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# **Research Article**

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# Molecular phylogeny based on six nuclear genes suggests that *Echinococcus granulosus*

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sensu lato genotypes G6/G7 and G8/G10 can be regarded as two distinct species

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# Abstract

Tapeworms of the species complex of Echinococcus granulosus sensu lato (s. l.) are the cause of a severe zoonotic disease - cystic echinococcosis, which is listed among the most severe parasitic diseases in humans and is prioritized by the World Health Organization. A stable taxonomy of E. granulosus s. l. is essential to the medical and veterinary communities for accurate and effective communication of the role of different species in this complex on human and animal health. E. granulosus s. l. displays high genetic diversity and has been divided into different species and genotypes. Despite several decades of research, the taxonomy of E. granulosus s. l. has remained controversial, especially the species status of genotypes G6-G10. Here the Bayesian phylogeny based on six nuclear loci (7387 bp in total) demonstrated, with very high support, the clustering of G6/G7 and G8/G10 into two separate clades. According to the evolutionary species concept, G6/G7 and G8/G10 can be regarded as two distinct species. Species differentiation can be attributed to the association with distinct host species, largely separate geographical distribution and low level of cross-fertilization. These factors have limited the gene flow between genotypic groups G6/G7 and G8/G10, resulting in the formation of distinct species. We discuss ecological and epidemiological differences that support the validity of these species.

## Introduction

Tapeworms belonging to the species complex of *Echinococcus granulosus sensu lato* (*s. l.*) have been identified as the cause of a chronic zoonotic disease known as cystic echinococcosis (CE), a disease that has considerable impact on both livestock and human health worldwide (Craig *et al.*, 2017; WHO, 2017). The general life cycle of *E. granulosus s. l.* involves various carnivores as definitive hosts for the adult stage, including mostly dogs in both rural and urban areas, and wolves (Moks *et al.*, 2006; Schurer *et al.*, 2014; Laurimaa *et al.*, 2015*a*; Thompson, 2017). Both domesticated and wild large mammalian herbivores act as intermediate hosts for the larval stage. The larval stage is in the form of hydatid cysts that are predominantly located in the liver and/or lungs of the intermediate hosts. Cysts can be fertile or sterile, depending on presence or absence of protoscoleces, respectively. While these cysts can cause significant health problems for the infected intermediate hosts, the infection in the definitive host is usually asymptomatic (Thompson, 2017).

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The taxonomy of E. granulosus s. l. has been a challenging issue for decades. It is well established that this parasite complex displays high genetic diversity and on the basis of mitochondrial DNA (mtDNA) it has been divided into eight different genotypes (G1, G3, G4-G8; and G10; Bowles et al., 1992, 1994; Lavikainen et al., 2003; Kinkar et al., 2017). Several of these recognized mitochondrial genotypes have differences in their lifecycles, hosts ranges and morphology (Thompson and McManus, 2002; Romig et al., 2017; Thompson, 2017). These differences have provided grounds to consider some of these genotypes as distinct species: G1 and G3 as E. granulosus sensu stricto (s. s.; Kinkar et al., 2017), G4 as Echinococcus equinus and G5 as Echinococcus ortleppi (Thompson and McManus, 2002; Lymbery, 2017). The analytical power has been low in most studies as the analyses have been based largely on short sequences of mtDNA, most often on a fragment of a single gene (e.g. Casulli et al., 2012; Andresiuk et al., 2013). Recent studies based on considerably longer mtDNA sequences (Kinkar et al., 2016, 2018a, 2018b; Laurimäe et al., 2016) have yielded significantly deeper insight into the phylogeny and phylogeography of different genotypes. For example, using sequences of nearly complete mitochondrial genomes and three nuclear genes, Kinkar et al. (2017) have revised the status of E. granulosus s. s. and demonstrated that genotypes G1 and G3 are distinct mitochondrial genotypes, whereas G2 is not a separate genotype or even a monophyletic cluster, but belongs to G3. On the other hand, nuclear data revealed no genetic separation of G1 and G3, suggesting that these genotypes form a single species due to ongoing gene flow. The authors concluded that in the taxonomic sense, genotypes G1 and G3 can be treated as a single species E. granulosus s. s., and that G1 and G3 should be regarded as distinct genotypes only in the context of mitochondrial data, whereas G2 was recommended to be excluded from the genotype list (Kinkar et al., 2017). A recently discovered isolate from Ethiopia is tentatively retained in E. granulosus s. s. as a genotype distant from G1/G3 awaiting taxonomic positioning (Wassermann et al., 2016). In contrast, the species status of E. equinus (G4), E. ortleppi (G5) and Echinococcus felidis has, to date, remained undisputed (Hüttner et al., 2008; Thompson, 2008; Saarma et al., 2009; Knapp et al., 2011; Romig et al., 2015; Lymbery, 2017).

The species status of the four E. granulosus s. l. genotypes G6, G7, G8 and G10, however, has remained uncertain. In addition to genetic differences, there are also various ecological and epidemiological differences between these mitochondrial genotypes. Genotypes G6 and G7 are known to be perpetuated predominantly in a domestic life cycle involving goats, camels or pigs as intermediate hosts, and dogs as definitive hosts; however, a recent study found members of the G6/G7 cluster to be widespread in wild mammals of southern Africa (Romig et al., 2017). Genotypes G8 and G10 are mostly circulating in a sylvatic cycle with moose and reindeer acting as intermediate hosts, and wolves as definitive hosts. Moreover, these four genotypic groups are largely allopatric. The distribution range of G6 and G7 covers more southern areas such as Western Europe, the Mediterranean area, Africa, South and Central America, and the Middle East (e.g. Varcasia et al., 2006, 2007; Lymbery et al., 2015). Genotypes G8 and G10 have been found to coexist in the northern hemisphere - mostly in northern part of Europe (e.g. Estonia, Finland, Sweden and Latvia), Northern Asia and Canada (Moks et al., 2006, 2008; Konyaev et al., 2013; Schurer et al., 2014; Marcinkute et al., 2015; Oksanen and Lavikainen, 2015).

