Genomic Characterization of Nipah Virus, West Bengal, India

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An intrafamilial outbreak in West Bengal, India, involving 5 deaths and person-to-person transmission was attributed to Nipah virus. Full-genome sequence of Nipah virus (18,252 nt) amplified from lung tissue showed 99.2% nt and 99.8% aa identity with the Bangladesh-2004 isolate, suggesting a common source of the virus.

Nipah virus (NiV) causes encephalitis or respiratory signs and symptoms in humans, with high death rates (1–4). NiV outbreaks have been reported from Malaysia, Bangladesh, Singapore, and India (1,5–13). Cases in humans have been attributed to zoonotic transmission from pigs and bats (1,4). We describe a full genome sequence of NiV from an outbreak in India.

The Study

During April 9–28, 2007, five persons became ill and died within a few days at village Belechuapara, Nadia district, West Bengal, India, which borders Bangladesh. The index case-patient (case-patient 1) was a 35-year-old male farmer addicted to country liquor derived from palm juice. Hundreds of bats were observed hanging from the trees around his residence, which strongly suggested association with the infection of the index case-patient and possibility of contamination of the liquor with bat excreta or secretions.

Three patients were close relatives of case-patient 1: his 25-year-old brother (case-patient 2), his 30-year-old wife (case-patient 3), and his 39-year-old brother-in-law (case-patient 4). They became ill 12, 14, and 14 days, respectively, after contact with case-patient 1. In another person, a 28-year-old man (case-patient 5) who collected blood samples from and performed a computed tomography scan of the brain of case-patient 1, symptoms developed 12 days after contact.

No samples were available from the first 2 case-patients. Brain and lung tissues from case-patient 3, cerebrospinal fluid (CSF) from case-patient 4, and urine and CSF from case-patient 5 were collected. Blood samples were obtained from case-patients 4 and 5 and from 34 asymptomatic contacts from the village. Serum samples from these persons were tested for immunoglobulin (Ig) M and IgG antibodies to NiV (IgM/IgG anti-NiV) with ELISA by using reagents provided by the Centers for Disease Control and Prevention (Atlanta, GA, USA). To detect NiV RNA, urine (250 μL), CSF (100 μL), or autopsied brain or lung tissue (100 mg) were used, and RNA was extracted by using TRIzol LS and TRIzol reagents (Invitrogen Life Technologies, Carlsbad, CA, USA). Nested reverse transcription–PCR (RT-PCR) was conducted by using nucleocapsid (N) gene–based primers (8).

Attempts to isolate NiV in Vero E6 cell lines or infant mice were unsuccessful. The full-length genomic sequence was obtained from the lung of case-patient 3 by using 36 sets of primers (online Appendix Table, www.cdc.gov/EID/content/17/5/907-appT.htm), Superscript II RNase reverse transcriptase for reverse transcription (Invitrogen), and Pfx polymerase for amplification (Invitrogen). The PCR products of predicted molecular size were gel eluted (QIAquick PCR Purification Kit; QIAGEN, Hilden, Germany) and sequenced by using BigDye Terminator cycle sequencing Ready Reaction Kit (Applied Biosystems, Foster City, CA, USA) and an automatic Sequencer (ABI Prism 3100 Genetic Analyzer; Applied Biosystems). PCR products were sequenced in both directions. To determine the genotypic status, phylogenetic analysis was conducted by using partial N gene and full NiV genome sequences with the Kimura 2-parameter distance model and neighbor-joining method available in MEGA version 3.1 software (www.megasoftware.net). The reliability of phylogenetic groupings was evaluated by the bootstrap test with 1,000 bootstrap replications.

Patients’ signs and symptoms included high fever (103°F–105°F [39.4°C–49.6°C]) with and without chills, severe occipital headache, nausea, vomiting, respiratory distress, pain in calf muscles, slurred speech, twitching of facial muscles, altered sensorium, (focal) convulsions, unconsciousness, coma, and death. The first 3 case-patients died within 2–3 days after symptom onset; case-patients 4 and 5 died after 5 and 6 days, respectively. Clinical investigations could be conducted for case-patients 4 and 5. Results of serologic tests for malaria parasite, typhoid, anti-dengue IgM, HIV, and hepatitis B surface antigen were negative. Peripheral blood profiles were within...
reference limits. Alanine aminotransferase, aspartate aminotransferase, creatine phosphokinase, and C-reactive protein were elevated. Blood gas analysis in case-patient 4 (case-patient 5 values in parentheses) showed oxygen saturation 49.4% (71%), pO₂ 36.1 mm Hg (44.8 mm Hg), pCO₂ 44.4 mm Hg (44.1 mm Hg), HCO₃⁻ 15.7 mmol/L (19.1 mmol/L), and pH 7.166 (7.255). Lumbar puncture of case-patient 5 showed opening pressure within reference range (1 drop/second), 2 cells/cm; all cells observed were lymphocytes. Chest radiograph indicated pulmonary edema, which suggested acute respiratory distress syndrome. At the time of admission, computed tomography and magnetic resonance imaging scans of the brain showed no abnormality.

