Emergence of Japanese encephalitis virus genotype V in the Republic of Korea

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Abstract

Background: Japanese encephalitis virus (JEV) genotype V reemerged in Asia (China) in 2009 after a 57-year hiatus from the continent, thereby emphasizing a need to increase regional surveillance efforts. Genotypic characterization was performed on 19 JEV-positive mosquito pools (18 pools of Culex tritaeniorhynchus and 1 pool of Cx. bitaeniorhynchus) from a total of 64 positive pools collected from geographically different locations throughout the Republic of Korea (ROK) during 2008 and 2010.

Findings: Two regions of the JEV genome were sequenced from 19 pools; the envelope gene and the nonstructural protein 5 (NS5)/3'-untranslated region (UTR). Eighteen pools of Culex tritaeniorhynchus and one pool of Cx. bitaeniorhynchus were positive for genotype I and genotype V, respectively. Sequence alignment of the complete E gene from Cx. bitaeniorhynchus showed high amino acid similarity (98.8%) to the Muar strain, characterized as the first report of genotype V, isolated from an encephalitis patient in Malaysia in 1952.

Conclusion: This study represents the first report of JEV genotype V in the ROK. The reemergence of genotype V in Asia (China and ROK) after more than a half-century and its discovery in Cx. bitaeniorhynchus, a mosquito species previously unknown to carry JEV in the ROK, emphasizes the need for enhanced JE surveillance to monitor the dynamics of JEV strains within the region. Future findings may have implications with regard to JEV vaccination/prevention strategies.

Keywords: Japanese encephalitis virus, genotype I, genotype V, Culex tritaeniorhynchus, Culex bitaeniorhynchus, Muar

Background

Japanese encephalitis virus (JEV) is a mosquito-borne member of the family Flaviviridae, genus Flavivirus, and a primary cause of viral encephalitis in humans within its range [1]. The positive-sense RNA viral genome is approximately 11 kb in length and is translated into three structural proteins [Capsid (C), Membrane (M), and Envelope (E)] and seven nonstructural (NS) proteins (NS1, NS2A, NS2B, NS3, NS4A, NS4B, and NS5) with untranslated regions (UTR) at the 5' and 3' ends of the genome [2]. Historically, Culex tritaeniorhynchus was implicated as the primary vector of JEV in the Republic of Korea (ROK) and much of Asia [3,4]. However, JEV has since been detected in additional culicine species throughout its range, including Cx. bitaeniorhynchus from the ROK [5]. JEV strains are generally classified into five genotypes (genotypes I, II, III, IV, and V) based on similarities in the E gene nucleotide sequence [6]. Previously, only genotype I was detected on the Korean peninsula [7]. Therefore, we characterized JEV-positive pools of Cx. tritaeniorhynchus and Cx. bitaeniorhynchus to determine whether the unexpected finding of JEV in Cx. bitaeniorhynchus in the ROK may have coincided with the appearance of an additional genotype.

