Japanese encephalitis virus RNA detected in Culex pipiens mosquitoes in Italy

P Ravanini (paolo.ravanini@gmail.com), E Huhtamo, V Ilaria, M G Crobu, A M Nicosia, L Servino, F Rivasi, S Allegrini, U Miglio, A Magri, R Minisini, O Vapalahti, R Boldorini

1. Laboratory of Molecular Virology, Azienda Ospedaliero-Università di Carità, Novara, Italy
2. Infection Biology Research Programme, Department of Virology, Haartman Institute, Faculty of Medicine, University of Helsinki, Helsinki, Finland
3. Department of Diagnostics, Laboratory Service and Forensic Medicine, Section of Pathological Anatomy, University of Modena and Reggio Emilia, Italy
4. Department of Pathological Anatomy, Faculty of Medicine and Surgery, University Amedeo Avogadro del Piemonte Orientale, Novara, Italy
5. Department of Translational Medicine, University of Eastern Piedmont ‘Amedeo Avogadro’, Novara, Italy

Mosquitoes collected in northern Italy were screened for flavivirus RNA. Positive amplicons were sequenced and found most similar to insect flavivirus (ISF), Usutu virus (USUV) and surprisingly also to Japanese encephalitis virus (JEV). The sequence (167 bp), obtained from one pool of Culex pipiens, was found identical to JEV strains from bats in China. Unfortunately additional sequence data or virus isolations were not obtained in this study. Confirmation of potential introduction of JEV to Italy and other European countries is urgently needed.

In the course of a small-scale preliminary study screening for the presence of flavivirus RNA in mosquitoes in Italy, we obtained sequences of three different flaviviruses; an insect-specific flavivirus (ISF) related to cell fusing agent virus, Usutu virus (USUV) and surprisingly also to Japanese encephalitis virus (JEV). The sequence (167 bp), obtained from one pool of Culex pipiens, was found identical to JEV strains from bats in China. Unfortunately additional sequence data or virus isolations were not obtained in this study. Confirmation of potential introduction of JEV to Italy and other European countries is urgently needed.

Sample collection

Following the active circulation of WNV and USUV, the recent detection of novel ISFs in Italy and elsewhere in Europe, and the detection of dengue virus in southern France and Croatia [1,7,10,11], the aim of this study was to screen mosquitoes for flavivirus RNA using a system allowing the detection of all flaviviruses. Female mosquitoes were collected in late summer of 2010 and 2011 in rural areas near Modena and Bologna in Emilia-Romagna region (Figure 1), using CO2-baited traps.

Mosquitoes were identified using morphological characteristics [5], pooled by species (identification at subspecies level was not done), date and site of collection (with a maximum of 27 individuals per pool) and stored at -80°C until processed. The mosquito species collected included mainly C. pipiens, and additionally Aedes albopictus, A. caspius and A. vexans. A total of 62 pools were studied; 52 had been collected in 2010 (all C. pipiens) and 10 in 2011.

Molecular analysis

The mosquitoes were ground manually using sterile sand and Dulbecco’s phosphate-buffered saline. Nucleic acids were extracted using EasyMag (bioMérieux) and examined by RT-PCR targeted to a conserved region of the flavivirus NS5 gene [6]. The PCR products were sequenced directly and cloned when necessary (CloneJET PCR Cloning Kit, Fermentas). The obtained sequences were identified using BLAST (blast.ncbi.nlm.nih.gov/Blast.cgi).

Of the 62 pools, five were found positive for flavivirus RNA. The sequences (Box) were identified as (i) ISF from A. albopictus, 2011 (two sequences; lengths 133 and 87 bp; identical in the overlapping region), (ii) USUV from C. pipiens, 2010 and 2011 (two sequences; lengths 133 and 167 bp; identical in the overlapping

Citation style for this article:
region) and (iii) JEV (one sequence; 167 bp) from one pool of C. pipiens, 2010 (Figure 2). The ISF (collected in Sasso Marconi) and USUV (collected in Pontecchio Marconi and Pianoro) sequences were identical to other sequences previously reported from Italy [1,7]. The JEV sequence (genomic position: 9,109–9,275) was obtained from mosquitoes collected in Sasso Marconi. It showed 100% similarity to four sequences in Genbank, all of them representing JEV genotype III viruses isolated from bats in China between 1986 and 2009 (JN711458, JN711459, JF706285, JF185036).

