

Letter



Insights into Genetic Diversity and Divergence Time of Human-Derived *Echinococcus granulosus* Isolates in Qinghai, China

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Echinococcosis is a zoonotic disease caused by the larval stages of tapeworms belonging to the genus *Echinococcus*, and it presents substantial global health and economic challenges^[1]. Qinghai Province, situated on the northeastern Qinghai-Tibet Plateau (QTP) with an average altitude exceeding 3,000 m^[2], is characterized by unique environmental features such as hypobaric hypoxia, intense ultraviolet radiation, and low temperatures. These conditions create an environment suitable for the transmission of echinococcosis, rendering the region a highly endemic area. The 2023 National Epidemiological Survey on Echinococcosis in China reported a prevalence of 0.05% for humans, with cystic echinococcosis (CE) accounting for 62.61% of cases^[3]. The prevalence in Qinghai Province is significantly higher than the national average, imposing a heavy burden on both the health of the local population and socioeconomic development of the region.

CE in China is predominant in pastoral and semi-pastoral regions, with the QTP being a highly endemic area, encompassing provinces including Sichuan, Gansu, and Qinghai. Nationwide surveillance shows notable disparities in prevalence across demographic groups, with higher burdens in females, the elderly, Tibetan populations, and herders^[3]. This indicates the high degree of geographical clustering of the disease and its distinct epidemiological characteristics across different populations. In molecular diagnostics, cytochrome c oxidase subunit 1 (COX1) serves as a reliable marker to accurately identify sequence variations among

parasite species, demonstrating high credibility in amplification results^[4]. Therefore, based on molecular genetic analysis of the COX1 gene, the genetic evolutionary patterns of CE in the Qinghai region were systematically investigated in this study.

All samples in this study were collected consecutively from Yushu Prefecture People's Hospital in Qinghai Province, China between 2014 and 2024, comprising 20 confirmed cases of CE that met the inclusion criteria. No artificial selection or exclusion was performed during sample collection. Polymerase chain reaction amplification and sequencing were conducted using the COX1 gene as a molecular marker. The obtained sequences were aligned using NCBI BLAST. Haplotype diversity was analyzed using DnaSP v6, and a haplotype network was constructed using PopArt. Using *Taenia multiceps* as the outgroup, a phylogenetic tree was generated using the maximum likelihood method in IQ-TREE v2.2.0, and node divergence times were estimated using BEAST v2.7.4. To enable an integrative analysis of molecular genetics and geoinformatics, geographical mapping was performed using ArcGIS. All maps were generated using the standard map services provided by the National Platform for Common Geospatial Information Services (<https://www.tianditu.gov.cn/>), with the base map review number GS(2024)0650. The basic geographic boundaries of the base map have not been altered in any way.

Echinococcus DNA was successfully identified in all 20 cyst fluid samples, yielding a COX1 band that was approximately 800 bp in length (Supplementary

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Figure S1). All sequences exhibited 98.35%–99.76% identity with *Echinococcus granulosus* (GenBank ID: MN787531). Demographic analysis of enrolled cases showed that most patients were Tibetan herders. The distribution may be associated with the high density of dogs and livestock in pastoral counties, limited access to clean water, and suboptimal personal hygiene practices among herders. CE prevalence was slightly higher in males than in females, which contrasts with some previously reported findings^[5]. This discrepancy may be attributed to differences in case detection methods. The inclusion criteria of our study resulted in a higher proportion of male patients who, due to their engagement in heavy physical labor, developed more severe complications and sought medical attention. In contrast, most community-based surveys are better equipped to comprehensively capture asymptomatic or mild infections.

Eleven haplotypes of *E. granulosus* were identified in this study (Figure 1). Among them, 7 were found in Yushu Tibetan Autonomous Prefecture. Given the local parasite population with high haplotype diversity (0.905), nucleotide diversity of 0.03035, and genetic background characterized by 182 variable sites in our study, we hypothesized that Yushu Prefecture may serve as an important center for the genetic differentiation of *E. granulosus* in the Qinghai region. In this study, EgH4 was determined to be the dominant haplotype (accounting for 25%, 5/20), followed by EgH1 (accounting for 20%, 4/20),

and then EgH5 and EgH8 (each accounting for 10%, 2/20). The remaining haplotypes accounted for 5% each. All 11 *E. granulosus* haplotypes clustered into a single clade together with sheep- and cattle-derived *E. granulosus* isolates retrieved from GenBank (Figure 2), indicating that *E. granulosus* in Qinghai Province belongs to the typical pastoral type and shares a common evolutionary origin with strains from livestock hosts. The GenBank accession numbers used in Figure 2 of this study are listed in Supplementary Table S1. Comparisons with isolates from surrounding regions indicated that the diversity indices reported in some areas of Xizang were similar to those obtained in this study^[6], whereas in parts of western Sichuan, the diversity tended to be higher^[7]. This geographical gradient may be attributable to the Qinghai population acting as a transitional zone for dispersal from Xizang to Sichuan, with its genetic composition incorporating lineages from different sources.