Materials and Methods

Nineteen JEV-positive mosquito pools, from a total of 64 JEV-positive pools collected during 2008 and 2010 in the ROK (18 pools of Cx. tritaeniorhynchus and 1 pool of Cx. bitaeniorhynchus), and one JEV culture received from
USAMRIID (United States Army Medical Research Institute of Infectious Diseases, USA) were genotypically characterized (Table 1, Figure 1). Total RNA was extracted from mosquito homogenate using Trizol reagent (Invitrogen, USA) in accordance with the manufacturer’s instructions and was resuspended in 50 µl of RNase-free water containing 10 units of RNasin™ Plus RNase Inhibitor (Promega, USA). RNA was used as the template for cDNA synthesis using the SuperScript III first strand synthesis system (Invitrogen, USA) with a random hexamer primer. The synthesized cDNA was then used for PCR amplification using iProof™ High-Fidelity DNA polymerase (Bio-Rad, USA). The NS5 gene/3'UTR and envelope (E) gene of 19 JEV-positive pools were amplified using EMF1/VD8 primers [8] and 940S/1720A primers [9], respectively. Products were purified using the QIAquick PCR purification kit (Qiagen, USA) and sequenced using the Applied Biosystems, USA. The sequences were edited and assembled using the Sequencer program v4.1.4 (Applied Biosystems, USA). Multiple sequence alignments and phylogenetic analysis were performed using ClustalX version 2.0.11 and MEGA version 5 programs [10,11]. Percent sequence similarity/divergence was calculated using the MegAlign program found in the Lasergene v.8 software (DNASTAR, Inc., Madison, WI, USA). Phylogenetic analysis of the partial E gene (705 bp) was performed using the neighbor-joining method and Tamura-Nei model of nucleotide substitution. The maximum likelihood (ML) tree was constructed from the NS5/3'UTR nucleotide sequences (550 bp) by PhyML software v 3.0 [12] using the best fit model with aLRT branch support [13]. The ML tree for the complete E gene used the Tamura-Nei model with bootstrap analysis (2,000 replicates) for testing the reliability of the tree using the MEGA5 (version 5) program (The Biodisgn Institute, Tempe, Arizona) [11].

Results

The phylogenetic relationships among 19 JEV strains and JEV sequences retrieved from GenBank representing genotypes I-V were analyzed. The ML tree for the NS5/3'UTR (550 bp) and the neighbor-joining tree for the partial E gene (705 bp) showed similar branching patterns with high bootstrap support. Therefore, the ML tree is only presented in this report (Figure 2). Two genotypes were identified among the 19 JEV strains. JEV strains from 18 Cx. tritaeniorhynchus mosquitoes grouped into genotype I. These genotype I strains were closely-related to strains isolated from China, Korea, Japan, Vietnam, and Thailand from the early 1980s to the present (Figure 2). The remaining strain from Cx. bitaeniorhynchus (10-1827) grouped into genotype V together with the Muar strain which was isolated from an encephalitis patient in Malaysia in 1952.

The complete E gene was sequenced from a subset of strains in genotype I (A10.825, A10.881, A8.789) and genotype V (10-1827). The ML tree constructed from the complete E gene of these strains together with representative JEV genotype I-V sequences is shown in Figure 3. This ML tree supports the phylogenetic analysis results performed on the NS5/3'UTR (Figure 2) and the partial E gene previously mentioned. The ML tree in Figure 3 shows that the 10-1827 strain grouped with the Muar strain with 79% bootstrap support, while the remaining sequences clustered in genotype I together with K01-JN and K05-GS strains that were isolated from Cx. tritaeniorhynchus in the ROK in 2001 and 2005, respectively.

Sequence analysis of 18 strains shows minimal sequence variation among viruses in genotype I, with nucleotide sequence similarity of 97.5-100% for the NS5/3'UTR (Figure 2) and 99.6-100% for the E gene (Figure 3). In an earlier study, genetic stability was also observed among JEV strains isolated from mosquito vectors in the ROK between 1994 and 2005 [7]. Examination of the complete E sequence of 10-1827 strain (genotype V) showed less similarity to the other genotypes, with nucleotide similarity approximately 77.3% (91.3% for amino acids) to genotype I (K01-JN, K05-GS), 78.1% (91.0% for amino acids) to genotype II (FU strain), 77.7% (90.4% for amino acids) to genotype III (Nakayama), and 77.8% (91.0% for amino acids) to genotype IV (JKT6468) (Table 2). However, nucleotide and amino acid similarities to the Muar strain were 90.0% and 98.8%, respectively (Table 2). Likewise, the XZ0934 strain, a JEV genotype V recently isolated from China (2009), showed E gene nucleotide and amino acid sequence similarities to the Muar strain of 86.0% and 93.2%, respectively [14].

