Phylogenetic evidence for multiple intertypic recombinations in enterovirus B81 strains isolated in Tibet, China

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Enterovirus B81 (EV-B81) is a newly identified serotype within the species enterovirus B (EV-B). To date, only eight nucleotide sequences of EV-B81 have been published and only one full-length genome sequence (the prototype strain) has been made available in the GenBank database. Here, we report the full-length genome sequences of two EV-B81 strains isolated in the Tibet Autonomous Region of China during acute flaccid paralysis surveillance activities, and we also conducted an antibody seroprevalence study in two prefectures of Tibet. The sequence comparison and phylogenetic dendrogram analysis revealed high variability among the global EV-B81 strains and frequent intertypic recombination in the non-structural protein region of EV-B serotypes, suggesting high genetic diversity of EV-B81. However, low positive rates and low titers of neutralizing antibodies against EV-B81 were detected. Nearly 68% of children under the age of five had no neutralizing antibodies against EV-B81. Hence, the extent of transmission and the exposure of the population to this EV type are very limited. Although little is known about the biological and pathogenic properties of EV-B81 because of few research in this field owing to the limited number of isolates, our study provides basic information for further studies of EV-B81.

Enteroviruses (EVs) belong to the family Picornaviridae within the new order Picornavirales. The genome of the small non-enveloped viruses is a single-stranded, positive-sense RNA molecule of approximately 7500 nucleotides consisting of a single open reading frame flanked by 5’ and 3’ untranslated regions (UTRs). A single polyprotein translated from the RNA strand is first cleaved into three polyprotein precursors: P1, P2, and P3. P1 is processed to yield four structural proteins: Viral protein 1–4 (VP1–VP4); P2 and P3 are precursors of the nonstructural proteins 2A–2C and 3A–3D, respectively.

Most EV infections are asymptomatic or cause only mild symptoms. However, some EVs can also cause a broad spectrum of other clinical illnesses, including acute flaccid paralysis (AFP); acute hemorrhagic conjunctivitis; encephalitis; aseptic meningitis; and hand, foot, and mouth disease1–4. Human EVs now comprise more than 110 serotypes, which are currently classified into four species, EV-A to EV-D, on the basis of their molecular and biological characteristics5. Although generally reliable, the neutralization test—a traditional method for EV typing—has gradually been replaced because it is labor-intensive and time-consuming; furthermore, it may fail to identify an isolate because of aggregation of virus particles, antigenic drift, or the presence of multiple viruses in the specimen6. Current EV classification is based on the high nucleotide sequence divergence within the VP1 capsid-coding region, which has been shown to correspond with serotype neutralization7. According to the recommended specific criteria for the interpretation of VP1 sequence data, EVs are classified into the same type if they have more than 75% nucleotide similarity (85% amino acid similarity) and into different types if they have less than 70% nucleotide similarity in this region. While 70–75% nucleotide similarity in VP1 region has been considered as a “grey zone” of molecular typing of EVs, and in this instance, additional information such as complete P1 region sequences nucleotide similarity or neutralization profile should be obtained to decide the EV serotype. A large number of new EV types have been discovered after molecular typing methods became available.

Enterovirus B81 (EV-B81) is a newly identified serotype within the species EV-B. The prototype strain (USA/CA68-10389) of EV-B81 was isolated in the USA in 19688. Subsequently, several other EV-B81 strains were isolated from AFP patients or healthy individuals during AFP case surveillance in China9, Bangladesh10, India11, 12.
distinct EV-B81 strains differ considerably. The complete genome within the EV-B species. The nucleotide sequences available for the strains and the EV-B81 prototype strain and other prototype strains EV-B81 prototype strain. Table 1 shows the nucleotide sequence and sites were predicted based on the full-length genome sequence of the overall base compositions of strains 99279 and 99298c were at position 7328 and a nucleotide deletion at position 7341. The at position 179. In the 3' insertions at positions 101, 102, and 118 and a nucleotide deletion (USA/CA68-10389) showed that they all had the same genomic EV-B81 strains with the genome of the EV-B81 prototype strain followed by a poly (A) tail composed of a long sequence of adenine nucleotides. Alignment of the full-length genomes of the two Tibetan isolates were then partially sequenced using molecular typing methods and analyzed by an online enterovirus genotyping tool. Both isolates were identified as EV-B81.