Based on divergence time estimation (Figure 3), the split between *E. granulosus* and *E. shiquicus* occurred approximately 2.51 million years ago (Mya; 95% CI: 2.23–2.79 Mya). This time point closely coincides with the onset of intensified global glaciation^[8]. Climatic changes directly influence the lineage evolution of the primary hosts of these parasites, thereby indirectly driving adaptive diversification in the parasites themselves. Given this association and the temporal congruence, we propose that the cascade of

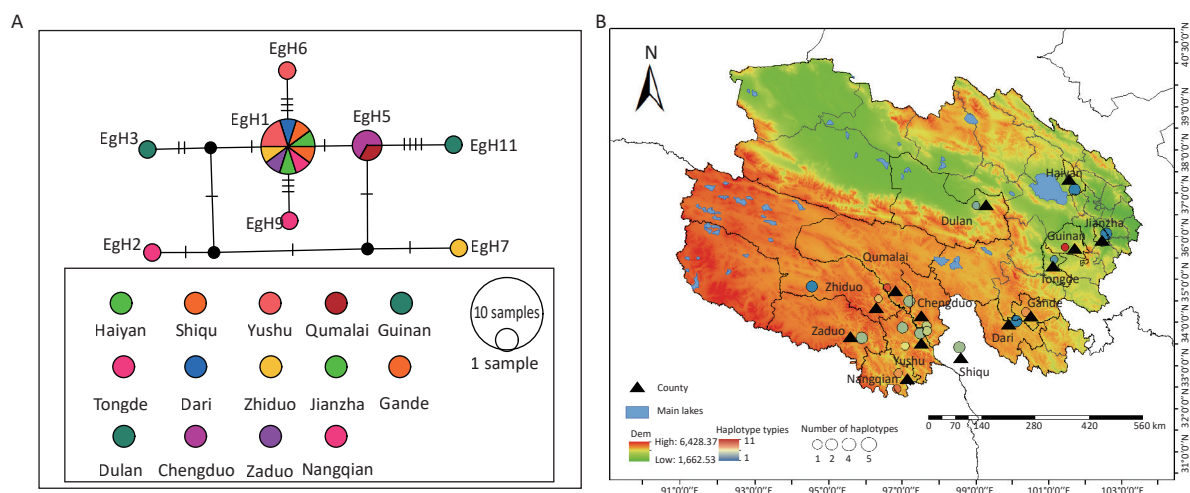


Figure 1. Eleven haplotypes of *E. granulosus* were identified. (A) Haplotype network constructed based on haplotype data of human-derived *E. granulosus* in Qinghai Province. (B) EgH1: Haiyan, Dari, Zhiduo, Jianzha; EgH2: Tongde; EgH3: Dulan; EgH4: Yushu, Zaduo, Qumalai, Shiqu; EgH5: Chengduo; EgH6: Yushu; EgH7: Zhiduo; EgH8: Nangqian, Gande; EgH9: Nangqian; EgH10: Qumalai; EgH11: Guinan; The area of each circle is proportional to the number of haplotypes. The basic map review number is GS(2024)0650.

ecological effects triggered by cold climates may have driven the divergence between these two *Echinococcus* species.

E. granulosus and *E. canadensis* are recognized as sister species in the genus *Echinococcus*^[9]. Their divergence is estimated to have occurred approximately 2.72 Mya, leading us to infer that they likely diverged in the early Pleistocene and have since evolved as sister species through coordinated differentiation. This divergence time falls within the period of the intense uplift of the QTP around 3 Mya^[10]. Such profound geographic and climatic changes may have served as a key driver of their speciation.

Although this study employs molecular genetics approaches and provides a preliminary analysis of the population structure of human-derived *E. granulosus* in Qinghai, the relatively small sample size is a major limitation. A small sample size may overrepresent rare haplotypes while underestimating low-frequency haplotypes that naturally occur in the population but were not sampled. Therefore, the 11 haplotypes and their relative frequencies identified in this study likely

reflect only a subset of the full genetic diversity present in the Qinghai region. Our findings indicate that the human-derived *E. granulosus* in Qinghai belongs predominantly to the G1 genotype, with EgH4 as the dominant haplotype. Based on an analysis of the regional geological and evolutionary context, we hypothesized that the divergence of *E. granulosus* may be associated with the intense uplift of the Tibetan Plateau. This study provides important scientific evidence for tracing the origin of *E. granulosus* in the Qinghai region, clarifying its transmission routes, and formulating targeted prevention and control strategies.