Previous studies have mostly focused on mtDNA to resolve the phylogeny and taxonomic status of genotypes G6–G10 (e.g. Moks *et al.*, 2008; Nakao *et al.*, 2013; Addy *et al.*, 2017). These studies demonstrated that the cervid genotype G10 is a sister taxon to the camel–pig genotypes G6/G7, rather than assuming a sister

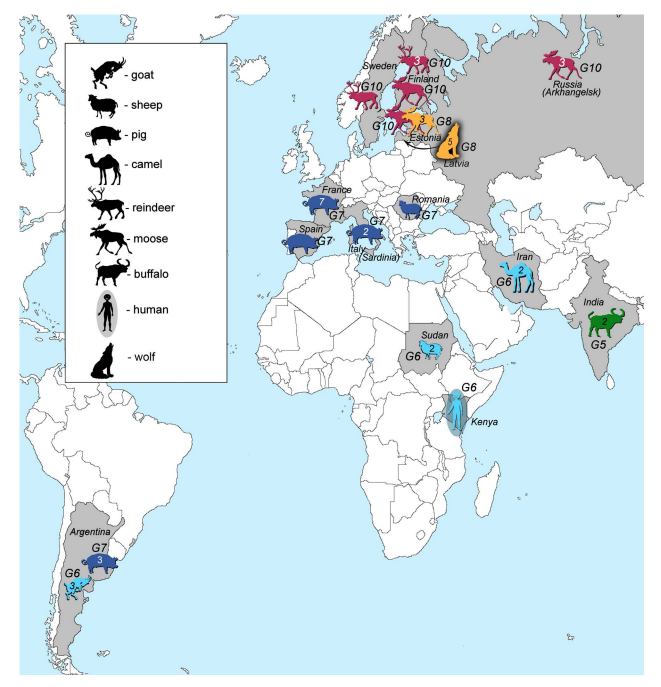
position with the other cervid genotype G8. It was therefore suggested to combine G6-G10 into a single species which, in terms of priority, should be E. canadensis (Nakao et al., 2007; Hüttner et al., 2008). Moreover, the mitochondrial studies placed E. multilocularis in the midst of the E. granulosus genotypes, rendering the E. granulosus complex paraphyletic and contradicting the classical taxonomy of the genus (reviewed in Knapp et al., 2015). Although mtDNA sequences are widely used and various parasite identification methods have been developed based on these (e.g. Boubaker et al., 2013; Laurimaa et al., 2015b) one has to be cautious when interpreting the results. MtDNA represents the evolution of the maternal linage, which can have different trajectories than that of the species. As argued in Saarma et al. (2009), once a new mtDNA mutation becomes fixed in a population, the new mitochondrial lineage separates from the ancestral one; from this point onwards, mutations continue to fix progressively in an independent manner in both the new and ancestral mitochondrial lineages, and mitochondrial lineages continue to diverge. However, this does not mean that these separate mitochondrial lineages have necessarily become separate biological entities - genetic exchange between different taxa can only be assessed with nuclear markers. Thus, it was clear that nuclear data are needed to clarify the taxonomy of the genus Echinococcus. Indeed, a phylogeny radically different from that of mtDNA data was inferred by using sequences of five nuclear genes (5086 bp in total); this analysis placed G8 and G10 as sister taxa, and E. multilocularis clearly separate from the E. granulosus s. l. complex (Saarma et al., 2009). However, in this work G6/G7 were represented by isolates from cattle and pig, and it was not evident which of these two genotypes these isolates belonged to (probably G7). Since the clear distinction between G6 and G7 was not made in this study, the exact phylogenetic relations between G6 and G7 remained obscure. On the other hand, the analysis performed by Knapp et al. (2011) based on a different set of nuclear loci suggested (in line with the mtDNA data), that E. granulosus s. l. complex could be paraphyletic. Unfortunately, this study did not include G10 and therefore the exact phylogenetic relations in the G6-G10 group remained unresolved. Thus, despite numerous attempts to revise the phylogeny and taxonomy of genotypes G6-G10, no consensus has been reached. Some authors have proposed to treat G6-G10 provisionally as a single species E. canadensis awaiting further data from the nuclear genome (e.g. Nakao et al., 2007; Moks et al., 2008; Nakao et al., 2013; Romig et al., 2015; Addy et al., 2017), while others as two distinct species: G6/G7 as E. intermedius and G8/G10 as E. canadensis (Thompson, 2008; Saarma et al., 2009) or even as three species: G6/G7 as E. intermedius, G8 as E. borealis and G10 as E. canadensis (Lymbery et al., 2015).

The main aim of this study was to use a more extensive range of nuclear loci and include all four genotypes (G6–G8 and G10) in a phylogenetic analysis to resolve their taxonomic status.

#### **Materials and methods**

#### Parasite material

Samples of *E. granulosus* genotypes G5–G10 used in this study (41 in total) originated from various regions and intermediate or final hosts (Fig. 1, Table 1). Genotype G5 was represented by two samples from India. Samples of genotype G6 (n = 8) were from three continents: South America (Argentina, n = 3), Africa (Kenya, n = 1; Sudan, n = 2) and Eurasia (Iran, n = 2). Samples of genotype G7 (n = 14) were from South America (Argentina, n = 3), and Eurasia (Spain, n = 1; France, n = 7; Italy, n = 2; Romania, n = 1). Samples that belonged to genotype G8 (n = 8) originated from Eurasia (Estonia, n = 3; Latvia, n = 5), whereas



**Fig. 1.** Geographic locations and host species (intermediate or final) of all of the analysed samples in this study. Numbers inside the animal figures stand for the number of samples collected. Green colour represents *E. ortleppi* (G5) samples, cyan genotype G6 samples, dark blue genotype G7 samples, orange G8 and pink G10 samples.

genotype G10 was represented by nine samples from Eurasia (Sweden, n = 1; Finland, n = 4; Russia, n = 3; Estonia, n = 1). DNA of the Finnish, Swedish and Russian specimens were provided by Antti Lavikainen. These specimens were already used and their genotypic identities defined in previously published studies (Lavikainen *et al.*, 2006; Nakao *et al.*, 2013). Samples were ethanol-preserved at -20 °C until further use.

# PCR amplification and sequencing of six nuclear loci

High Pure PCR Template Preparation Kit (Roche Diagnostics, Mannheim, Germany) was used to extract DNA from either protoscoleces or cyst membranes, following the manufacturer's protocols. Six nuclear genes were chosen for PCR amplification and sequencing: transforming growth factor beta receptor kinase (*tgf*; 1137 bp), calreticulin (*cal*; 1138 bp), elongation factor 1 alpha (*ef1*; 1055 bp), ezrin–radixin–moesin-like protein (*elp*; 780 bp), phosphoenolpyruvate carboxykinase (*pepck*; 1506 bp) and DNA polymerase delta (*pold*; 1771 bp). For further details on cycle parameters for PCR and sequencing see Saarma *et al.* (2009; *tgf*, *cal*, *ef1*, *elp*) and Knapp *et al.* (2011; *pepck*, *pold*). Nuclear sequences were deposited in GenBank (MG766944–MG767169). Consensus sequences were assembled using Codon Code Aligner v6.0.2. BioEdit v7.2.5 (Hall, 1999) was used for multiple alignments with Clustal W and for manual correction of sequences.