Figure 4 shows the amino acid sequence alignment of the complete E gene derived from strains A10.825, A10.881, A8.789, and 10-1827 and reference sequences (Muar, K01-JN, K05-GS). The E protein of the strains in genotype I is very conserved with few amino acid changes detected: A10.825 (from S = serine to N = asparagine at position 123) and A8.789 (from L = leucine to M = methionine at position 371). The alignment reveals differences in 6 amino acid residues between the Muar and 10-1827 strains (Figure 4). The eight Muar signature amino acid residues in domain III comprising a putative receptor binding region [15] were also identified in the 10-1827 strain along with the critical amino acid residue thought to be involved in receptor binding activity (Q = glutamine at position 327) [16]. Table 3 provides a complete listing of the strains that are referenced in this study.
| Collection Serial No. | Collection Date | Collection Sites (US Military Bases, Villages/Cities) | Province     | Species               | Accession no.               |
|----------------------|-----------------|------------------------------------------------------|--------------|-----------------------|----------------------------|
| A8.789               | 29-Jul-08       | Haenam                                               | Jeonnam      | Cx. tritaeniorhynchus | JNS587257, JNS587261       |
| A10.825              | 28-Sep-10       | Changnyeong                                          | Gyeongnam    | Cx. tritaeniorhynchus | JNS587255, JNS587259       |
| A10.881              | 21-Oct-10       | Jinju                                                | Gyeongnam    | Cx. tritaeniorhynchus | JNS587256, JNS587260       |
| 10-1742              | 1-Sep-10        | Warrior Base* (Munsan)                               | Gyeonggi     | Cx. tritaeniorhynchus | JNS587241                 |
| 10-1748              | 1-Sep-10        | Warrior Base* (Munsan)                               | Gyeonggi     | Cx. tritaeniorhynchus | JNS587242                 |
| 10-1728              | 31-Aug-10       | Daeseongdong                                         | Gyeonggi     | Cx. tritaeniorhynchus | JNS587240                 |
| 10-1937              | 11-Sep-10       | Daeseongdong                                         | Gyeonggi     | Cx. tritaeniorhynchus | JNS587245                 |
| 10-2044              | 14-Sep-10       | Daeseongdong                                         | Gyeonggi     | Cx. tritaeniorhynchus | JNS587248                 |
| 10-2097              | 21-Sep-10       | Daeseongdong                                         | Gyeonggi     | Cx. tritaeniorhynchus | JNS587249                 |
| 10-2130              | 21-Sep-10       | Daeseongdong                                         | Gyeonggi     | Cx. tritaeniorhynchus | JNS587250                 |
| 10-2357              | 13-Oct-10       | Daeseongdong                                         | Gyeonggi     | Cx. tritaeniorhynchus | JNS587252                 |
| 10-1827              | 8-Sep-10        | Daeseongdong                                         | Gyeonggi     | Cx. bitaeniorhynchus  | JNS587243, JNS587258       |
| 10-1835              | 8-Sep-10        | CP Humphreys* (Pyeongtaek)                            | Gyeonggi     | Cx. tritaeniorhynchus | JNS587244                 |
| 10-1291              | 5-Aug-10        | Gunsan Air Base* (Gunsan)                            | Jeonbuk      | Cx. tritaeniorhynchus | JNS587239                 |
| 10-2004              | 8-Aug-10        | Gunsan Air Base* (Gunsan)                            | Jeonbuk      | Cx. tritaeniorhynchus | JNS587251                 |
| 10-1990              | 30-Aug-10       | Gwangju Air Base* (Gwangju)                          | Jeonnam      | Cx. tritaeniorhynchus | JNS587246                 |
| 10-1992              | 30-Aug-10       | Gwangju Air Base* (Gwangju)                          | Jeonnam      | Cx. tritaeniorhynchus | JNS587247                 |
| 10-2378              | 2-Sep-10        | Gwangju Air Base* (Gwangju)                          | Jeonnam      | Cx. tritaeniorhynchus | JNS587253                 |
| 10-2397              | 27-Sep-10       | Gwangju Air Base* (Gwangju)                          | Jeonnam      | Cx. tritaeniorhynchus | JNS587254                 |

