West Nile (WN) virus outbreaks were recorded in Israel during the 1950s and 1970s (2-4); however, in the last decade, diagnosis ceased and no clinical cases were reported, although seroepidemiologic surveys indicated that the virus continued to circulate (5). Following reports of illness in birds in 1998, the Central Virology Laboratory (CVL) of the Public Health Services reestablished the capability to diagnose WN virus based on serologic assays, including virus neutralization, immunofluorescence, and enzyme-linked immunosorbent assays (ELISA) with immunoglobulin (Ig) G and IgM.

This led to the identification of acute human cases beginning in the fall of 1999 (6). High seroprevalence was found in the southern region of Israel (Eilot region) in the fall and winter of 1999-2000, with IgG levels ranging from 21% to 82% and IgM levels ranging from 0% to 73% in seven communities. Some IgM-positive cases were associated with clinical symptoms compatible with WN fever. Additional acute cases were diagnosed in the central region of Israel during the spring and summer of 2000 (H. Bin, unpub. data).

During the late summer and fall of 2000, an outbreak occurred in the central and northern parts of Israel. Between August 1 and November 30, 439 patients with clinical symptoms compatible with WN fever were admitted to hospitals. The clinical details in records of the patients who died were >65 (Quarterly Report No. 3 of the Department of Epidemiology, Ministry of Health, Jerusalem). Epidemiologic and clinical aspects of the outbreak were also described by Weinberger et al. (7) and Chowers et al. (8), respectively. We describe the isolation and characterization of four viral strains from human serum obtained during this outbreak.

The Study

During the outbreak, patients’ samples (serum, cerebrospinal fluid [CSF], or both) submitted to the CVL were immediately divided into two aliquots. One aliquot was immediately used to test for IgM antibodies, and the other was stored at -70°C until further processing. Virologic studies were performed on the frozen acute-phase samples from patients who seroconverted from IgM negative to IgM positive, as well as on CSF from fatal cases. Reverse transcription-polymerase chain reaction (RT-PCR) and virus isolation were attempted simultaneously on 32 patients’ samples (17 serum and 15 CSF). RT-PCR and an indirect immunofluorescence assay (IFA) were used to confirm the presence of WN virus in cell culture. Direct sequencing of RT-PCR amplified fragments was performed to characterize the genome of isolated viruses.

Patient samples were analyzed for WN virus by RT-PCR using primer sequences for the envelope gene, as described by Lanciotti et al. and Shi et al. (9,10). Viral RNA was extracted by using the QIAamp Viral RNA Mini Kits (QIAGen GmbH, Hilden, Germany), and the RT-PCR was performed with Ready to Go Beads (Amersham Pharmacia, Buckinghamshire, England) according to manufacturer’s instructions. The primers Kun 108, Kun 848, Kun 998c, and Kun 1830c were used in RT-PCR for sequence analysis (11). Sequence analysis was performed on a 1648-bp fragment of the WN virus genome encoding 309 nt upstream from the pre-membrane protein (preM), the entire preM and membrane protein (M) genes, and 811 nt of the 5' portion of the envelope glycoprotein (E) gene. Purification of the RT-PCR product and sequence and phylogenetic analyses were performed as described (12). Both strands of the amplified PCR products were sequenced. RT-PCR conditions used were 42°C for 45 min, 95°C for 5 min, 60°C for 2 min, 72°C for 2 min, 34 cycles at 93°C for 45 sec, 55°C for 45 sec, and 72°C for 90 sec, followed by 72°C for 7 min; samples were then left at 4°C. PCR products were visualized by staining with ethidium bromide after electrophoresis on 2% agarose gels. With the RT-PCR assay, we can detect as low as 0.01 PFU based on serially diluted titered WN virus isolated from a White-eyed Gull.

Virus isolation was performed on Vero cell monolayers (ATCC CCL-81) by using the tube method. Vero cell monolayers (80%-90% confluent) were washed twice with phosphate-buffered saline, then infected with 100- to 200-μL patient samples. Patient samples were allowed to adsorb for 1 hour at 37°C with gentle swirling every 15 min. Eagle's
West Nile Virus

Virus was isolated from serum from four nonfatal WN virus IgM-negative Israeli patients who seroconverted 1 to 2 weeks later. Patient 1 (WN-0043), a 51-year-old woman from the northcentral region, was hospitalized; CNS disease did not develop. Patient 2 (WN-0233), a 20-year-old man from the north, was hospitalized for fever of unknown origin and neutropenia; CNS disease did not develop. Patient 3 (WN-0247), a 5-year-old boy residing in the central region, had meningoencephalitis and was hospitalized. Patient 4 (WN-0304), a 55-year-old woman from the north, had high fever and myalgia and no CNS symptoms; she was not hospitalized.

