might lead to recruitment of more host and inflammatory cells that further amplify viral replication and organ injury (6). Downregulation of several factors highlights the damage. For instance, the VEGF-A levels mirror the impairment of spermatogonia, primary spermatocytes, and Sertoli cells upon Zika virus infection (4). However, the decrease in CXCL-1, CXCL-8, and CXCL-10 levels in semen during infection could indicate a local immunosuppressive state induced by infection, limiting immune cell infiltration in the MRT and potentially virus dissemination throughout the body.

The different kinetics of virus replication and cytokine secretion in semen samples raises questions about the semen secretome in cases of couples aiming for conception and the necessity to extend the convalescence period beyond Zika disease recovery. In fact, at high concentration, most of these factors might alter the integrity of the mucosal barriers within the female reproductive tract and increase a woman’s susceptibility to infection (7). They might also promote peroxidation and affect sperm function, potentially resulting in infertility (8).

The limitations of our study include small sample size. Further investigations with a larger cohort of patients and controls are warranted.

In summary, a profound disruption in the cytokine network is evident in plasma and semen starting at the earliest stage of Zika virus infection and is maintained over time even after viral clearance. Studies to characterize the mechanism involved in the establishment of compartmentalization and develop efficient antiviral therapies that interfere with virus replication in the MRT are needed.

Acknowledgment
We thank Isabelle Da Silva for technical assistance.

About the Author
Dr. Mansuy is a medical virologist at the University Hospital of Toulouse in Toulouse, France. His research interests include emerging viral diseases and respiratory and enteric diseases.

References

Detection of Epizootic Hemorrhagic Disease Virus Serotype 1, Israel

Natalia Golender,1 Velizar Y. Bumbarov1

Author affiliation: Kimron Veterinary Institute, Beit Dagan, Israel

DOI: https://doi.org/10.3201/eid2504.180149

During September 2016–February 2017, we detected epizootic hemorrhagic disease virus (EHDV) in ruminants in Israel. BLAST and phylogenetic analyses of segment 2 in 6 EHDVs isolated from field samples indicated a close relationship to the EHDV serotype 1 strain in Nigeria. Affected cattle had mostly mild or asymptomatic disease.

Epizootic hemorrhagic disease is an infectious, non-contagious viral disease of ruminants, transmitted by insects of the genus Culicoides; it mostly affects white-tailed deer and cattle. Epizootic hemorrhagic disease virus (EHDV) belongs to the genus Orbivirus within the family Reoviridae and is closely related to bluetongue virus (BTV) and African horse-sickness virus. At least

1Both authors share first authorship.
7 EHDV serotypes are currently recognized worldwide (1). (EHDV serotype 1 has been isolated from cattle in the Northern Territory of Australia (2), Nigeria, Ecuador, French Guiana, Réunion Island (in the Indian Ocean) (3), and the United States, where it was also isolated from white-tailed deer (4). In cattle in Israel, EHDV serotype 7 first was recognized in 2006, when it manifested in serious clinical signs (5). EHDV serotype 6 caused an outbreak in 2015, when clinical signs were milder than those associated with EHDV serotype 7 (6). However, EHDV serotype 6 RNA was found in placenta and brains of aborted cattle fetuses during the latter outbreak (6).

In the summer and fall of 2016, several arbovirus infections were registered simultaneously in diseased domestic and wild ruminants in Israel. Included were infections with BTV serotypes 2, 3, 4, 8, and 15; Shuni and Akabane viruses, belonging to the Simbu serogroup of genus Orthobunyavirus in the Peribunyaviridae family; and EHDV. BTV serotype 8 caused a large outbreak that caused heavy losses of livestock in some sheep and cattle farms.

During September 2016–February 2017, we routinely tested 265 field samples by using specific EHDV real-time reverse transcription PCR (rRT-PCR), as described previously (6). The tested field samples included 10 spleen samples from wild and zoo ruminants taken from 4 mountain gazelles, 1 giraffe, 1 Arabian oryx, and 4 Nubian ibexes, as well as 13 spleen samples and 242 whole-blood EDTA samples from diseased cattle. We similarly tested 22 aborted cattle fetuses. We obtained the first EHDV-positive whole-blood sample from a diseased dairy cow on September 12, 2016, and the most recent one on February 26, 2017.

A total of 81 EHDV-positive field samples originated from northern Israel (the Golan Heights, Galilee, the Sharon Plain, and the Jordan Valley) and central Israel (the Coastal Plain). Positive samples included 1 spleen sample from a wild mountain gazelle that was found dead as a result of a head wound near the Sea of Galilee and 2 spleen samples and 78 whole-blood samples from infected cattle. However, the deaths of EHDV-positive cattle probably were not caused by EHDV. In 1 such case, in an adult dairy cow, Escherichia coli from all tested internal organs was isolated, and Babesia spp. were found during microscopic examination. In another case, in a 4-month-old calf, we identified BTV by using a BTV-specific rRT-PCR test (VetMAX BTV NS3 All Genotypes Kit, LSI; Thermo Fisher Scientific, https://www.thermofisher.com), which led to isolation of BTV serotype 8.