Locations are presented in Figure 1.
* US military training site or installation.
Figure 1 Locations of JEV-positive mosquito pools collected during 2008 and 2010 in the Republic of Korea. Daeseongdong is a village near the military demarcation line (MDL) (center of the 4-Km wide demilitarized zone separating North and South Korea); Warrior Base training area is approximately 5 km north of Munsan; Camp Humphreys is in a rural area of Pyeongtaek; Gunsan Air Base is located near the small city of Gunsan; Gwangju Air Base is located near the metropolitan city of Gwangju; Haenam, Jinju, and Changnyeong sites are beef/swine farms near the small cities. Pool = number of sequenced samples/total JEV-positive samples.
Conclusion

This study is the first report of JEV genotype V in the ROK and represents the third report of genotype V in Asia, with the most recent findings from *Cx. tritaeniorhynchus* collected in Tibet, China (2009) [14]. The fact that JEV genotype V, first reported from an encephalitis patient in Malaysia in 1952 (Muar strain), came long before the discovery of its reemergence in China in 2009 and now its subsequent appearance in the ROK may mark the beginning of a genotypic shift in JEV within the region. Additionally, the emergence of this strain in *Cx. bitaeniorhynchus*, a mosquito
species previously unknown to carry JEV in the ROK, underscores the need to step-up surveillance efforts within the ROK. The reemergence of this genotype after 57 years may have future implications with regard to JEV vaccination effectiveness and policy among civilian and military populations, as well as with preventive strategies designed to reduce the health impact and incidence of JEV among at risk Asian populations.

Table 2 Nucleotide sequence similarity and divergence of the complete E gene from ROK mosquito pools

|                | A10.825 | A10.881 | A8.789 | JE_USAMRIID | K01-JN | K05-GS | FU | Nakayama | JKT6468 | 10-1827 | Muar |
|----------------|---------|---------|--------|-------------|--------|--------|----|----------|---------|---------|------|
| A10.825 (1)    | 99.4    | 99.3    | 94.3   | 98.2        | 99.3   | 89.0   | 87.8| 81.8     | 77.2    | 76.8    |
| A10.881 (1)    | 0.6     | 99.3    | 94.1   | 98.1        | 99.2   | 89.0   | 88.0| 82.2     | 77.4    | 76.5    |
| A8.789 (1)     | 0.7     | 0.7     | 94.3   | 98.2        | 99.3   | 89.1   | 87.8| 82.1     | 77.2    | 76.6    |
| JE_USAMRIID (1)| 6.0     | 6.3     | 6.0    | 94.1        | 94.3   | 88.0   | 86.6| 82.1     | 77.2    | 76.6    |
| K01-JN (1)     | 1.8     | 1.9     | 1.8    | 6.2         | 98.3   | 89.1   | 87.6| 81.8     | 77.3    | 76.8    |
| K05-GS (1)     | 0.7     | 0.8     | 0.7    | 6.0         | 1.8    | 89.4   | 88.0| 81.8     | 77.4    | 76.6    |
| FU (2)         | 12.2    | 12.2    | 12.1   | 13.6        | 12.2   | 11.7   | 88.1| 81.9     | 78.1    | 77.0    |
| Nakayama (3)   | 13.8    | 13.5    | 13.7   | 15.3        | 14.1   | 13.6   | 13.3| 83.0     | 77.7    | 77.5    |
| JKT6468 (4)    | 21.8    | 21.1    | 21.3   | 21.3        | 21.7   | 21.7   | 21.5| 20.1     | 77.8    | 77.1    |
| 10-1827 (5)    | 27.6    | 27.4    | 27.7   | 27.6        | 27.5   | 27.3   | 26.9| 26.8     | 90.0    |        |
| Muar (5)       | 28.2    | 28.6    | 28.7   | 28.4        | 28.1   | 28.5   | 27.9| 27.2     | 27.7    | 11.1    |