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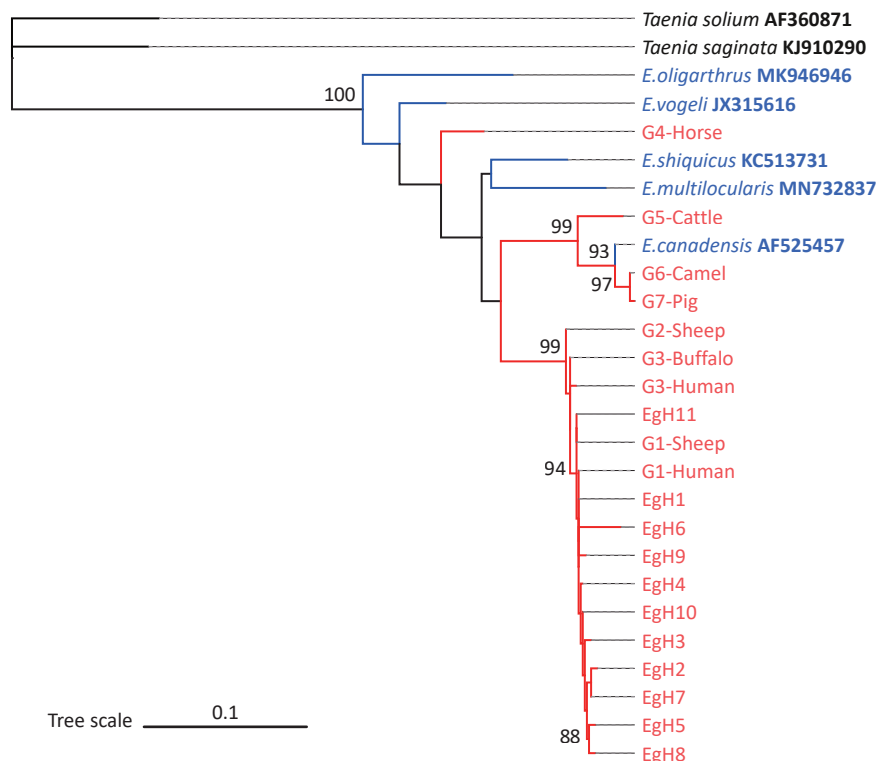


Figure 2. Maximum likelihood phylogenetic tree of *E. granulosus* based on the COX1 gene. The text in black font represents the outgroup, the text in blue represents the genus *Echinococcus*, the text in red represents *E. granulosus*, and Eg1-Eg11 are the fine-grained *E. granulosus* haplotypes obtained in this study.

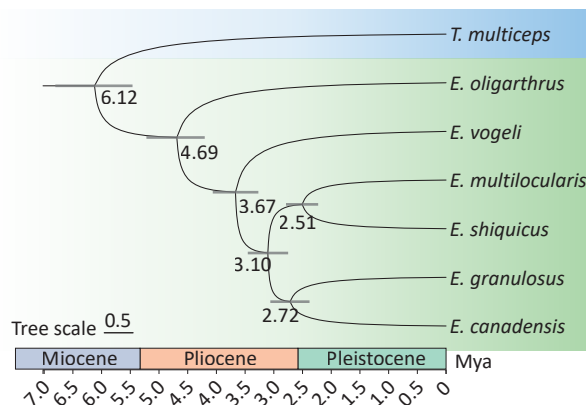


Figure 3. Divergence time estimation of *Echinococcus* species based on the COX1 gene.

Competing Interests The authors declare no competing interests.

Ethics This study was conducted in accordance with the Declaration of Helsinki and has obtained approval from the Ethics Committee of Qinghai University (PJ202501-102).

Authors' Contributions Conceptualization, Project administration, Data analysis, and manuscript drafting: Yong Fu, Hailong Zhao; Executing laboratory experiments, Data analysis, and manuscript drafting: Hongrun Ge; Sample collection, Environmental data collection, Data analysis, and review & Editing: Ru Meng, Zhi Li, Hong Duo, Yuanqing Lin, Suoang Qiupei, Xihuo You, Qinyi He. All authors read and approved the final manuscript.

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