As polymorphic sites where the same mutations are shared between genotypes G6 and G10 have been found in *pepck* and *pold* genes (Yanagida *et al.*, 2017), we also checked our aligned sequences for polymorphic sites that discriminate between

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|---|--|-----------------|------------------|-------------------|----------------------------|-------------------------|------------------|-------------------------|-------------------------|------------------------------|----------------------|
| (6)         (6) <th>Sample no.</th> <th>nDNA</th> <th>Genotype</th> <th>Host</th> <th>Origin</th> <th>Accession no <i>ef1</i></th> <th>Accession no cal</th> <th>Accession no <i>tgf</i></th> <th>Accession no <i>elp</i></th> <th>Accession no<br/><i>pepck</i></th> <th>Accession no<br/>pold</th>  | Sample no.   | nDNA            | Genotype         | Host              | Origin                     | Accession no <i>ef1</i> | Accession no cal | Accession no <i>tgf</i> | Accession no <i>elp</i> | Accession no<br><i>pepck</i> | Accession no<br>pold |
| (5)         (5)         (10)         (   | 1  | G5              | G5               | Buffalo           | India                      | MG766944                | MG766984         | MG767024                | MG767064                | MG767098                     | MG767135             |
| (6) $(6)$ <t< td=""><td>2</td><td>G5</td><td>G5</td><td>Buffalo</td><td>India</td><td>MG766945</td><td>MG766985</td><td>MG767025</td><td>MG767065</td><td>MG767099</td><td>MG767136</td></t<>   | 2  | G5              | G5               | Buffalo           | India                      | MG766945                | MG766985         | MG767025                | MG767065                | MG767099                     | MG767136             |
| (6)         (6)         (7)         (7)         (7)         (7)         (7)         (7)           (6)   | 3  | G6              | G6               | Camel             | Iran, Isfahan              | MG766946                | MG766986         | MG767026                | I                       | 1                            | MG767137             |
| (6)         (6)         (60)         Agentin, Nouquen         (67603)         (67703)         (67703)         (67703)           (6)   | 4  | G6              | G6               | Camel             | lran, Isfahan              | MG766947                | MG766987         | MG767027                | MG767066                | MG767100                     | MG767138             |
| (6) $(6)$ <t< td=""><td>5</td><td>G6</td><td>G6</td><td>Goat</td><td>Argentina, Neuquen</td><td>MG766948</td><td>MG766988</td><td>MG767028</td><td>MG767067</td><td>MG767101</td><td>MG767139</td></t<>   | 5  | G6              | G6               | Goat              | Argentina, Neuquen         | MG766948                | MG766988         | MG767028                | MG767067                | MG767101                     | MG767139             |
| 6c $6c$ $acentha, Neudration         a756950 a77030 a77030 a77030 6c 6c Human ecyn, Tuchana         a767632 a77732 a77732 6c 5rep 5rehn 5rehn 5rehn a76732 a77732 a77732 6c 5rep 5rehn a766432 a76732 a75732 a75732 7c 7c Propolicin         a766932 a767332 a75732 a75732 7cr 7cr Propolicin         a766932 a776332 a75732 a75732 7cr 7cr 7cr a776332 a776332 a75732 a75732 7cr 7cr 7cr a776332 a7766332 a77732 a 775732 7cr 7cr 7cr a776932 a776932 a 77732 7cr 7cr a776332 a 776632 a 77732 a 77732 7cr 7cr 7cr a7$   | 9  | G6              | G6               | Goat              | Argentina, Neuquen         | MG766949                | MG766989         | MG767029                | MG767068                | MG767102                     | MG767140             |
| 66         66         Human         Kenya, Tukkaa         M 676691         M 676703 $-$ 166         66         Shep         Sudan, Nyala         M 676692         M 677032 $-$ 166         66         Shep         Sudan, Nyala         M 676695         M 677032 $-$ 176         7         Pig         Agentia, Buenos         M 676695         M 677033         M 677013           176         7         Pig         Agentia, Buenos         M 676695         M 676703         M 677013           176         7         Pig         Agentia, Buenos         M 76695         M 676703         M 677013           176         7         Pig         Agentia, Buenos         M 76695         M 676703         M 676703           176         7         Pig         M 76695         M 676703         M 676703         M 676703           176         7         Pig         M 766695         M 676704         M 676703         M 676703           176         17         Pig         M 766695         M 676704         M 676703         M 676703           176         17         17695         M 766695         M 6767040         M 676703         M 676704<  | 7  | G6              | G6               | Goat              | Argentina, Neuquen         | MG766950                | MG766990         | MG767030                | MG767069                | MG767103                     | MG767141             |
| 66 $66$ $8uedan$ , $Nabla$ $M076693$ $M077032$ <th< td=""><td>8</td><td>G6</td><td>G6</td><td>Human</td><td>Kenya, Turkana</td><td>MG766951</td><td>MG766991</td><td>MG767031</td><td>I</td><td>MG767104</td><td>I</td></th<>   | 8  | G6              | G6               | Human             | Kenya, Turkana             | MG766951                | MG766991         | MG767031                | I                       | MG767104                     | I                    |
| (6) $(6)$ $(8)$ $(8)$ $(8,76)$ $(8,76)$ $(8,76)$ $(8,76)$ $(7)$ $(7)$ $(7)$ $(7)$ $(7)$ $(7,70)$ $(7,70)$ $(7,70)$ $(7)$ $(7)$ $(7)$ $(7)$ $(7,70)$ $(7,70)$ $(7,70)$ $(7,70)$ $(7)$ $(7)$ $(7)$ $(7,70)$ $(7,70)$ $(7,70)$ $(7,70)$ $(7)$ $(7)$ $(7)$ $(7,70)$ $(7,70)$ $(7,70)$ $(7,70)$ $(7,7)$ $(7,7)$ $(7,70)$ $(7,70)$ $(7,70)$ $(7,70)$ $(7,70)$ $(7,7)$ $(7,7)$ $(7,7)$ $(7,70)$ $(7,70)$ $(7,70)$ $(7,70)$ $(7,7)$ $(7,7)$ $(7,7)$ $(7,7)$ $(7,70)$ $(7,70)$ $(7,70)$ $(7,7)$ $(7,7)$ $(7,7)$ $(7,7)$ $(7,7)$ $(7,70)$ $(7,70)$ $(7,7)$ $(7,7)$ $(7,7)$ $(7,7)$ $(7,7)$ $(7,7)$ $(7,70)$ $(7,7)$ <td>6</td> <td>G6</td> <td>G6</td> <td>Sheep</td> <td>Sudan, Nyala</td> <td>MG766952</td> <td>MG766992</td> <td>MG767032</td> <td>I</td> <td>MG767105</td> <td>MG767142</td>  | 6  | G6              | G6               | Sheep             | Sudan, Nyala               | MG766952                | MG766992         | MG767032                | I                       | MG767105                     | MG767142             |
| Gfe         Gf         Pig         Agentina, Buenos         Gr6695         MG76034         MG   | 10   | G6              | G6               | Sheep             | Sudan, Nyala               | MG766953                | MG766993         | MG767033                | MG767070                | MG767106                     | MG767143             |
| GTe         Fig.         Agentia         Increases         Incres         Incres  | 11   | G7e             | 67               | Pig               | Argentina, Buenos<br>Aires | MG766954                | MG766994         | MG767034                | MG767071                | MG767107                     | MG767144             |
| 67c $67$ $8c$ <  | 12   | G7e             | G7               | Pig               | Argentina                  | MG766955                | MG766995         | MG767035                | MG767072                | MG767108                     | MG767145             |
| GTC         Fig.         Central Spain, Segoira         MG76697         MG76703         MG76703           GTd         GT         Sheep         Romaina         MG76696         MG76703         MG76703           GTd         GT         Pig         Ruy, Sardinia         MG76696         MG76703         MG76703           GTd         GT         Pig         Ruy, Sardinia         MG76700         MG76704         MG76704           GTd         GT         Pig         Ruy, Sardinia         MG76704         MG76704         MG76704           GTd         GT         Pig         Fance, Corsica         MG76704         MG76704         MG7707           GTd         GT         Pig         Fance, Corsica         MG76702         MG76702         MG76704         MG7707           GTd         GT         Pig         Fance, Corsica         MG76702         MG76704         MG7707         MG7707           GTd         GT         Pig         Fance, Corsica         MG76902         MG76704         MG7707         MG7707           GTd         GT         Pig         Fance, Corsica         MG76902         MG76704         MG7707         MG7707           GTd         GT         Pig         Fance, C   | 13   | G7c             | G7               | Pig               | Argentina, Neuquen         | MG766956                | MG766996         | MG767036                | I                       | MG767109                     | MG767146             |
| GTd         GT         Sheep         Romaria         MG76658         MG76036         MG76703         -           GTd         GT         Pig         taly, Sardinia         MG76909         MG76704         MG76704         MG76707           GTd         GT         Pig         taly, Sardinia         MG76504         MG76704         MG76707         MG76707           GTd         GT         Pig         Fance, Corsica         MG76504         MG76704         MG76707         MG76707           GTd         GT         Pig         Fance, Corsica         MG766963         MG76704         MG76707         MG76707           GTd         GT         Pig         Fance, Corsica         MG766963         MG76706         MG76707         MG76707           GTd         GT         Pig         Fance, Corsica         MG766963         MG76706         MG76707         MG76708           GTd         GT         Pig         Fance, Corsica         MG766963         MG76706         MG76708         MG76708           GTd         GT         Pig         Fance, Corsica         MG766963         MG76706         MG76708         MG76708           GTd         GT         Pig         Fance, Corsica         MG766964  | 14   | G7c             | G7               | Pig               | Central Spain, Segovia     | MG766957                | MG766997         | MG767037                | MG767073                | MG767110                     | MG767147             |
| GTd         GT         Pig         Italy, Sardinia         MG76059         MG76039         MG76103         MG76104           GTd         GT         Pig         Italy, Sardinia         MG76906         MG76100         MG76109         MG76107           GTd         GT         Pig         Italy, Sardinia         MG76961         MG76101         MG76101         MG76102           GTd         GT         Pig         Fance, Corsica         MG76961         MG76102         MG76103         MG76103           GTd         GT         Pig         Fance, Corsica         MG769612         MG76102         MG76104         MG76108           GTd         GT         Pig         Fance, Corsica         MG76051         MG76102         MG76103         MG76103           GTd         GT         Pig         Fance, Corsica         MG76056         MG76106         MG76108         MG76108           GTd         GT         Pig         Fance, Corsica         MG760505         MG76104         MG76108         MG76108           GTd         Pig         Fance, Corsica         MG766965         MG76106         MG76104         MG76108           GTd         Pig         Fance, Corsica         MG766965         MG76106  | 15   | G7d             | G7               | Sheep             | Romania                    | MG766958                | MG766998         | MG767038                | I                       | I                            | I                    |
| (7d) $(7)$ <t< td=""><td>16</td><td>G7d</td><td>G7</td><td>Pig</td><td>Italy, Sardinia</td><td>MG766959</td><td>MG766999</td><td>MG767039</td><td>MG767074</td><td>MG767111</td><td>MG767148</td></t<>  | 16   | G7d             | G7               | Pig               | Italy, Sardinia            | MG766959                | MG766999         | MG767039                | MG767074                | MG767111                     | MG767148             |
| GTd         G7         Pig         France, Corsica         MG76501         MG76701         MG76704         MG76704           G7d         G7         Pig         France, Corsica         MG765063         MG76703         MG767043         MG767043           G7d         G7         Pig         France, Corsica         MG765063         MG767043         MG767043         MG767043           G7d         G7         Pig         France, Corsica         MG765064         MG767040         MG767043         MG767043           G7d         G7         Pig         France, Corsica         MG765064         MG767040         MG767043         MG767043           G7d         G7         Pig         France, Corsica         MG766064         MG767040         MG767043         MG767043           G7d         G7         Pig         France, Corsica         MG766064         MG767040         MG767043         MG767043           G7d         G7         Pig         France, Corsica         MG766064         MG767040         MG767043         MG767043           G7d         G7         MG766061         MG767004         MG767049         MG767043         MG767043           G8         G8         Moose         Estonia         MG7669   | 17   | G7d             | G7               | Pig               | Italy, Sardinia            | MG766960                | MG767000         | MG767040                | MG767075                | MG767112                     | MG767149             |
