Imported dengue virus serotype 1 from Madeira to Finland 2012

Huhtamo, E.

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Imported dengue cases originating from the Madeiran outbreak are increasingly reported. In 2012 five Finnish travellers returning from Madeira were diagnosed with dengue fever. Viral sequence data was obtained from two patients. The partial C-preM sequences (399 and 396 bp respectively) were found similar to that of an autochthonous case from Madeira. The partial E-gene sequence (933 bp) which was identical among the two patients grouped phylogenetically with South American strains of dengue virus serotype 1.

Between August and December 2012, the hospital district of Helsinki and Uusimaa laboratory (HUSLAB) which solely performs dengue diagnostics in Finland, received samples from 16 patients with clinically suspected dengue fever who had visited the Autonomous Region of Madeira, Portugal, where an outbreak of dengue fever is taking place since the beginning of October 2012 [1,2].

Confirmation of dengue virus infection

The samples of the patients were analysed by dengue non-structural protein (NS)1 antigen (Ag) strip test (BioRad), IgM-enzyme immunoassay (EIA) (Focus Technologies) and an in-house IgG immunofluorescence assay (IFA) using dengue virus (DENV)-3 infected Vero E6 cells as antigens. The dengue diagnosis was confirmed in five patients by serological tests. Three of five serologically-confirmed patients were also positive for NS1 Ag.

The earliest available samples of the five serologically-confirmed dengue patients were studied by reverse transcription-polymerase chain reaction (RT-PCR) to obtain information of the infective virus strain. RNA was extracted from serum samples using QIAamp viral RNA mini kit (Qiagen) according to manufacturer’s instructions. The RNA was used as a template in dengue typing RT-PCR [3] and in an RT-PCR amplifying a partial sequence of the envelope gene (E-gene) [4]. RT-PCR products were obtained from two patients, for which the typing RT-PCR product size corresponded to DENV serotype 1 (DENV-1) [3].

The obtained RT-PCR products were purified enzymatically using ExoSAP-IT (USB laboratories) and directly sequenced (in case of E-gene, using DENV-1 E-gene specific primers). Primer sequences are available from the authors upon request. The obtained sequences of the E-gene (933 bp) from the two patients were identical to one another and the sequence information was deposited in GenBank under accession number: KC567679. Sequences from the capsid-pre-membrane (C-preM) region [3] derived from the first patient (399 bp, GenBank accession number: KC616348) and the second patient (396 bp, GenBank accession number: KC616349) differed from each other by one nucleotide.

The sequences from the patients were compared to available DENV sequences in GenBank database and the basic local alignment search tool (BLAST) results confirmed that the infective virus was DENV-1. This finding was in line with the typing RT-PCR results and with previous reports [1,5,6].

Phylogenetic analysis of the viral sequence derived from the patients

All of the four known serotypes of DENV are further genetically categorised into several distinct genotypes. Two different ways of categorising DENV-1 strains have been used: into three [7-9] or into five different genotypes [10-12]. The DENV-1 viruses include a sylvatic strain originating from Malaysia (designated as sylvatic genotype or genotype III). In the five genotype
classification, some strains originating from Thailand in the 1950s and 1960s are designated as genotype II.

All the rest of the epidemic strains can be divided into three main groups corresponding to a three genotype classification. These groups include: (i) mainly Asian strains and some east African strains designated as genotype I (or Asian I genotype in the five genotype classification), (ii) strains originating mainly from the west Pacific islands and Australia designated as genotype II (or South Pacific genotype IV in the five genotype classification), and (iii) strains originating from Africa, some Asian strains and all strains originating from the Americas designated as genotype III (or American/African genotype V in the five genotype classification). To identify dengue virus strain(s) responsible for a given outbreak, characterisation of the infecting virus by sero- and genotyping is needed. Although dengue viruses currently have a global circulation, their genetic characterisation likely provides clues of their origins.

The E-gene sequence derived from the patients was aligned with sequences sharing highest nucleotide homologies in GenBank and with a global collection of representatives of different recent epidemic DENV-1 genotypes using multiple sequence comparison by log expectation (MUSCLE) (www.ebi.ac.uk/Tools/msa/muscle/) and a neighbour-joining phylogenetic tree was estimated with the maximum composite likelihood method using the programme MEGA (www.megasoftware.net/) (Figure).

The results suggested that the Madeiran strain represented the American/African genotype (designated as genotype III or V). In the phylogenetic tree, the Madeiran strain was clustered together with strains from Colombia and Venezuela. The highest nucleotide similarities were shared between the Madeiran strain and a strain from Colombia from 2008 (99.9%). The Madeiran strain had a single nucleotide synonymous substitution (T®C) in comparison to the Colombian strain at position 2,040 (GenBank accession number: GQ868570).