Serum samples from case-patients 4 and 5 were positive for IgM anti-NiV. Of the clinical samples screened, brain and lung tissues of case-patient 3, CSF of case-patient 4, and urine of case-patient 5 were NiV RNA positive. Of the 34 asymptomatic contacts, 1 was positive for IgG anti-NiV and negative for IgM anti-NiV and did not report any major illness in the past. This positivity may reflect a previous subclinical infection or cross-reactivity in ELISA needing further follow-up.

Before this report, similar cases had not been reported from the village or the surrounding area. Partial N gene sequences confirmed NiV in the clinical specimens from all 3 case-patients. Phylogenetic analysis showed that similar to findings from the 2001 outbreak study (8), viruses from Bangladesh and India clustered and diverged from the viruses from Malaysia. (Figure, panel A). The length of the full genome of the isolate from India was 18,252 nt. The sequence of this virus (INDNipah-07–1, GenBank accession no. FJ513078) was closer to the virus from Bangladesh (Figure, panel B), with 99.2% (151 nt substitutions) and 99.80% (17 aa substitutions) identity at nucleotide and amino acid levels respectively. Of the 151 nt substitutions, 9 occurred in the N open reading frame (ORF), 11 in the phosphoprotein ORF, 8 in the matrix ORF, 11 in the fusion glycoprotein ORF, 7 in the attachment protein ORF, and 47 in the large polymerase ORF. Fifty-eight substitutions occurred in nontranslated regions at the beginning and the end of each ORF. The intergenic sequences between gene boundaries were highly conserved in the isolate from India, compared with the isolate from Bangladesh, which showed 1 change (GAA to UAA) between the attachment protein and large polymerase genes. No change was observed in the leader and the trailer sequences.

The Table compares amino acid substitutions in the different regions of the genome of the isolate from India with those of the viruses from Bangladesh and Malaysia. Of the 17 aa substitutions, 7 were unique to the isolate from India, and 10 were similar to the isolates from Malaysia. Overall, however, the isolate from India was closer to the isolate from Bangladesh, although distinct differences were observed.

To our knowledge, this is the second report of an NiV outbreak in India, identified within 1 week of the investigation. The first outbreak affected mainly hospital staff or persons visiting hospitalized patients; the 74% case-fatality rate strongly suggested person-to-person transmission (8). Both outbreaks (2001 and 2007) occurred in the state of West Bengal bordering Bangladesh wherein several outbreaks of the disease have been reported (7,9–13). However, fruit bats from West Bengal have not been screened for evidence of NiV infection. This state needs to create awareness about NiV and obligatory testing of suspected case-patients.

Conclusions

NiV caused an intrafamilial outbreak with a 100% case-fatality rate, which confirmed person-to-person transmission.
transmission. The NiV strains from India and Bangladesh were closer than the Malaysian viruses. Although the outbreaks occurred in neighboring geographic areas, NiV outbreaks in Bangladesh and India were not caused by the same virus strain or by spillover.

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Table. Regionwise amino acid substitutions in the Nipah virus genome*

| Region and amino acid position | India | Bangladesh | Malaysia |
|-------------------------------|-------|------------|----------|
| Phosphoprotein 228            | K     | R          | R        |
| 276                           | S     | G          | G        |
| 285                           | R     | H          | R        |
| 310                           | R     | G          | G        |
| Nucleocapsid protein 188      | E     | D          | E        |
| 211                           | R     | Q          | Q        |
| Matrix protein 13              | I     | M          | M        |
| Fusion protein 19              | I     | M          | M        |
| 207                           | L     | S          | L        |
| 252                           | D     | G          | D        |
| Attachment protein 304        | V     | I          | I        |
| Large polymerase protein 94   | I     | T          | I        |
| 112                           | K     | R          | K        |
| 632                           | N     | S          | N        |
| 639                           | N     | D          | N        |
| 665                           | T     | I          | T        |
| 1748                          | I     | V          | I        |

*GenBank accession numbers of isolates examined: India (FJ513078), Bangladesh (AY988601), and Malaysia (AY029767,AY029768, and AJ564623). Boldface indicates unique amino acids in the isolate from India.