| G7d $F_1$   | 18   | G7d             | G7               | Pig               | France, Corsica            | MG766961                | MG767001         | MG767041                | MG767076                | MG767113                     | MG767150             |
| G7d $G7$ $Pig$ $Fance, Corsica$ $MG76903$ $MG76704$ $MG76704$ $MG76704$ $G7d$ $G7d$ $Pig$ $Fance, Corsica$ $MG768964$ $MG76704$ $MG76704$ $MG76704$ $G7d$ $G7d$ $Pig$ $Fance, Corsica$ $MG76066$ $MG76706$ $MG76704$ $MG76708$ $G7d$ $G7$ $Pig$ $Fance, Corsica$ $MG76066$ $MG76706$ $MG767046$ $MG76708$ $G7d$ $G7$ $Pig$ $Fance, Corsica$ $MG76606$ $MG76706$ $MG76704$ $MG76708$ $G7d$ $G7$ $Pig$ $Fance, Corsica$ $MG76606$ $MG76706$ $MG76708$ $MG76708$ $G7d$ $G8$ $Moose$ $Etonia$ $MG76708$ $MG76708$ $MG76708$ $MG76708$ $G8$ $G8$ $Moose$ $Etonia$ $MG76609$ $MG76708$ $MG76704$ $MG76708$ $G8$ $G8$ $Moose$ $Etonia$ $MG76609$ $MG76709$ $MG76704$ $MG76708$ $G8$ $G8$ $Moose$ $Etonia$ $MG76697$ $MG76709$ $MG76704$ $MG76708$ $G8$ $G8$ $Woff$ $Latuia$ $MG76697$ $MG76709$ $MG76704$ $MG76708$ $G8$ $G8$ $Woff$ $Latuia$ $MG76097$ $MG76704$ $MG76708$ $MG76708$ $G8$ $G8$ $Woff$ $Latuia$ $MG76097$ $MG76704$ $MG76708$ $MG76708$ $G8$ $G8$ $Woff$ $Latuia$ $MG76097$ $MG76709$ $MG76708$ $MG76708$ $G8$ $G8$  | 19   | G7d             | G7               | Pig               | France, Corsica            | MG766962                | MG767002         | MG767042                | MG767077                | MG767114                     | I                    |
| GTd         GT         Pig         France, Corsica         MG F6666         MG F6704         MG 76704         MG 76709           GTd         GT         Pig         France, Corsica         MG 766665         MG 76706         MG 76709         MG 76708           GTd         GT         Pig         France, Corsica         MG 766965         MG 767006         MG 76708         MG 76708           GTd         GT         Pig         France, Corsica         MG 766966         MG 767007         MG 76708         MG 76708           GTd         GT         Pig         France, Corsica         MG 766967         MG 76709         MG 76708         MG 76708           GTd         Mose         Estonia         MG 766969         MG 76709         MG 76708         MG 76708           GTd         Mose         Estonia         MG 766969         MG 76709         MG 76708         MG 76708           GTd         GT         Mose         Estonia         MG 766970         MG 76709         MG 76708           GTd         GT         MG 766910         MG 767010         MG 76709         MG 76708         MG 76708           GTd         GT         Wose         Estonia         MG 766910         MG 767010         MG 77008         <   | 20   | G7d             | G7               | Pig               | France, Corsica            | MG766963                | MG767003         | MG767043                | MG767078                | MG767115                     | MG767151             |
| GTd         GT         Pig         France, Corsica         MG766965         MG767005         MG767096         MG767080           GTd         GT         Pig         France, Corsica         MG766965         MG767007         MG767046         MG767080           GTd         GT         Pig         France, Corsica         MG766965         MG767007         MG767080         MG767082           G8a         G8         Moose         Estonia         MG766969         MG767009         MG767082         MG767082           G8a         G8         Moose         Estonia         MG766910         MG767010         MG767083         MG767083           G8a         G8         Moose         Estonia         MG766910         MG767010         MG767083         MG767083           G8a         G8         Wolf         Latvia         MG766910  | 21   | G7d             | G7               | Pig               | France, Corsica            | MG766964                | MG767004         | MG767044                | MG767079                | MG767116                     | MG767152             |
| GTd         GT         Pig         France, Corsica         MG766966         MG767006         MG767046         MG767081           GTd         GT         Pig         France, Corsica         MG766966         MG767007         MG767046         MG767082           GRa         G8         G8         Moose         Estonia         MG766968         MG767008         MG767048         MG767082           G8a         G8         Moose         Estonia         MG766969         MG767009         MG767049         MG767083           G8a         G8         Moose         Estonia         MG766969         MG767009         MG767099         MG767083           G8a         G8         Wolf         Latvia         MG766971         MG767010         MG767085         MG767085           G8a         G8         Wolf         Latvia         MG766972         MG767012         MG767086         MG767086           G8a         G8         Wolf         Latvia         MG766973         MG767012         MG767085         MG767086           G8a         G8         Wolf         Latvia         MG766973         MG767012         MG767082         MG767086           G8a         Volf         Latvia         MG766973 <t< td=""><td>22</td><td>G7d</td><td>G7</td><td>Pig</td><td>France, Corsica</td><td>MG766965</td><td>MG767005</td><td>MG767045</td><td>MG767080</td><td>MG767117</td><td>MG767153</td></t<>  | 22   | G7d             | G7               | Pig               | France, Corsica            | MG766965                | MG767005         | MG767045                | MG767080                | MG767117                     | MG767153             |
| G7d         G7         Pig         France, Corsica         MG 766967         MG 767007         MG 767042         MG 767082           G8a         G8         Moose         Estonia         MG 766969         MG 767009         MG 767049         MG 767083           G8a         G8         Moose         Estonia         MG 766969         MG 767009         MG 767049         MG 767084           G8a         G8         Moose         Estonia         MG 766969         MG 767010         MG 767084         MG 767084           G8a         G8         Work         Estonia         MG 766970         MG 767010         MG 767084         MG 767084           G8a         G8         Work         Latvia         MG 766971         MG 767011         MG 767085         MG 767085           G8a         G8         Work         Latvia         MG 766972         MG 767011         MG 767085         MG 767085           G8a         G8         Work         Latvia         MG 766973         MG 767012         MG 767085         MG 767085           G8a         Work         Latvia         MG 766973         MG 767012         MG 767085         MG 767085           G8a         Work         Latvia         MG 766973         MG 76702   | 23   | G7d             | G7               | Pig               | France, Corsica            | MG766966                | MG767006         | MG767046                | MG767081                | MG767118                     | MG767154             |
| G8a         G8         Moose         Estonia         MG766968         MG767008         MG767048         MG767038           G8a         G8         Moose         Estonia         MG766969         MG767009         MG767049         MG767084           G8a         G8         Moose         Estonia         MG766969         MG767010         MG767039         MG767084           G8a         G8         Worle         Latvia         MG766971         MG767010         MG767050         MG767086           G8a         G8         Worle         Latvia         MG766971         MG767010         MG767050         MG767086           G8a         G8         Worle         Latvia         MG766973         MG767013         MG767053         MG767086           G8a         G8         Worle         Latvia         MG766973         MG767013         MG767053         MG767088           G8a         G8         Worle         Latvia         MG766974         MG767013         MG767085           G8a         G8         Worle         Latvia         MG766974         MG767013         MG767053         MG767088           G8a         G8         Worle         Latvia         MG766974         MG767014         MG76705   | 24   | G7d             | G7               | Pig               | France, Corsica            | MG766967                | MG767007         | MG767047                | MG767082                | MG767119                     | MG767155             |
| G8a         G8         Moose         Estonia         MG766969         MG767009         MG767049         MG767084           G8a         G8         Moose         Estonia         MG766970         MG767010         MG767050         MG767085           G8a         G8         Wolf         Latvia         MG766971         MG767011         MG767051         MG767085           G8a         G8         Wolf         Latvia         MG766972         MG767012         MG767085         MG767086           G8a         G8         Wolf         Latvia         MG766973         MG767012         MG767050         MG767086           G8a         G8         Wolf         Latvia         MG766973         MG767013         MG767053         MG767088           G8a         G8         Wolf         Latvia         MG766974         MG767013         MG767053         MG767088           G8a         G8         Wolf         Latvia         MG766974         MG767013         MG767053         MG767088   | 25   | G8a             | G8               | Moose             | Estonia                    | MG766968                | MG767008         | MG767048                | MG767083                | MG767120                     | MG767156             |
| G8a         G8         Moose         Estonia         MG766970         MG767010         MG767050         MG767085           G8a         G8         Wolf         Latvia         MG766971         MG76701         MG767051         MG767086           G8a         G8         Wolf         Latvia         MG766972         MG76701         MG767051         MG767086           G8a         G8         Wolf         Latvia         MG766973         MG767013         MG767053         MG767088           G8a         G8         Wolf         Latvia         MG766973         MG767013         MG767053         MG767088           G8a         G8         Wolf         Latvia         MG766974         MG767013         MG767053         MG767088   | 26   | G8a             | G8               | Moose             | Estonia                    | MG766969                | MG767009         | MG767049                | MG767084                | MG767121                     | MG767157             |
| G8a         G8         Wolf         Latvia         MG766971         MG767011         MG767051         MG767086           G8a         G8         Wolf         Latvia         MG766972         MG767012         MG767052         MG767087           G8a         G8         Wolf         Latvia         MG766973         MG767013         MG767053         MG767088           G8a         G8         Wolf         Latvia         MG766974         MG767013         MG767053         MG767088           G8a         G8         Wolf         Latvia         MG766974         MG767014         MG767053         MG767089  | 27   | G8a             | G8               | Moose             | Estonia                    | MG766970                | MG767010         | MG767050                | MG767085                | MG767122                     | MG767158             |
| G8a         G8         Wolf         Latvia         MG766972         MG767012         MG767052         MG767087           G8a         G8         Wolf         Latvia         MG766973         MG767013         MG767053         MG767088           G8a         G8         Wolf         Latvia         MG766974         MG767014         MG767054         MG767089  | 28   | G8a             | G8               | Wolf              | Latvia                     | MG766971                | MG767011         | MG767051                | MG767086                | MG767123                     | MG767159             |
| G8a         G8         Wolf         Latvia         MG766973         MG767013         MG767053         MG767088           G8a         G8         Wolf         Latvia         MG766974         MG767014         MG767059         MG767089   | 29   | G8a             | G8               | Wolf              | Latvia                     | MG766972                | MG767012         | MG767052                | MG767087                | MG767124                     | MG767160             |
| G8a G8 Wolf Latvia MG766974 MG767014 MG767054 MG767089  | 30   | G8a             | G8               | Wolf              | Latvia                     | MG766973                | MG767013         | MG767053                | MG767088                | MG767125                     | MG767161             |
|   | 31   | G8a             | 68               | Wolf              | Latvia                     | MG766974                | MG767014         | MG767054                | MG767089                | MG767126                     | MG767162             |

