimäe <i>et al.</i> (2023)
	Finland	H19	Moose	OQ161111	Laurimäe <i>et al.</i> (2023)
	Finland	H19	Moose	OQ161112	Laurimäe <i>et al.</i> (2023)
	Finland	H19	Reindeer	OQ161113	Laurimäe <i>et al.</i> (2023)
	Finland	H19	Reindeer	OQ161114	Laurimäe <i>et al.</i> (2023)
	Finland	H19	Reindeer	OQ161115	Laurimäe <i>et al.</i> (2023)

(Continued)

Table 3. Diversity and neutrality indices of *E. canadensis* G10 based on complete *cox1* gene (1608 bp)

Region/country	<i>n</i>	<i>n_H</i>	Diversity indices		Neutrality indices	
			<i>H_d</i> ± s.d.	<i>N_d</i> ± s.d.	Tajima's <i>D</i>	Fu's <i>F_s</i>
Europe	40	3	0.0987 ± 0.0638	0.0078 ± 0.0099	-2.0034*	-1.0744
Sakha	42	15	0.7526 ± 0.0647	0.0694 ± 0.0434	-1.6215*	-7.1116*
Eastern Russia ^a	47	17	0.7817 ± 0.0576	0.0786 ± 0.0478	-1.5262*	-8.0627*
Northeast Asia ^b	54	19	0.8204 ± 0.0449	0.0943 ± 0.0554	-1.5289*	-8.0256*
Alaska (USA)	5	3	0.8000 ± 0.1640	0.0313 ± 0.0291	0.2431	-0.4754
Total (Global)	99	23	0.7858 ± 0.0334	0.1118 ± 0.0634	-1.2954	-7.1149*

n, number of isolates (sequences); *n_H*, number of haplotypes; *H_d*, haplotype diversity; *N_d*, nucleotide diversity; s.d., standard deviation.

^aIncluding sequences from Sakha and additional from eastern Russia.

^bIncluding sequences from eastern Russia, Mongolia, and Tibet.

*Significant at *P* < 0.05.

Population diversity and neutrality indices

Analyses of the haplotype (*H_d*) and nucleotide diversity (*N_d*) indices show low values in Europe, compared to the diversity found in northeast Asia, which are 8.3 (*H_d*) and 12.1 (*N_d*) times higher, respectively (Table 3). The value of European diversity *H_d* was 0.0987 and that of nucleotide diversity *N_d* was 0.0078. The indices from Sakha determined in the present study are almost 1 order of magnitude higher compared to Europe (*H_d*: 0.7526; *N_d*: 0.0694) and increase further with the inclusion of the other sequences from eastern Russia, Mongolia and China (*H_d*: 0.8204; *N_d*: 0.0943). The Alaskan samples show also high haplotype diversity values (*H_d*: 0.8000); the nucleotide diversity (*N_d*: 0.0313) is lower than that in northeast Asia, but still higher compared to Europe.

Statistically significant negative Tajima's *D* values were found in all populations except for Alaska, which had a positive value, but this should be interpreted with caution due to the small number of samples. The Fu's *F_s* values were all negative, but only the northeast Asian populations were significant (Table 3).

Population differentiation (*F_{ST}*)

The fixation index (*F_{ST}*) was calculated to estimate the population differentiation between the European, the different groups of northeast Asian and Alaskan sequences. The *F_{ST}* values between the European population and all others were significantly high, similar to between Alaska and northeast Asia, which means that the populations are quite separate and there is little genetic exchange (Table 4). In comparison, the *F_{ST}* of Alaska and northeast Asia is considerably smaller at 0.6875, but still represents a

Table 4. *F_{ST}* values between *E. canadensis* G10 populations based on complete *cox1* gene (1608 bp)

	Europe	Sakha	Eastern Russia	Northeast Asia
Europe	-	-	-	-
Sakha	0.6731*	-	-	-
Eastern Russia ^a	0.6349*	-0.0191	-	-
Northeast Asia ^b	0.5666*	-0.0017	-0.0053	-
Alaska (USA)	0.9628*	0.7697*	0.7426*	0.6875*

^aIncluding sequences from Sakha and additional from eastern Russia.