Results

Serotyping and molecular typing of the Tibetan isolates. The Tibetan isolates (strain 99279/XZ/CHN/1999 and strain 99298c/XZ/CHN/1999, hereafter referred to as 99279 and 99298c, respectively) were initially characterized using a standard pool of EV typing antisera (RIVM, the Netherlands) distributed by the World Health Organization. However, neither of the isolates could be neutralized by any of the antisera (data not shown). Therefore, the isolates were initially identified as “untypeable” non-polio EVs. The VP1 capsid-coding regions of the two Tibetan isolates were then partially sequenced using molecular typing methods and analyzed by an online enterovirus genotyping tool. Both isolates were identified as EV-B81.

Full-length genomic characterization of Chinese EV-B81 strains. The full-length genomes of the two EV-B81 strains were sequenced. Both were sequenced in length, encoding a polypeptide of 2191 amino acids. The coding sequences were flanked by a non-coding 5' UTR of 741 nucleotides and a non-coding 3' UTR of 100 nucleotides followed by a poly (A) tail composed of a long sequence of adenine nucleotides. Alignment of the full-length genomes of the two Tibetan EV-B81 strains with the genome of the EV-B81 prototype strain (USA/CA68-10389) showed that they all had the same genomic organization and collinear order of genomic regions. However, in the 5' UTR, Tibetan EV-B81 strains contained three nucleotide insertions at positions 101, 102, and 118 and a nucleotide deletion at position 179. In the 3' UTR, they contained a nucleotide deletion at position 7328 and a nucleotide deletion at position 7341. The overall base compositions of strains 99279 and 99298c were 28.03% and 28.11% A, 24.36% and 24.35% G, 23.82% and 23.73% C, and 23.79% and 23.81% U, respectively. The polypeptide cleavage sites were predicted based on the full-length genome sequence of the EV-B81 prototype strain. Table 1 shows the nucleotide sequence and deduced amino acid sequence identities between the Tibetan EV-B81 strains and the EV-B81 prototype strain and other prototype strains within the EV-B species. The nucleotide sequences available for the distinct EV-B81 strains differ considerably. The complete genome nucleotide sequence similarity between these two Tibetan EV-B81 strains is 99.6%, and they displayed 79.1% and 79.2% nucleotide identity and 94.7% and 95.2% amino acid identity with the prototype EV-B81 strain, respectively.

Phylogenetic analysis of the Chinese EV-B81 strains and other EV-B genomes. Phylogenetic trees were generated from the 5'-nucleotide (nucleotide 2567–2818) partial VP1 coding region of the two Tibetan EV-B81 strains and eight other EV-B81 strains available in the GenBank database (Fig. 1). The two Tibetan strains displayed great genetic distance to another Chinese strain (strain 142-98-YN/CHN/1998 isolated in the Yunnan province in 1998) and clustered with the strain BAN-10491 isolated from Bangladesh. Three strains isolated from India clustered together, and most branches displayed long genetic distances.

To investigate the genetic relationship between the Tibetan EV-B81 strains, the EV-B81 prototype strain, and other EV-B prototype strains available in the GenBank database, we constructed phylogenetic trees based on the VP1, P1, P2, and P3 regions of the genome. The phylogenetic tree based on the VP1 region also contained an EV-B81 strain (N-428/IND/2008) isolated from India, for which the entire VP1 region was available in the GenBank database (Fig. 2). In the VP1 and P1 capsid regions, the two Tibetan EV-B81 strains clustered together with the EV-B81 prototype strain and the Indian EV-B81 strain, confirming the preliminary molecular typing results. In the VP1 region, the nucleotide identity between the Tibetan strains