The PCR product yielding the NS5 sequence related to JEV was amplified from the original material twice and sequenced in three separate laboratories. Additional sequence data would be needed for detailed characterisation of the viral strain and sequence analysis, but unfortunately the attempts to amplify longer sequences from the JEV-positive pool using primers targeted to E, NS5 and NS3 regions in nested and semi-nested protocols remained negative. Attempts to isolate the virus from the JEV- and USUV-positive pools on Vero and on C6/36 insect cells were not successful.

Discussion
While the potential risk of JEV spreading to Europe has been acknowledged before [8], and despite the active surveillance for flaviviruses such as WNV and USUV, to the best of our knowledge, this is the first report of a JEV-like sequence in mosquitoes in Europe. The JEV-like sequence was detected within a small scale preliminary study, and some details of the field work along
with mosquito subspecies identification were unfortunately not documented in detail. Laboratory contamination as the source of the obtained JEV sequence was highly unlikely, as no JEV virus, RNA or PCR products had ever been handled in the laboratories where the mosquitoes were processed or where the RT-PCRs were performed. Interestingly, Mani et al. have reported detecting in 1996-97 JEV antibodies and RNA in Italian birds [9], but unfortunately no further information is currently available about the sequences found in that study.

Recently, autochthonous dengue virus (DENV) infections have been detected in France [10] and Croatia [11]. While these viruses are most likely to have been imported there from endemic regions, most probably through viraemic travellers or via materials harbouring infected mosquitoes, eggs or larvae, JEV could have been introduced to Italy through waterfowl or wild waterbirds. Future arbovirus surveillance should include JEV-specific or pan-flavivirus detection methods, and it should be noted that due to cross-reactions, serological assays with the exception of seroneutralisation are probably unable to differentiate an immune response to JEV from one to WNV and USUV.

Conclusions

A partial genomic sequence of JEV was detected in Italian C. pipiens mosquitoes for the first time, but confirmation of the finding by additional sequence data or virus isolation has not yet been successful. The authors are aware that these findings are preliminary, and confirmation of the results is necessary. Further evidence of JEV circulation is required for evaluating the possible need for precautionary measures against JEV transmission in Italy and other European countries.

Box

Viral nucleotide sequence fragments obtained from mosquitoes collected in Italy, summer 2010 and 2011 (n=3)

<table>
<thead>
<tr>
<th>Virus</th>
<th>Pool</th>
<th>Length</th>
<th>Sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>JEV</td>
<td>pool_M20</td>
<td>167 bp</td>
<td>TCATGTCGTGGGCGCAATCTCAGGTTAGGCTGAGGAGAATTCAGGAGGTGGAGTGGAAAGGCTCAGGCGTCCAAAAGCTAGGATACATCCTCCGTGACATAGCAGGAAAGCAAGGAGGGAAAAATGAC</td>
</tr>
<tr>
<td>USUV</td>
<td>pool_M7</td>
<td>167 bp</td>
<td>TCATGTCGTGGGCGCAATCTCAGGTTAGGCTGAGGAGAATTCAGGAGGTGGAGTGGAAAGGCTCAGGCGTCCAAAACCTGGTTGTCACATTGTGACATTGGTGGAGGAGGAGGAGGTGGAGTGGAAAGGCTCAGGCGTCCAAAAGCTAGGATACATCCTCCGTGACATAGCAGGAAAGCAAGGAGGGAAAAATGAC</td>
</tr>
<tr>
<td>ISF</td>
<td>pool_M2B</td>
<td>133 bp</td>
<td>CTCGGAAGTCGTTTTCTGGGCTGAGGAGGAGGAAAAATGAC</td>
</tr>
</tbody>
</table>

Figure 2

Phylogenetic tree based on a 122 bp region of flavivirus NS5 sequences obtained from mosquitoes collected in Italy, summer 2010 and 2011 (n=3)
Acknowledgments

We thank Alberto Palandri for the graphic support.

References


