the recombination with nonpolio enteroviruses is important. These VDPVs have major implications for the cessation of immunization with OPV after certification that wild PV has been eradicated.

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West Nile Virus Infection in Crocodiles

To the Editor: Recently West Nile virus (WNV) infection has been reported in three alligators (Alligator sp.) from central Florida (1) and one captive crocodile monitor (Varanus salvadori) with neurologic signs from the District of Columbia and Maryland area (2). These first reports of the virus in American reptiles highlight the possible role of this group of vertebrates in the WNV life cycle. To our knowledge, WNV in a reptile was reported only once before in a serosurvey conducted in Israel from 1965 to 1966, in which 22 reptiles and 96 amphibians were tested for hemagglutination-inhibiting antibodies against several viruses, including WNV; one turtle (Clemmys caspica) was seropositive (3). Experimental infection of the lake frog (Rana ridibunda) with a Russian strain of WNV resulted in high levels of viremia (4). At present, the role of reptiles and amphibians in the life cycle and epidemiology of WNV is not known.

We report, for the first time, WNV infection in crocodiles (Crocodylus niloticus). To assess the potential role of crocodiles in the life cycle of WNV in Israel, serum specimens were collected from 20 healthy crocodiles on a commercial farm in the Negev Desert, in southern Israel (31°14′N, 34°19′E). The crocodiles came from two separate breeding farms (32°03′N, 35°26′E and 30°18′N, 35°07′E) in the Syrian-African Rift Valley, which is on the main route of bird migration from Africa to Europe. Five males and 15 females, 1–2.5 years of age, were examined. Blood was withdrawn from the crocodiles’ ventral caudal vein, separated by centrifugation, and kept at –20°C until analyzed. Neutralizing antibody titers were determined against WN-goose-98 (5) and attempts to isolate the virus were performed by using Vero cell culture (6) and by using direct reverse transcription–polymerase chain reaction (RT-PCR) on the serum specimens. To eliminate the possibilities of nonspecific reaction, all serum samples were concurrently tested for the only other flavivirus known to be present in Israel; Israeli turkey meningoencephalitis virus (ITV) (7). Because ITV does not produce cytopathic effects (CPE) in Vero cells, virus neutralization was conducted on BHK cells for both WNV and ITV by using WN-goose-98 and ITV (vaccine strain). In this case, the virus stocks (10-4.2 50% tissue culture infective dose) were diluted 1:400, and virus neutralization titers were checked 3 days later.

Viral RNA was extracted from serum samples with the QIAamp RNA blood kit (QIAGEN, Valencia, CA), according to the manufacturer’s protocol and resuspended in 30 μl of RNase-free water. The primer pair WN240-Kun848 (respective genome positions 5′: 848 and 1,645) was used to synthesize an 800-bp product in the E gene region (8,9). The resulting DNA fragment was visualized on

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1.5% agarose gel stained with ethidium bromide.

The seroprevalence rate in the first set of virus neutralization assays in Vero cells was 14/20 (70%, with titers ranging from 1:20 to 1:320 [3x1:20, 3x1:40, 3x1:80, 2x1:160, 3x1:320]). No differences were discernible in either the seroprevalence rate or in the average titers of crocodiles from two different breeding farms. In BHK cells, a similar seroprevalence rate was observed, with titers ranging from 1:40 to 1:1,280 (3x1:40, 2x1:80, 1x1:160, 4x1:320, 3x1:640, 1x1:1280). All serum samples, except one, were <1:10 against ITV virus, which had a titer of 1:640 against WNV and 1:10 against ITV. Viremia was not detected in any of the 20 samples in Vero cell culture or by RT-PCR.

These results demonstrate a high rate of infection with WNV in crocodiles in Israel. The crocodiles may have been exposed to the virus during the summer at their present location, since no difference in prevalence was seen between the two groups (which differed only in the farm of origin) and since the younger crocodiles had been hatched in the spring of 2002. Furthermore, a cross-reaction with the other prevalent flavivirus in Israel, ITV, was ruled out. Preliminary results from an equine seroprevalence study (involving 800 horses over a 3-year period) of virus neutralization antibodies to WNV collected during fall 2002, indicate that most horses sampled in Israel’s Arava Valley (a desert in the Syrian-African Rift near the Jordanian and Egyptian borders) and the Gulf of Aqaba/Eilat (30°59’N, 35°18’E to 29°34’N, 34°57’E) (85%, 79/90) were positive (A. Steinman and S. Tal, unpub. data.). WNV was also isolated from mosquitoes in the same region (10). The seroprevalence of WNV antibodies among horses and local birds from the Negev Desert is not known nor is the time when the horses acquired WNV infection. However, the isolation of WNV from mosquitoes (10) and the presence of antibodies to WNV in young crocodiles demonstrate arboviral activity in this region in the summer of 2002, although clinical cases were few. That virus was not isolated from crocodiles in late November (past outbreaks of WNV in Israel mainly occurred between August and October) (6,11).