Among the EHDV rRT-PCR–positive samples, 18 were also rRT-PCR–positive for BTV and 1 was positive in pan-Simbu rRT-PCR (7), showing Akabane virus infection. All tested aborted cattle fetuses were EHDV-negative by rRT-PCR. We attempted virus isolation on all EHDV- and BTV-positive samples. We isolated 6 EHDV in embryonated chicken eggs, which consequently passed on BHK-21 cells, as previously described (6).

We sequenced 4 partial and 2 complete sequences of segment 2 (encoding viral protein 2) by using the standard Sanger method and submitted them to GenBank (accession

![Figure](https://www.cdc.gov/eid/) Phyllogenetic analysis based on full-length sequences of segment 2 in 2 EHDV serotype 1 isolates from Israel with global EHDVs and BTV-8 from GenBank. We analyzed 24 nucleotide sequences and inferred phylogenetic relationship by using the neighbor-joining method. Numbers below branches indicate bootstrap values. Recent isolates from Israel are marked with black circles. Viruses are identified by GenBank accession number, virus and serotype, 3-letter code of country (and additional information in some cases), isolate identification (in most cases), and year of isolates. Evolutionary analyses were conducted in MEGA7 (8). Scale bar indicates number of nucleotide substitutions per site. BTV, bluetongue virus; EHDV, epizootic hemorrhagic disease virus.

Emerging Infectious Diseases • www.cdc.gov/eid • Vol. 25, No. 4, April 2019
nos. MG808405–MG808410). BLAST (https://blast.ncbi.nlm.nih.gov/Blast.cgi) and phylogenetic analyses of the segment 2 sequences showed that the EHDV recently isolated in cattle in Israel belongs to serotype 1 and is closely related to the lbaAr22619 strain from Nigeria, with which it shares 95.72%–95.76% identity (Figure).

Retrospective analysis of clinical signs in EHDV-1–infected cattle enabled us to conclude that in many farms EHDV infection was asymptomatic or subclinical; milk-yield reduction, fever, and recumbency were the only prominent clinical signs observed during the outbreak. However, animals with BTV and EHDV co-infections showed more severe clinical signs, including fever, abortion, lameness, subcutaneous emphysema, and death.

During recent years, several new arboviruses have been detected in Israel that were not identified previously. BTV serotype 3 was first identified in 2016 but probably was present in Israel since at least 2013 (9), EHDV serotype 6 was identified in 2015 (6), EHDV serotype 1 was found in 2016, and Shuni virus was detected in 2014 (10). These findings showed that new introductions of arthropodborne viral infections into the Middle East region had occurred. Molecular epidemiologic data indicate the viruses originated in Africa, as ours and other studies (5,6) have shown. Molecular diagnostics, vector-control strategies, and epidemiologic studies should be implemented in Israel to mitigate potential risk for future outbreaks.

About the Authors
Drs. Golender and Bumberov are virologists in the Virology Department of the Kimron Veterinary Institute, Beit Dagan, Israel. Their primary research interests include the investigation of arboviral infections caused by viruses of the Reoviridae (Orbiviruses) family and the Peribunyaviridae (Orthobunyavirus, Simbu serogroup viruses) family, which affect ruminant populations, and developing diagnostic methods to detect these viruses.

References


Address for correspondence: Natalia Golender, Kimron Veterinary Institute, Beit Dagan 50250, POB 12, Israel; email: golendeder@moag.gov.il


Maite Aubry, Mike Kama, Jessica Vanhomwegen, Anita Teissier, Teheipuaura Mariteragi-Helle, Stephane Hue, Martin L. Hibberd, Jean-Claude Manuguerra, Ketan Christi, Conall H. Watson, Eric J. Nilles, Colleen L. Lau, John Aaskov, Didier Musso, Adam J. Kucharski, Van-Mai Cao-Lormeau

Author affiliations: Institut Louis Malardé, Papeete, French Polynesia (M. Aubry, A. Teissier, T. Mariteragi-Helle, D. Musso, V.-M. Cao-Lormeau); Fiji Centre for Communicable Disease Control, Suva, Fiji (M. Kama); The University of the South Pacific, Suva (M. Kama, K. Christi); Institut Pasteur, Paris, France (J. Vanhomwegen, J.-C. Manuguerra); London School of Hygiene and Tropical Medicine, London, UK (S. Hue, M.-L. Hibberd, C.H. Watson, A.J. Kucharski); World Health Organization Division of Pacific Technical Support, Suva (E.J. Nilles); Harvard Medical School and Brigham and Women’s Hospital, Boston,

Emerging Infectious Diseases • www.cdc.gov/eid • Vol. 25, No. 4, April 2019