^bIncluding sequences from eastern Russia, Mongolia and Tibet.

*Significant at *P* < 0.01.

clear differentiation. The negative values when comparing the Sakha population with total eastern Russia or total northeast Asia can be considered as zero and show no genetic differentiation.

Discussion

The taxonomic status of the 4 genotypes G6/7, G8 and G10, provisionally grouped under the name *Echinococcus canadensis* (Vuitton *et al.*, 2020) the phylogenetically youngest *Echinococcus* species, is not yet fully resolved. As mentioned above, differences can be seen between the genotypes with respect to life cycle, geography and genetics (Romig *et al.*, 2017). The genotypes 6/7 involve primarily domesticated animals in its cycle, whereas G8 and G10 both are linked to transmission by wild animals (Oksanen and Lavikainen, 2015; Romig *et al.*, 2017). Compared to G6/7, only a few studies were carried out with genotypes 8 and 10, and these only with small sample sizes as acquisition of samples from wild animals is difficult and therefore only a few samples are usually available for evaluation. Even more importantly, the majority of these specimens came from northern Europe, very few from northeast Asia and Alaska. A recent study investigated the genetic diversity of European specimens of *E. canadensis* G8 and G10 using the whole mitochondrial genome and found only a very small number of genetic variants within the 2 genotypes (3 haplotypes among 14 G8 samples and 5 among 15 G10 samples) (Laurimäe *et al.*, 2023). All varied only in 1–3 nucleotide exchanges, except for 1 G10 isolate from Estonia, which had a higher variation with 24 exchanges. Considering that this variation occurred in the entire mitochondrial genome with more than 13 500 bp, the numbers of exchanges are small compared to the results of the Asian samples in the present study. The variation of the *cox1* gene (1608 bp) alone is thus remarkable and unexpected. The largest distance between the haplotypes found in Sakha was 8 nucleotide exchanges. In total, i.e. the 42 sequences analysed here and the 5 additional sequences from the GenBank, which also originate from eastern Russia, 17 gene variants could be detected. Including the sequences from China and Mongolia, 19 haplotypes can be found in 54 northeast Asian sequences. The European *cox1* sequences showed only 3 haplotypes, whereby the above-mentioned Estonian sequence was the most distant with a difference of 4 bases to the major European haplotype. Interestingly, this gene variant was also described from the very east of Sakha (Konyaev *et al.*, 2013); and conversely, the European main variant was also found in 2 intermediate hosts from Sakha. Even as this indicates some minor genetic exchange between Europe and

eastern Asia, it concerns only 3 out of 99 sequences and is therefore at an extremely low level.

In general, there is a surprisingly high genetic variation of G10 in northeast Asia. Even more so when one considers that the 38 worms from the present study came from only 10 wolves. In some wolves, all the worms analysed had the same haplotype, but 6 wolves possessed worms with different gene variants. In 2 cases, each of the 5 isolated worms showed a different haplotype. This suggests a wide distribution and high prevalence of the parasite in the intermediate hosts in this region.

Unfortunately, we could not obtain the complete *cox1* sequence from the remaining isolates despite multiple repetitions. In those cases, DNA isolation using NaOH was followed by the classical digestion and the phenol–chloroform extraction method as well as a commercial extraction kit. In none of the cases could an amplification product be obtained afterwards, even with large *Taenia* worms (results not shown), so it is assumed that the DNA was degraded during storage for unknown reasons. This prevented also the analyses of further genes of the samples with complete *cox1* sequence, since multiple repetitions were needed to obtain this single gene and DNA solutions were depleted in most cases of adult worms. The analyses of other mitochondrial and in particular nuclear genes would have been desirable, as this could have helped to clarify the question of whether it could be justified to split *E. canadensis* into 2 or 3 species.