The upper triangle represents similarity while the lower triangle represents divergence. A8.789, A10.825, A10.881, and 10-1827 represent the ROK mosquito pools. JEV reference strains are shown for 5 genotypes: I (K01-JN, K05-GS), II (FU), III (Nakayama), IV (JKT6468), and V (Muar). Percent similarity/divergence was computed using the MegAlign program (Lasergene v.8 software, USA). Numbers in parentheses represent the JEV genotype.
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Authors’ contributions
RT and BPE conceived the study, the design, and drafted the manuscript. RT, BT, and AK carried out all molecular work. HCK, WJL, and TAK collected the mosquitoes and assisted in drafting the manuscript. JG appropriated funding (program protocols) and participated in conducting the study. All authors read and approved the final manuscript.

Competing interests
The authors declare that they have no competing interests.

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Table 3 Origin of 30 JEV strains referenced in this study

| Strain     | Location     | Year   | Host   | Genotype | Accession no. |
|------------|--------------|--------|--------|----------|---------------|
| 1070/82_Subin | Thailand     | 1982   | Human  | 1        | GQ902059      |
| 90VN70     | Vietnam      | 1990   | Human  | 1        | HM228921      |
| B-0860/82  | Thailand     | 1985   | Swine  | 1        | GQ902058      |
| Beijing-1  | China        | 1949   | Mosquito | 3     | L48961, FJ872376 |
| Bennett    | Korea        | before 1951 | Human | 2        | FJ51927      |
| FU         | Australia    | 1995   | Human  | 2        | AF217620      |
| GZ56       | China        | 2008   | Human  | 1        | HM366552      |
| Ishikawa   | Japan        | 1998   | Mosquito | 1     | AB051292      |
| JEV/sw/Mie/40/2004 | Japan | 2004 | Swine  | 1        | AB241118      |
| JEV40783   | Korea        | before 1971 | Human | 3        | FJ51923      |
| JKT6468    | Indonesia    | 1987   | Mosquito | 4     | AY184212      |
| JKT7003    | Indonesia    | 1981   | Mosquito | 4     | U70408       |
| JX61       | China        | 2008   | Swine  | 1        | GU56217       |
| KO1-JN     | Korea        | 2001   | Mosquito | 1     | FJ938222      |
| K87P39     | Korea        | 1987   | Mosquito | 3     | AY585242      |
| K98A07     | Korea        | 1988   | Mosquito | 3     | FJ938227      |
| K91P55     | Korea        | 1991   | Mosquito | 4     | U34928       |
| K94P05     | Korea        | 1994   | Mosquito | 1     | AF045551      |
| K05-GS     | Korea        | 2005   | Mosquito | 1     | FJ938223      |
| KPP82-39-214CT | Thailand | -     | Mosquito | 3     | GQ902063      |
| KV1899     | Korea        | 1999   | Swine  | 1        | AY316157      |
| Muar       | Malaysia     | 1952   | Human  | 5        | HM596272      |
| MVE-1-51   | Australia    | 1951   | Human  | 3        | AF161266      |
| Nakayama   | Japan        | 1935   | Human  | 3        | EF571853      |
| SC04-17    | China        | 2004   | Mosquito | 1     | GU187972      |
| SH17M      | China        | 2007   | Mosquito | 1     | EU429297      |
| T1P1       | Taiwan       | 1997   | Mosquito | 3     | AF254553      |
| XJ69       | China        | 2007   | Mosquito | 1     | EU880214      |
| XJP613     | China        | 2007   | Mosquito | 1     | Eu938999      |
| XZ0938     | China        | 2009   | Mosquito | 1     | HQ652538      |

All strains are JEV, with the exception of MVE-1-51, a strain of Murray Valley encephalitis virus.

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