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Published in:
Genome Announcements

Link to article, DOI:
10.1128/genomeA.01015-17

Publication date:
2017

Document Version
Publisher's PDF, also known as Version of record

Link back to DTU Orbit

Citation (APA):
Ariel, E., Subramaniam, K., Imnoi, K., Sriwanayos, P., Ahasan, M. S., Olesen, N. J., ... Waltzek, T. B. (2017). Genomic Sequencing of Ranaviruses Isolated from Edible Frogs (Pelophylax esculentus). Genome Announcements, 5(38), e01015-17. DOI: 10.1128/genomeA.01015-17
Genomic Sequencing of Ranaviruses Isolated from Edible Frogs (*Pelophylax esculentus*)

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ABSTRACT Ranaviruses were isolated from wild edible frogs (*Pelophylax esculentus*) during epizootics in Denmark and Italy. Phylogenomic analyses revealed that these isolates are closely related and belong to a clade of ranaviruses that includes the *Andrias davidianus* ranavirus (ADRV), common midwife toad ranavirus (CMTV), *Testudo hermanni* ranavirus (THRV), and pike-perch iridovirus (PPIV).

The earliest reported occurrence of ranaviral disease in amphibians in Europe resulted in recurrent, low-level mortality in wild-caught edible frogs in a commercial operation in Croatia from 1970 to 1981 (1). Other ranaviral isolates from edible frogs have been obtained from tadpoles collected from the wild during disease outbreaks in Italy in 2002 (REV 282/I02) and from free-living adults collected in Denmark in 2008 (PEV_DK1) (2, 3). Mortality events in *Rana temporaria* in the United Kingdom in the 1990s (4) and, more recently, in common midwife toads (*Alytes obstetricans*) in Spain (5) and *Pelophylax* spp. in The Netherlands (6) have been added to the list of amphibian ranaviral outbreaks in Europe.

The two isolates PEV_DK1 and REV 282/I02 were amplified in epithelioma papulosum cyprini cells until the cytopathic effect was complete, and then the resulting supernatant was clarified and the total nucleic acid was purified from the clarified supernatant, as previously described (7). DNA libraries were prepared using the Nextera XT DNA kit (Illumina), and sequencing was performed using a v3 chemistry 600-cycle kit on an Illumina MiSeq platform. De novo assembly of the paired-end reads in SPAdes (8) produced contiguous consensus sequences of 107,392 bp with G/H11001C content of 55.31% and 107,444 bp with G/H11001C content of 56.03% for PEV_DK1 and REV 282/I02, respectively.

The genomes of the two isolates were annotated using Genome Annotation Transfer Utility (GATU) (9) with *Frog virus 3* (GenBank accession no. NC_005946) as the reference genome. Additional putative open reading frames (ORFs) were identified using GenemarkerS (10), and gene functions were predicted based on BLASTP searches against the NCBI GenBank non-redundant protein sequence database. A total of 99 putative ORFs were predicted in PEV_DK1 and 101 in REV 282/I02 compared to 101 ORFs in *Andrias davidianus* ranavirus (ADRV) (Genbank accession no. KC865735), 104 ORFs in common midwife toad ranavirus (CMTV) (GenBank accession no. JQ231222 and KP056312), 75 in *Testudo hermanni* ranavirus (THRV) (GenBank accession no. KP266741), and 109 in pike-perch iridovirus (PPIV) (GenBank accession no. KX574341). Comparative genomic analyses revealed that these two ranaviruses are closely related, except for a nonsense mutation in a gene encoding a hypothetical protein (orthologous to ORF51.

Received 14 August 2017 Accepted 18 August 2017 Published 21 September 2017


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in REV 282/102) and the absence of a gene encoding a hypothetical protein (orthologous to ORF95 in REV 282/102) in PEV_DK1. An analysis of locally collinear blocks (LCB) in Mauve (11) revealed that the genomes of PEV_DK1 and REV 282/102 display the same genome arrangement as ADRV, CMTV, THRV, and PPIV (12). Maximum likelihood phylogenetic analyses based on the concatenated genome-wide LCB alignments revealed that the Italian and Danish ranaviral isolates from edible frogs belong to a clade of ranaviruses that includes ADRV, CMTV, THRV, and PPIV.

The repeated isolation, separated by time and space, of nearly identical ranaviral strains from edible frogs indicates that this species may serve as an important natural host. The detection of closely related strains in Chinese giant salamander (Andrias davidianus), common midwife toad (Alytes obstetricans), Hermann’s tortoise (Testudo hermanni), and pike-perch (Sander lucioperca) underscores the low host specificity of these ranaviruses.

Accession number(s). The complete genome sequences of PEV_DK1 and REV 282/102 have been deposited in GenBank under the accession numbers MF538627 and MF538628, respectively.

ACKNOWLEDGMENT
We thank Patrick Thompson for his technical assistance.

REFERENCES