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| 32   | G8a   | G8                         | Wolf                                 | Latvia   | MG766975 | MG767015 | MG767055 | MG767090 | MG767127 | MG767163 |
|--|---|----------------------------|--------------------------------------|--|----------|----------|----------|----------|----------|----------|
| 33   | G8b <sup>a</sup>  | G8                         | Moose                                | USA  | I        | I        | I        | I        | FN567995 | FN568366 |
| 34   | GIOd  | G10                        | Moose                                | Estonia  | MG766976 | MG767016 | MG767056 | 1        | I        | 1        |
| 35   | G10c <sup>b</sup>   | G10                        | Moose                                | Finland  | EU834907 | EU834940 | EU834916 | EU834895 | I        | I        |
| 36   | G10c  | G10                        | Reindeer                             | Finland  | MG766977 | MG767017 | MG767057 | MG767091 | MG767128 | MG767164 |
| 37   | G10c  | G10                        | Reindeer                             | Finland  | MG766978 | MG767018 | MG767058 | MG767092 | MG767129 | MG767165 |
| 38   | G10c  | G10                        | Reindeer                             | Finland  | MG766979 | MG767019 | MG767059 | MG767093 | MG767130 | MG767166 |
| 39   | G10c  | G10                        | Reindeer                             | Sweden   | MG766980 | MG767020 | MG767060 | MG767094 | MG767131 | MG767167 |
| 40   | G10c  | G10                        | Moose                                | Russia, Arkhangelsk  | MG766981 | MG767021 | MG767061 | MG767095 | MG767132 | I        |
| 41   | G10c  | G10                        | Moose                                | Russia, Arkhangelsk  | MG766982 | MG767022 | MG767062 | MG767096 | MG767133 | MG767168 |
| 42   | G10c  | G10                        | Moose                                | Russia, Arkhangelsk  | MG766983 | MG767023 | MG767063 | MG767097 | MG767134 | MG767169 |
| <sup>a</sup> Sample sequence<br><sup>b</sup> Sample sequence | es of <i>pepck</i> and<br>s of <i>ef1</i> , <i>cal</i> , tg | 1 <i>pold</i> obtained fro | im the GenBank di<br>from the GenBan | <sup>a</sup> Sample sequences of <i>pepck</i> and <i>pold</i> obtained from the GenBank database (Knapp <i>et al.</i> , 2011).<br><sup>b</sup> Sample sequences of <i>et1, cal, tgf</i> and <i>etp</i> obtained from the GenBank database (Saarma <i>et al.</i> , 2009). |          |          |          |          |          |          |