| Region | EV-B81 strain 99279 | Proteins of other EV-B | EV-B81 strain 99298c | Proteins of other EV-B |
|--------|---------------------|------------------------|----------------------|------------------------|
| 5' UTR | 78.3                | 66.3–89.4              | 78.4                 | 66.3–89.3              |
| VP4    | 78.2 (95.6)         | 67.6–79.7 (71.0–82.6)  | 78.2 (95.6)          | 67.6–79.7 (71.0–82.6)  |
| VP2    | 79.6 (96.5)         | 64.5–73.4 (73.3–84.2)  | 79.6 (96.5)          | 64.3–73.5 (73.3–83.9)  |
| VP3    | 79.8 (100.0)        | 62.1–73.0 (67.5–87.4)  | 79.8 (100.0)         | 62.0–72.8 (67.5–87.4)  |
| VP1    | 79.1 (94.7)         | 53.8–68.8 (53.3–79.5)  | 79.2 (95.4)          | 53.9–69.2 (53.6–79.5)  |
| 2A     | 79.3 (93.3)         | 75.1–82.8 (85.3–96.6)  | 79.3 (93.3)          | 75.1–82.8 (86.0–97.3)  |
| 2B     | 77.5 (94.0)         | 72.6–79.8 (91.0–96.0)  | 77.2 (94.0)          | 72.6–79.8 (91.0–96.0)  |
| 2C     | 80.9 (96.9)         | 78.0–85.7 (95.1–99.1)  | 81.1 (97.2)          | 78.2–85.8 (95.4–98.4)  |
| 3A     | 75.2 (92.1)         | 73.7–90.2 (88.7–97.7)  | 75.2 (92.1)          | 73.7–90.2 (88.7–97.7)  |
| 3B     | 77.2 (100.0)        | 72.7–89.3 (90.9–100.0) | 77.2 (100.0)         | 72.7–89.3 (90.9–100.0) |
| 3C     | 77.9 (95.6)         | 75.0–88.7 (92.8–99.4)  | 78.1 (95.6)          | 75.2–88.7 (92.8–99.4)  |
| 3D     | 79.4 (96.7)         | 77.8–86.2 (94.5–98.0)  | 79.6 (96.7)          | 77.9–86.1 (94.5–98.0)  |
| 3' UTR | 84.6                | 75.9–90.2              | 84.6                 | 75.9–90.2              |

Figure 1 | Phylogenetic relationships based on partial VP1 region sequences of enterovirus B81 (EV-B81). Two Tibetan EV-B81 strains isolated in this study (indicated by circles) and other EV-B81 strains (available in the GenBank database) were analyzed based on the 252-nucleotide (nucleotide 2567–2818) partial VP1 coding region sequence. The strain indicated by a diamond is the EV-B81 prototype strain.
Figure 2 | Phylogenetic relationships based on the VP1, P1, P2, and P3 genome regions of enterovirus B (EV-B). Two Tibetan EV-B81 strains (indicated by solid circles) and 55 other EV-B prototype strains were analyzed by nucleotide sequence alignment using the Neighbor-Joining algorithms implemented in the MEGA 5.0 program. Numbers at the nodes indicate bootstrap support for that node (percent of 1000 bootstrap replicates). The open triangle indicates the India EV-B81 which has the entire VP1 sequence in the GenBank database, and the solid diamond indicates EV-B81 prototype strain. The scale bars represent the genetic distance. All panels have the same scale. (a) VP1 coding sequences; (b) P1 coding sequences; (c) P2 coding sequences; and (d) P3 coding sequences.
99279 and 99298c and the Indian strain was 88.8% and 89.1%, respectively. However, in the non-capsid regions, the phylogenetic trees differed greatly from those in the capsid regions. In the P2 and P3 regions, the two Tibetan EV-B81 strains shared the highest similarity with the prototype strains of EV-B111 and EV-B86, respectively. This surprised us because EV-B111 is a recently reported type of enterovirus\(^2\). The phylogenetic analysis indicated that recombinations between Chinese EV-B81 strains and other EV-B serotypes might have occurred.

**Recombinant structure of the Chinese EV-B81 strains.** Similarity plots and bootscanning analyses were performed to confirm the recombinations between the Tibetan EV-B81 strains and other EV-B prototype strains. Because the two Tibetan EV-B81 strains share high nucleotide identity in their full-length genomes (99.6%), only one of the strains (99279) was used as a query sequence. It was compared with the EV-B81 prototype strain (USA/CA68-10389) and other EV-B prototype strains. In the P1 coding region, strain 99279 possessed the highest similarity with the EV-B81 prototype strain as expected (Fig. 3). However, in the 5’ UTR, P2, P3, and 3’ UTR regions, strain 99279 was apparently not related to the EV-B81 prototype strain, which further confirmed the occurrence of recombination in these regions. Interestingly, a relatively high similarity within the 3’ end of the 2C region to the 5’ end of the 3C region was found between strain 99279 and the EV-B111 prototype strain, which was supported by the bootscanning analysis. And the bootscanning analysis also suggested the possibility of recombination of small-sized fragment in 5’ UTR, 3D, and 3’ UTR region between Chinese EV-B81 strains and other EV-B serotypes such as EV-B86, EV-B107, and EV-B87. (Fig. 3).

**Seroprevalence of EV-B81 in Tibet.** Among the 50 serum samples surveyed, 16 were seropositive for EV-B81 (>1:8), with a total positive rate of 32.0% and geometric mean titers (GMTs) of 1:27 among the positive sera samples. The composition ratios for the anti-EV-B81 antibody titers of <1:8, 1:8–1:64, and >1:64 were 68%, 32%, and 0, respectively. Compared with seroepidemiology studies of other EVs in China, the positive rate and GMTs of EV-B81 are apparently lower than that of other EVs such as EV-A71 and CVA16 in the same age group (1–5 years old)^16,17.