Additional sequences from the C-preM region [3] were also compared to Genbank sequences and to the available sequence of the autochthonous case from Madeira that became recently available [6]. The sequence from the first patient was identical to the sequence from the autochthonous case within the alignable region of 385 bp, and the sequence from the second patient had a single nucleotide change (position 82, GenBank accession number: KC248375). The BLAST search results for the C-preM sequences were in line with those of the E-gene sequence, showing highest sequence homologies to strains from South America.

Discussion and conclusion
Our results are in line with previous reports [1,5,6], and all data thus far suggests that possibly a single strain of DENV-1 likely originating from South America caused the Madeiran epidemic, although more sequence data, e.g. complete genome data and sequences from more patients of the outbreak would be needed to confirm these findings.

Interestingly, DENV-1 was also identified in 2010 from autochthonous cases in southern France [13] and in Croatia [14,15]. The available sequences from Croatia [15] did not represent the same region as our sequence, and thus could not be included in our analysis. The analysis of the sequences from Croatia was based on short E-NS1 junction sequences (222 bp). The results suggested that the Croatian DENV-1 also represented genotype III (a genotype also designated as American/African genotype V), but grouped together with strains from India [15]. As the Croatian sequences were associated with strains from India rather than strains from South America, it seems unlikely that the Croatian strains would be closely related to the strain circulating in Madeira. However, more comprehensive sequence data would be needed from the autochthonous cases found previously in Europe for comparisons.

Aside from the autochthonous cases of dengue fever in southern France [13] and Croatia [14], the Madeiran event is the first epidemic of dengue fever in Europe since 1928, when Greece was affected by an outbreak [16,17]. The 2012 Madeiran epidemic has had a great impact on local residents, but has also affected visitors of the island. The latest available data from February 2013 reports that 78 visitors from European countries got infected in Madeira [18]. These countries, listed according to decreasing number of cases, include the United Kingdom, Germany, mainland Portugal, Finland, Sweden, and France, followed by Denmark, Austria and Norway, and finally Croatia, Slovenia, Spain and Switzerland. The number of imported cases in Europe is however likely to be an underestimate. Also the actual number of persons in Finland, who got infected by DENV in Madeira may be higher than the number reported here as the travel history information was not available for all the suspect dengue patients whose samples were tested at the HUSLAB.

Whereas the risks for possible endemic transmission of dengue virus (DENV) in Europe have been mainly associated with the spreading of Stegomyia albopicta (=Aedes albopictus) [13,14], interestingly, the vector associated with the Madeiran outbreak was reported to be Stegomyia aegypti (=Aedes aegypti) [1] as was the 1928 outbreak in Greece [16,17] (mosquito nomenclature according to Reinert et al., 2009 [19]).

Similarly to Madeira, it has been suggested that island locations that are major tourist attractions such as Mauritius [20] may be at elevated risk for dengue introduction via travellers if suitable vectors are available. Although significant efforts have been taken to map and collect the information of the relevant mosquito vectors of mosquito-transmitted diseases in Europe.
Phylogenetic analysis of partial envelope gene sequence derived from two imported dengue cases from Madeira to Finland, 2012

The tree, determined by neighbour-joining method, was based on partial (933 bp) envelope gene sequences of dengue viruses of serotype 1. Bootstrap support values >60% are shown at the nodes. The scale bar represents nucleotide substitutions per site. The GenBank accession numbers of the sequences used as well as the country of origin and year are shown on the tree. The strain from the case imported from Madeira in 2012 is indicated in bold and clusters with strains of genotype III (a genotype also designated as American/African genotype V).
by e.g. the European Centre for Disease Prevention and Control (ECDC) VBORNET, this information is not available from all locations. For example, according to ECDC VBORNET mosquito maps [21] no information is currently available on competent dengue vectors, (Stegomyia albopicta (=Aedes albopictus) or Stegomyia aegypti (=Aedes aegypti)) for some locations situated relatively closely to Madeira, such as the Canary Islands. The Madeirian outbreak highlights the importance for obtaining up-to-date information of the vector species distribution in Europe and urges the need for planning countermeasures and surveillance at virological, medical and environmental levels [22].

Although the Madeiran dengue outbreak was widely reported in the international news, dengue may still not be generally suspected in intra-European travelers. The possible endemisation of dengue in Europe highlights the need for diagnostic preparedness and awareness of clinicians to suspect and recognise dengue originating from within Europe. We conclude that the detailed molecular characterisation of the DENV strains from travellers, autochthonous cases and outbreaks are needed for studying the molecular epidemiology and transmission chains of dengue virus strains in Europe.

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Conflict of interest
None declared.

Authors’ contributions
All authors contributed to the study design, interpretation of the results, writing of the manuscript and approved the final version. EMK performed RNA extractions and RT-PCR tests. EH performed the NS1 antigen tests, sequence analysis and wrote the first version of the manuscript. OV collected the diagnostic and clinical data.

References