genotypes G6 and G10, as well as for positions where mutations were shared between these genotypes.

# Bayesian phylogeny

Bayesian phylogenies were constructed for two datasets, both based on six nuclear genes (7387 bp in total): (1) Dataset 1 (a total of 40 sequences): 39 samples of G6–G8 and G10 analysed in this study, and one additional G8 sample from GenBank, originating from the USA (accession numbers for *pepck* and *pold* were FN567995 and FN568366, respectively; Knapp *et al.*, 2011); (2) Dataset 2 (a total of 42 sequences): the same set of samples as in Dataset 1 and two additional sequences of genotype G5.

The best-fit nucleotide substitution model was selected on the basis of BIC (Bayesian Information Criterion) scores using jModelTest 2 (Guindon and Gascuel, 2003; Darriba *et al.*, 2012). Bayesian phylogenetic analysis was performed in BEAST 1.8.4 (Drummond *et al.*, 2012) using StarBeast (Heled and Drummond, 2010). Posterior distributions of parameters were estimated by using the MCMC (Markov Chain Monte Carlo) sampling. Total length of the chain was 10 000 000 and the parameters were logged every 1000 generations. The resulting phylogenetic trees were summarized and annotated using TreeAnnotator 1.8.4 and visualized with FigTree 1.4.3 (http://tree.bio.ed.ac.uk/software/figtree).

# Results

Total length of the alignment based on six nuclear loci was 7387 bp: *ef1* 1055 bp, *cal* 1138 bp, *tgf* 1137 bp, *elp* 780 bp, *pepck* 1506 bp and *pold* 1771 bp. However, a few of the samples did not yield positive results for all analysed nuclear loci, but as BEAST allows analysis with some missing data, these samples were also included in the analysis (Table 1). All of the samples were homozygotes at all six nuclear loci.

Across the six nuclear loci, 12 polymorphic positions were found that discriminated between G6/G7 and G8/G10. However, in the *pepck*, mutations in three positions were shared between two G7 isolates (samples 13 and 14) and G8/G10 isolates. According to GenBank reference FN567995 (Knapp *et al.*, 2011) these positions were: 236; 1435–1436; 1513.

The best-fit nucleotide substitution model used for the nuclear DNA (nDNA) data was GTR + I + G. The Bayesian phylogeny for the Dataset 1 revealed that genotypes G6 and G7 formed one clade, whereas G8 and G10 another (Fig. 2). Posterior probability values for both nodes assigning G6/G7 and G8/G10 into two different clades were very high (1.00). According to the evolutionary (general lineage) species concept, they can be regarded as two distinct species.