Two viral isolates were detected from patients 3 and 4 on day 4 after inoculation on Vero cells; the other two isolates were detected from patients 1 and 2 on day 7 after inoculation. All four virus isolates were confirmed as WN virus by IFA. Only two original acute-phase serum samples (patients 3 and 4) were positive by RT-PCR. Negative RT-PCR results and lengthy time until appearance of CPE are apparently consistent with low viral load in patients’ serum (Table).

Sequence analysis showed that isolates WN-0233 (GenBank Accession Number AF375043) and WN-0304 (GenBank Accession Number AF375045) had identical sequences over 1,648 nt and isolates WN-0043 (GenBank Accession Number AF375042) and WN-0247 (GenBank Accession Number AF375044) differed by only 1 nt. Such high homology is similar to results reported by Lanciotti et al., who also described identical WN virus sequences from brain samples from two patients (14). WN-0247 differed from WN-0043 by 50 (3%) of 1,648 nts or 25 (2.9%) of 855 nt for the partial E gene sequence. Most differences in the E gene were in the third position of the codon (21 of 25); all of these were synonymous. All four differences in the first and second codon positions encoded different amino acids when isolate WN-0247 was compared with isolate WN-0304.

A 255-nt fragment of the WN virus E gene has previously been used for phylogenetic studies (15-17). A search of the EMBL/GenBank database using the equivalent fragment of the Israeli outbreak isolates indicated that isolate WN-0043 and WN-0247 were identical to WN-flamingo-NY99, while isolates WN-0233 and WN-0304 were most closely related to the WN-Romania-97 isolate AF130362 (3-nt difference, 1.2%) and less closely related to the WN-flamingo-Y99 (9-nt difference, 3.5%).

A similar search, using a 1,648-nt fragment encoding the preM, M, and part of the E gene, allowed the construction of a more detailed phylogenetic comparison (Figure). As with the 255-nt fragment, WN-0043 and WN-0247 were closest to WN-flamingo-NY99 (AF196835, 99.7% homology), while WN-0233 and WN-0304 most closely resembled a 1997 isolate from Romania (AF130362, 98.4% homology). Phylogenetic analysis showed that two lineages of WN virus circulate in Israel. The first is similar to the WN virus isolates from mosquito, horse, and flamingo during the 1999 NY outbreak. The other lineage is similar to the virus isolated from a mosquito pool during the 1997 Romanian outbreak and to the nucleotide sequences reported from the Russian outbreak in

Table. Analysis of West Nile patients’ isolates, Israel, 2000 outbreak

| Patients | Virus isolate | RT-PCR culture & tissue | Tissue culture | RT-PCR culture & tissue | Tissue culture | RT-PCR culture & tissue |
|----------|--------------|-------------------------|---------------|-------------------------|---------------|-------------------------|
|          |              | acute-phase serum       | (+/IFA)       | infected cell culture   | convalescent-phase serum | (+)          |
| 1        | WN-0043      | (-)                     | (+)           | (+)                     | ND            |
| 2        | WN-0233      | (-)                     | (+)           | (+)                     | ND            |
| 3        | WN-0247      | (+)                     | (+)           | (+)                     | ND            |
| 4        | WN-0304      | (+)                     | (+)           | (-)                     | (-)           |

+, positive result; -, negative result; ND, not done; RT-PCR, reverse transcription-polymerase chain reaction; CPE, cytopathic effect; IFA, indirect immunofluorescence assay.

Figure. Phylogenetic comparison of human West Nile virus isolates from the Israel 2000 outbreak with sequences from the EMBL/GenBank database. The PHYLIP DNA Maximum Likelihood program (bootstrap = 100) was used to compare a 1,648-nt sequence encoding the PreM, M gene, and part of the E gene from the four human outbreak isolates with nine sequences from the EMBL/GenBank database (accession numbers in parentheses) and one from a 1999 isolate from an Israeli White-eyed Gull. CSF = cerebrospinal fluid sample.
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Conclusions

Our sequence analysis shows that at least two lineages of WN virus infected the human population in Israel in 2000. Virus lineage and severity of symptoms were not clearly correlated, although more human isolates would be necessary to confirm this finding. More than one lineage can be found in areas where a virus is endemic and has been circulating for extended periods. More studies, using archived materials, are necessary to determine if there were more than two co-circulating lineages. Yet to be determined is whether changes in the virus genome resulted in a more virulent strain, which caused the high rates of illness and death during the 2000 Israeli outbreak.

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