Although both Lhasa City and Shigatse Prefecture showed low seroprevalence rates and low titers of anti-EV-B81 antibodies, there were some differences between the two areas. In Lhasa City, the seroprevalence rate and GMTs of the positive sera samples were 16% and 1:22.6, respectively, and in Shigatse Prefecture, they were 48% and 1:28.5, respectively. The positive rate in Shigatse is significantly higher than those in Lhasa City, while there is no significant difference between the GMTs of the two prefectures (seroprevalence rate: p = 0.0153, GMTs: p = 0.521).

**Discussion**

EV-B81 is a new type of enterovirus belonging to the species EV-B. The prototype of EV-B81 was isolated in the USA in 1968. Subsequently, several other EV-B81 strains were isolated from different countries in Asia and Africa, indicating global distribution of the serotype. Of the eight EV-B81 strains reported in the GenBank database to date, six were isolated from Southeast Asia, including from India, Bangladesh, and provinces in southwestern China, suggesting the possible circulation of EV-B81 in this area. Detailed information about the prototype strain, such as information about its host, was not indicated in the reference publications, but almost all the other EV-B81 strains were isolated from patients with AFP except for a strain from Cameroon\(^2\). Hence, there may be some correlation between EV-B81 and AFP. However, little research has been done on EV-B81 worldwide, and more data are necessary to unveil the biological and pathological properties of EV-B81.

The first EV-B81 strain reported in China was isolated from the Yunnan province in 1998, and no other case has since been reported in China\(^1\). In this study, we characterized the full-length genomes of two EV-B81 strains isolated in 1999 from Mangkang Prefecture of Tibet, located in the southeast of the Qinghai-Tibet Plateau. Although EV-B81 strains were reported in both Tibet and the Yunnan province of China, they showed great genetic diversity, the level of genetic diversity between two Tibetan EV-B81 strains and Yunnan strains was 20.4% and 20.9%, respectively, suggesting that EV-B81 from different lineages circulated separately in these two regions. For the entire VP1 sequences, the nucleotide identities between the two Tibetan EV-B81 strains 99279 and 99298c and the EV-B81 prototype strain were 79.1% and 79.2%, respectively, further indicating that EV-B81 strains have high nucleotide sequence diversity.

Recently, increasing numbers of studies have shown that intertypic recombination is common within the species EV-A, EV-B, and EV-C\(^3\)–\(^5\). The two EV-B81 strains characterized in this study are no exception. The clustering of the two strains in the phylogenetic trees based on the P2 and P3 regions was inconsistent with the clustering in the tree based on the P1 region, which suggested intertypic recombination between the two EV-B81 strains and other EV-B serotypes (probably EV-B86 or EV-B111) in the P2 and P3 regions. The result was further corroborated by similarity plot and bootscanning analyses. Previous studies have indicated that recombination between different serotypes may occur when different viruses infect and replicate in the same cell, and that recombination usually occurs among EV serotypes within a species\(^3\)–\(^5,18\). Hence, we can assume that the two Tibetan EV-B81 strains may have co-circulated with other EV-B serotypes, especially EV-B111, for a period before they were isolated. The EV-B111 prototype was also isolated in the Tibet Autonomous Region in the same period. However, more data are needed to define the exact serotype of the donor sequence.

Most EV infections are asymptomatic and are hence an under-estimated epidemic. To investigate the prevalence of EV-B81 in the Tibet Autonomous Region, we conducted a seroepidemiology survey. We found that the seropositive rate and GMT were higher in Shigatse Prefecture than in Lhasa City (p < 0.05), indicating a geographical difference in EV-B81 infection patterns. However, the seropositive rates and titers of anti-EV-B81 antibodies were overall very low compared with other EVs prevalent in China such as EV-A71 and CVA16\(^6\), suggesting that the extent of transmission and the exposure of the population to EV-B81 were limited. However, considering the small sample sizes (n = 50) of the survey and the limited number of EV-B81 strains, the possibility of cross reactivity between EV-B81 and antibody against other closely related EVs, in particular members within EV-B, could not be exclude, so further research will be required to draw firm conclusions.