Internal nodes for the clade G8/G10 also received high posterior probability values (0.98 and 1.00). It was shown that G8b (the GenBank sample from the USA) was a sister taxon to G10d (Estonia) and that G10c was a sister taxon to the G8b/G10d clade. Additionally, the tree topology indicated that G8a was positioned as a basal taxon in relation to the G10c/G10d/G8b clade. Similarly to G8/G10 clade, the internal nodes for G6/G7 also received high posterior probability values (0.96 and 1.00). The resultant tree topology shows that G6 is a sister taxon to G7e and that G7d is sister to G6/G7e. G7c occupied a basal position inside the G6/G7 clade.

We also performed a phylogenetic analysis for the Dataset 2 (included G5), as well as with only the samples for which all six nuclear loci were sequenced (Table 1). These analyses yielded essentially the same phylogenetic relations between G6 and G10 as with the larger dataset (Fig. 2; Supplementary Fig. S1–S3).

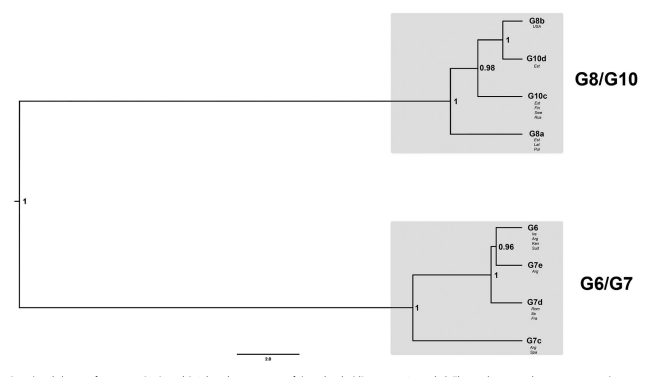


Fig. 2. Bayesian phylogeny of genotypes G6–G8 and G10, based on sequences of six nuclear loci (Dataset 1; 40 samples). The numbers on nodes represent posterior probability values. For further details on the included samples see Table 1.

## Discussion

A stable taxonomy of E. granulosus s. l. is essential to the medical and veterinary communities for accurate and effective communication of the role of different species in this complex on human and animal health. Despite several decades of research, the taxonomy of E. granulosus s. l. has remained controversial and a subject of intense discussion (Saarma et al., 2009; Knapp et al., 2011; Lymbery et al., 2015; Nakao et al., 2015). Most of the studies aiming to resolve the taxonomic status of genotypes G6-G10 have relied on mtDNA (e.g. Lavikainen et al., 2003; Nakao et al., 2007; Hüttner et al., 2008; Moks et al., 2008; Nakao et al., 2013; Addy et al., 2017). However, the mitochondrial genome can only reveal the evolutionary history of the maternal lineage, which can be different to that of the species. For species delimitation, a key component is the analysis of genetic exchangeability, which can be effectively studied only by using various loci from the nuclear genome (Saarma et al., 2009). Until recently, only two studies have analysed multiple nuclear loci to infer the phylogeny of E. granulosus s. l. (Saarma et al., 2009; Knapp et al., 2011), yielding contradictory results. Moreover, both studies did not include all genotypes of the G6-G10 complex.

## Nuclear data and taxonomy of G6-G10

The Bayesian phylogeny based on six nuclear loci clustered the camel-pig genotypes G6/G7 into one clade and the cervid genotypes G8/G10 into another clade (Fig. 2). This result provides strong support for the hypothesis according to which genotypes G6-G10 are divided into two species (Thompson, 2008; Saarma *et al.*, 2009). The internal division of the G6/G7 and G8/G10 clades provides evidence for gene flow between G6 and G7, as well as between G8 and G10, but non-existent or very limited gene flow between genotypic groups G6/G7 and G8/G10. The latter seems to be supported by a recent study based on two nuclear loci, which suggested some degree of gene flow between genotypic groups G6/G7 and G8/G10 (Yanagida *et al.*, 2017); however, their result could be also due to incomplete lineage sorting (see below). Since G6 and G7 are not distinct taxa based on nuclear data (notice in Fig. 2 that G7e forms a subclade with G6, while other isolates of G7 are sister to this), it demonstrates that the gene flow between G6 and G7 has been sufficient to guarantee that G6 and G7 have not diverged from each other. Exactly the same is valid for G8 and G10 (notice in Fig. 2 that G8 and G10 do not form separate subclades, but the isolates of both genotypes are not monophyletic).

Gene flow can occur under conditions of sympatry between both genotypic groups (i.e. G6/G7 and G8/G10). To date, none of the studies have demonstrated sympatry of all these four genotypes. Nevertheless, there are regions where at least some genotypes of these two genotypic groups are potentially sympatric. One such region is in north-eastern Europe, where G8 (this study) and G10 have been recorded from Latvia, and G7 in neighbouring Lithuania (Marcinkute et al., 2015). Considering that wolves (as a main definitive host species for G8/G10) can cover very long distances and their populations in Europe are connected over the distance of more than 800 km (Hindrikson et al., 2017), the possibility for gene flow between G7 and G8/G10 is potentially there, and yet the genotypic groups G6/G7 and G8/G10 are clearly separate on the nuclear phylogeny. Another region of potential sympatry is in eastern Russia, where G6 has been found in relative geographical proximity with G10 (>500 km between the reported cases) (Konyaev et al., 2013; Yanagida et al., 2017). Nevertheless, the phylogeny based on six nuclear loci (current study) shows also that gene flow between genotypic groups G6/G7 and G8/G10 has not been sufficient to merge all four genotypes into a single clade (species). A recent study by Yanagida et al. (2017) based on two nuclear loci (pepck and pold) indicated that some degree of gene flow might occur between G6/G7 and G8/G10 as they found few polymorphic sites where mutations were shared among G6/G7 and G8/G10. Based on this, they suggested that G6-G10 could be considered as one species. However, there were only a limited number of polymorphic characters in the two analysed loci, which may likely