WNV has been endemic in Israel since the early 1950s (12). More recently, in the summer of 2000, an extensive outbreak occurred, affecting hundreds of people (11), dozens of horses (6), and several flocks of geese (5). However, no deaths of crocodiles were reported. This contrasts with the report from Florida (1), where WNV was isolated from dead alligators, and where hundreds of cases of sudden death had been reported in previous years; these deaths are now suspected to result, at least in part, from WNV disease. The role of various reptile species in the epidemiology of other arboviruses such as western equine encephalitis, eastern equine encephalitis, and Venezuelan equine encephalitis is well documented (13–15). At present, the role of reptiles and amphibians in the life cycle and epidemiology of WNV is not known, and further research is necessary.

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Dejà Vu

"I must have made this molecule before; It is familiar to the core!"

Said a yeast cell emerging from mitosis, With no experience yet in synthesis. Then, guided by a transmigrant human gene, It assembled that "alien" protein.

Boghos L. Artinian
Rickettsia aeschlimannii in Spain: Molecular Evidence in H. marginatum and Five Other Tick Species that Feed on Humans

To the Editor: Rickettsia aeschlimannii is a pathogenic spotted fever group rickettsia first isolated from Hyalomma marginatum ticks collected in Morocco in 1997 (1). Later found in H. marginatum ticks from Zimbabwe, Niger, Mali, and Portugal (2), R. aeschlimannii has also been found in a Rhipicephalus appendiculatus tick attached to the right thigh of a patient in South Africa (3). These data suggest a broad geographic distribution for R. aeschlimannii and the possibility that tick species other than H. marginatum may also be suitable vectors for this rickettsia.

The pathogenicity of Rickettsia aeschlimannii in humans has been demonstrated by Raoult et al. (4) in a French patient who became ill after returning from a trip to Morocco. The patient exhibited symptoms similar to those of Mediterranean spotted fever (MSF) produced by R. conorii, with a tache noire–like eschar on his ankle, fever (39.5°C), and a generalized maculopapular skin rash. The second documented and most recent case of human infection by R. aeschlimannii occurred in a South African man who was bitten by R. appendiculatus; an eschar also developed around the tick attachment site on this patient (3). He was aware of the risk for tick-transmitted disease, so he removed the tick and administered doxycycline; he did not develop additional symptoms.

Over the past 6 years, throughout the region of Castilla y León, northwestern Spain, we have collected and identified 3,059 ticks belonging to 15 species (unpub. data) that were attached to persons living in this territory. We have systematically analyzed the ticks by polymerase chain reaction (PCR) to detect those infected with Rickettsia spp., we proceeded as described by La Scola and Raoult (6): All DNA samples were first tested for a fragment of the rickettsial gltA gene (7), and then, in the gltA-positive samples, a fragment of the rickettsial ompA gene (8) was amplified, sequenced, and compared with gene databases for identification. The gltA amplicon was sequenced only when the ompA was not successfully amplified. To prevent DNA contamination and the carryover of amplified products, we used sterile tools at all times and carried out each step of the analysis (extracting DNA, preparing the reaction mixture, and amplifying and analyzing the PCR product) in separated work areas. Two negative controls (Milli-Q water and DNA from laboratory-reared noninfected ticks) were included in each amplification trial. These controls were never amplified.

We obtained 21 ompA amplicons (629–632 bp) from 21 ticks. One of these annotated to Rickettsia aeschlimannii. The pathogenicity of R. aeschlimannii in humans has never been demonstrated; it is known to infect D. variabilis ticks in Morocco (1) and R. appendiculatus ticks in South Africa (3). In northwestern Spain, we found R. aeschlimannii in 35 tick specimens belonging to H. marginatum and to another five species.

During the 6-year study, ticks found on patients who sought medical advice in the hospitals and healthcare centers of Castilla y León were removed and referred to our laboratory for identification and analysis. Each tick was first disinfected by immersion in 70% alcohol, rinsed in sterile water, and dried on sterile filter paper. We then extracted DNA in 5% Chelex-100, according to the method of Guttman et al. (5). In searching for Rickettsia spp., we used the gltA gene (7), and then, in the gltA-positive samples, a fragment of the rickettsial ompA gene (8) was amplified, sequenced, and compared with gene databases for identification. The gltA amplicon was sequenced only when the ompA was not successfully amplified. To prevent DNA contamination and the carryover of amplified products, we used sterile tools at all times and carried out each step of the analysis (extracting DNA, preparing the reaction mixture, and amplifying and analyzing the PCR product) in separated work areas. Two negative controls (Milli-Q water and DNA from laboratory-reared noninfected ticks) were included in each amplification trial. These controls were never amplified.

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