In conclusion, we report the full-length genome sequences of two EV-B81 strains isolated during AFP surveillance in the Tibet Autonomous Region, China. Sequence analysis revealed high genetic diversity in the two strains compared with the EV-B81 prototype, as well as intertypic recombination in the non-structural protein region of both strains. Although little is known about the biological and pathogenic properties of EV-B81 because of few research in this field owing to a limited number of isolates, our study provides a basis for further study of EV-B81.

**Methods**

**Sample collection.** This study did not involve human participants or human experimentation; the only human materials used were stool samples collected from AFP patients or their close contacts at the instigation of the Ministry of Health P. R. of China for public health purposes, and written informed consent for the use of their clinical samples was obtained from their parents of all the patients involved in this study. This study was approved by the second session of the Ethics Review Committee of the National Institute for Viral Disease Control and Prevention, Chinese Center for Disease Control and Prevention.
The two EV-B81 strains (strain 99279 and 99298c) were isolated from stool samples from two children living in the same village in Mangkang Prefecture of the Tibet Autonomous Region, China. The samples were collected in 1999, during the course of poliovirus surveillance activities in support of the global polio eradication initiative.

Viral isolation and primary identification. Stool samples from AFP patients were collected in Lhasa City and Shigatse Prefecture. No children had any sign of disease at the time of sample collection.

Molecular typing. For molecular typing, viral RNA was extracted from the viral isolates using a QIAamp Viral RNA Mini Kit (Qiagen, Germany) and stored at −80°C until use. Reverse transcription polymerase chain reaction (RT-PCR) was performed to amplify the VP1 coding region using PrimeScript One Step RT-PCR Kit Ver.2 (TaKaRa, Dalian, China) with primer pairs 008 and 0137. The PCR products were purified using the QIAquick PCR purification kit (Qiagen, Germany) and then sequenced on an ABI 3130 Genetic Analyzer (Applied Biosystems, USA).

Neutralizing antibody detection. Neutralizing antibodies against EV-B81 were detected with a neutralization test using the microtechnique and the human RD cell line as previously described, with some modifications. Serum samples were inactivated at 56°C for 30 min before use, and sample dilutions of 1:2 to 1:8, and the GMT was calculated.

Full-length genomic sequencing. Two long-distance PCR amplifications were performed using the SuperScript III One-Step PCR-PCR system (Invitrogen). Reaction mixtures (50 μL) contained template RNA (5 μL), reaction buffer (25 μL),...
forward [0001548 (GGGGACAGTTTGCATACAAAAAGCGGGCTTT)4] or E989 (TGTGTTGTGGTCTCCGTTTCTGGG)4] and reverse [E982 (GGTGGTCGWTYGGCCTTGA)4] or 7500A (GGGGACATTCGATGAAACAGGTTGG(T)24)] primers (1.0 ng/L), and RT/Platinum Taq High Fidelity enzyme (5 U). cDNA synthesis and pre-denaturation were carried out with one cycle of 50 °C (30 min) and 94 °C (2 min).

Amplification was carried out with 40 cycles of 94 °C (15 s), 60 °C (30 s), and 68 °C (5 s), followed by a final incubation at 68 °C (5 min). The primers used for sequencing of the full-length genome were designed by a “primer-walking” strategy.

**Phylogenetic and bioinformatics analyses.** The nucleotide and deduced amino acid sequences of strains 99279 and 99298c were compared with those of the prototype EV-B strains by pairwise alignment using the MEGA program (version 5.03).

Phylogenetic trees were constructed by the neighbor-joining method implemented in the MEGA program using the Kimura 2-parameter model. Regions containing alignment gaps were omitted from the analysis. The branch lengths of the dendrogram were determined from the topologies of the trees and were obtained by majority rule consensus among 1000 bootstrap replicates. Bootstrap values greater than 80% were considered statistically significant for grouping. Similarity plot and bootscanning analyses were performed using the SimPlot program (version 3.5).

Stuart Ray, Johns Hopkins University, Baltimore, MD, USA). For similarity plot analyses, a 200-nucleotide window was moved in 20-nucleotide steps, and bootscanning analyses were run with the neighbor-joining method.

**Statistical analysis.** The titers of neutralization antibodies were log-transformed to calculate the GMTs. Chi-square test was used to compare the seroprevalence rates with an error probability of P < 0.05 were regarded as significant. All statistical analyses were performed with IBM SPSS Statistics software.

**Nucleotide sequence accession numbers.** The complete genomic sequences of the EV-B81 strains (99279 and 99298c) described in this study were deposited in the GenBank database under the accession numbers KJ755189 and KJ755190, respectively.

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