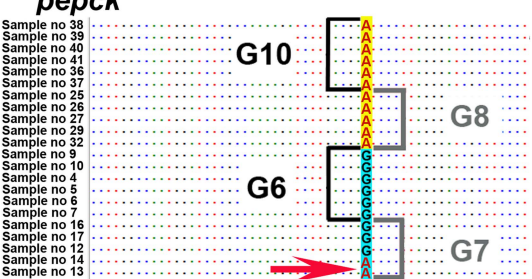
be the reason why their conclusion is not supported by the results of our study. One possible explanation for the shared characters reported in Yanagida et al. (2017) could be due to incomplete lineage sorting, which means that due to the relatively recent evolutionary divergence of G6/G7 and G8/G10, some loci have not had enough time to diverge and as a result there are still shared characters between different genotypes. This is actually evident also from our results. When we examined all six nuclear loci of our study (that include also pepck and pold used by Yanagida et al., 2017), there are several positions in the alignment where the same nucleotide is shared between different genotypic groups. For example, in pepck all isolates of G8 and G10 have A in the shown position in Fig. 3, but remarkably A is also found in two isolates of G7. And yet, despite of some shared mutations between different genotypes, there are a large number of characters specific only either to the genotypic group G6/G7 or to G8/G10, and as a result in the phylogenetic tree the camel-pig genotypes G6/G7 firmly form one clade and the cervid genotypes G8/G10 another (Fig. 2). While we cannot rule out the possibility that to some extent gene flow (hybridization) between these two genotypic groups can occur, as suggested in Yanagida et al. (2017), the nDNA evidence in our study that is based on a larger number of nuclear loci compared with Yanagida and co-authors, clearly shows the phylogenetic division of G6-G10 into two clades, G6/G7 and G8/G10. According to the evolutionary (general lineage) species concept, these two clades can be regarded as distinct species as they represent two distinct evolutionary lineages and other data also support this (see below).

Limited gene flow between species, i.e. hybridization, is in fact relatively common in nature. Possibly the most popular example is the hybridization between wolves and dogs (e.g. Hindrikson et al., 2012; Leonard et al., 2014). In general, it has been estimated that 10-30% of multicellular animal and plant species hybridize regularly (Abbott et al., 2013). Hybridization is also well-known among parasites, it is known for example between different species of helminths (Taenia, Trichinella, Schistosoma, Fasciola, Ascaris) and protozoa (Plasmodium, Leishmania, Toxoplasma and Trypanosoma) (Arnold, 2004; Detwiler and Criscione, 2010; King et al., 2015). Hybridization between closely related tapeworm species in Taeniidae has been demonstrated between T. saginata and T. asiatica (e.g. Okamoto et al., 2010; Yamane et al., 2013). The occurrence of hybridization does not mean that two hybridizing species, if clearly separate on the phylogeny, should therefore be regarded as a single species, it just means that reproductive barrier between species has not yet fully developed.

Although our study did not include samples from the whole geographical range of the genotypes, we argue that this is not a major limitation, since our result shows that even in close geographical proximity these genotypic groups maintain their genetic differences. Moreover, our data included samples from northeastern Europe where genotypes G7, G8 and G10 have been recorded in relatively close geographic areas. A need for including samples from the whole geographical range of the species would have been critical if the genetic data showed no differentiation on a smaller scale, but this was not the case here. Our results demonstrated that gene flow between G6/G7 and G8/G10 genotypic groups in relatively close geographical areas has been insufficient to merge them into a single clade, and instead they form two statistically well supported separate clades (species). One of the possible contributing factors for the limited gene flow between G6/G7 and G8/G10 could also be the reproduction mode of E. granulosus s. l. Although cross-fertilization can occur (e.g. Haag et al., 2011), the main mode of reproduction appears to be self-fertilization (Lymbery, 2017; Thompson, 2017). As the potential for outcrossing between genotypic groups is rare, the evolutionary potential for genetic differentiation and species divergence is high (Lymbery, 2017).

# Ecological, epidemiological and morphological differences of the two species

The division of G6-G10 into two separate species is also supported by other data that can be found in detail in Thompson (2008) and Saarma et al. (2009). Briefly, while G6/G7 is known to be typically circulating in the domestic cycle (camels, goats, pigs and dogs), G8/G10 cycles primarily in the sylvatic cycle, between cervids (moose, elk and reindeer) and wolves (Thompson and McManus, 2002; Lymbery, 2017). Although G6 is commonly involved in a cycle between goats/camels-dogs and G7 mainly pigs-dogs, these two also share some overlap in



### Fig. 3. Nucleotide position on pepck locus, where the same nucleotide A is shared between two samples of genotype G7 and the G8/G10 genotypic group. Depicted position according to FN567995 from the GenBank database is 236 (Knapp et al., 2011). Sample numbers correspond to sample numbers in Table 1.

pepck

their life cycles as both can infect the same intermediate hosts, such as goats and humans (Cardona and Carmena, 2013; Alvarez Rojas et al., 2014; Addy et al., 2017), and definitive host - dogs. Most likely dogs act as vectors for both genotypes, providing opportunities for outcrossing. Moreover, the geographical distribution of G6/G7 is largely different from G8/G10. G6 and G7 are sympatric in Turkey, Argentina and Peru (e.g. Moro et al., 2009; Šnabel et al., 2009; Soriano et al., 2010; Simsek et al., 2011; Lymbery et al., 2015). The recent discovery of the G6/G7 cluster in African wildlife is highly interesting on phylogeographical grounds and is currently further explored (Romig et al., 2017). The cervid strains G8 and G10 are, in contrast, distributed in the northern part of Eurasia and North America (Lavikainen et al., 2003; Thompson et al., 2006; Moks et al., 2008) and so far there are only a few recorded occurrences of G6 in northern latitudes (Konyaev et al., 2013; Yanagida et al., 2017). As G6/G7 circulate primarily in the domestic cycle and G8/G10 in the sylvatic cycle, the probability that parasites from different genotypic groups co-occur in the same definitive host and cross-fertilize is very low. On the other hand, since G6/G7 share the same final host (dog) cross-fertilization has apparently been frequent enough to guarantee that G6/G7 have not diverged. The same is valid for G8/G10, which utilize wild canids (mostly wolves) as definitive hosts. Thus, the association with distinct host species, largely separate geographical distribution and limited rate of cross-fertilization are the main factors that have limited the gene flow between genotypic groups G6/G7 and G8/G10. As a result, these genotypic groups can be regarded as distinct species.

Morphological comparisons of adult worms of G6/G7 and G8/G10 are scarce. Recently, it has been found that genotypes G6 and G7 share similar morphological characteristics, e.g. long terminal segment when compared with the total adult worm length, genital pore is generally anterior in the mature segment and rostellar hook morphometric data have also given similar results for both of these genotypes (Soriano et al., 2016). Based on the limited data, it has been suggested that there are some morphological differences in the reproductive anatomy between G6/G7 and G8/G10, and between rostellar hook morphology (Thompson et al., 2006; Lymbery et al., 2015; Soriano et al., 2016). However, these differences need to be further confirmed as neither direct comprehensive morphological nor extensive ecological comparisons between G6/G7 and G8/G10 have been made so far. Such studies would provide additional valuable information for species delimitation.

Based on priority, the species name for G8/G10 should be *E. canadensis*; however, the species name for G6/G7 warrants further discussions. It has been proposed to use *E. intermedius* for G6/G7 (Thompson, 2008; Saarma *et al.*, 2009); however, this name is highly problematic since the original description by Lopez-Neyra and Soler Planas (1943) did not describe intermediate host and no original type specimen for *E. intermedius* can be found (Nakao *et al.*, 2015).

**Supplementary material.** The supplementary material for this article can be found at https://doi.org/10.1017/S0031182018000719

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