The results of the present study show that the genetic diversity of *E. canadensis* G10 in Northeast Asia is far higher than elsewhere, particularly when compared to northern Europe. Intriguingly, northeast Asia is also the only region where all 4 genotypes of *E. canadensis* are known to occur. The northernmost identification of a G6 infection was done in Sakha by Konyaev *et al.* (2013) who found a reindeer infected with this genotype. Further south, in Mongolia and the Tibetan Plateau of China, *E. canadensis* G6/7 is common and G10 has also been identified (Ito *et al.*, 2014; Zhang *et al.*, 2014; Yang *et al.*, 2015; Wu *et al.*, 2018). The Tibetan Plateau is the southernmost area where G8 and G10 can still be found in addition to G6/7 (Wu *et al.*, 2018; Hua *et al.*, 2019). Despite this diversity, our data indicate that *E. canadensis* genotypes are not randomly scattered throughout the region, as only G10 was detected among 191 amplifiable worms from 35 wolves, and cysts from 4 cervids in Sakha, while not a single isolate of G8 could be found. We can only speculate about the reasons. In Europe, records of G8 are mainly from more temperate regions, with G10 venturing further north (Laurimäe *et al.*, 2023). Our study area in Sakha boasts some of the lowest temperature records outside Antarctica, with winter temperatures that may fall below -60°C . This certainly puts some stress on a parasite life cycle that includes environmental stages (eggs), but whether G10 is better adapted to such conditions than other genotypes is open to further investigations.

Two of the wolves were infected with *E. multilocularis*, indicating that wolves may be involved in the life cycle of this parasite in Sakha as it has been shown in the subarctic parts of North America (Geszy *et al.*, 2014).

We cannot draw any conclusion on the identity of the *Taenia* species that was present in 21 of the 35 wolves with amplifiable DNA. The sequence did not match any GenBank deposit, and only 95.5% identity rules out conspecificity with *T. multiceps*. The presence of only juvenile worms is likely due to sampling bias, as only *Echinococcus*-sized worms were targeted. Further investigations, also on cysticerci from wolf prey species, are required to identify this obviously cryptic species.

The exploration of the genetic diversity can shed light on the origin of species, as has been shown for *E. granulosus* s.s. Based on differences in haplotype diversity of *E. granulosus* s.s.

from different regions, this parasite is thought to have originated in the Middle East, in the area of the Fertile Crescent. There, the genetic diversity of the parasite is highest, and it decreases with increasing geographical distance (Casulli *et al.*, 2012; Yanagida *et al.*, 2012). As for *E. canadensis*, the genetic haplotype diversity has only been investigated from the G6/7 genotypic cluster. Although some geographical structuring can be seen with the 'G6' variants originating mostly from camel-raising regions and G6/7 contains a large number of haplotypes, most of these cluster closely together suggesting close relationship and recent ancestry (Addy *et al.*, 2017; Laurimäe *et al.*, 2018b). In contrast, many haplotypes of G10 analysed in this study are widely separated from each other as reflected by the far higher value of nucleotide diversity ($N_d = 0.0943$) as compared to G6/7 ($N_d \leq 0.00173$) (Addy *et al.*, 2017).

The remarkably high genetic diversity of G10 in northeast Asia, the occurrence of all *E. canadensis* genotypes in this region and the phylogenetically basal position of genotypes 8 and 10 may be circumstantial evidence that this parasite might have evolved in this region.

However, more samples are needed from northern Russia, covering the vast area between Sakha and Europe, and from North America. More importance should be given not only to G10, but also to the analysis of the even more elusive G8, the most basally localized genotype. Assuming our tentative hypothesis that the geographic origin of the entire *E. canadensis* cluster is northeast Asia, a gradual decline in diversity towards the West and East (and to North America) should be observed.

Data availability statement. The authors confirm that the data supporting the findings of this study are available within the article.

Author's contribution. M. W., L. K. and T. R. conceived and designed the study. L. K., I. O. and A. O. conducted the sample collection. M. W., S. L. and J. O. carried out the laboratory analyses. M. W. and F. A. performed the computer analyses. All authors wrote and revised the manuscript.

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