Re-emergence of old infection nipah virus

Abstract

Since the last 3 decades new infection including the re-emergence of old infections have become a public health concern as they tend to become pandemic with serious implication for the socio-economic and socio-political scenario of the global community. The facilitating factors include urbanization and destruction of natural habitats, climate change and changing ecosystems, changes in population of reservoir hosts or intermediate insect vectors and microbial genetic mutation, international trade and commerce, change in human demographics and behavior, lack of public health services and infrastructure, and antibiotic resistance.

Keywords: Henipavirus, nipah virus, environment, host

Introduction

According to WHO Re-emerging infectious disease “Infectious agents that have been known for some time, had fallen to such low levels that they were no longer considered public health problems & are now showing upward trends in incidence or prevalence worldwide or have appeared in areas where they were not previously found”.

Factors contributing to emergence

Agent: (1) Evolution of pathogenic infectious agents,(microbial adaptation & change), Mutations. (2) Development of resistance to drugs. (3) Resistance of vectors to pesticides.

Host: (1) Human demographic change (inhabiting new areas), increase contact with animals and natural environment. (2) Human behaviour (sexual & drug use- sharing needles, drug abuse, body piercing),Human susceptibility to infection,(Immunosuppression)- stress and lifestyle changesNutritional changes, more use of pesticides.

Environment: Climate & changing ecosystems, Economic development & Land use,(urbanization, deforestation) Technology & industry (food processing & handling), Changes in agricultural & food production, patterns- food-borne infectious agents (E. coli) (Figure 1).

Etiology:

Nipah virus was first identified in Malaysia in the year 1978 and considered to be a variant of Japanese Encephalitis it is so called after the village of “SungaiNipah” where it was first identified. Nipah is an envelope, negative sense, single-stranded RNA virus, with a genome sequence size of about 18,000 nucleotides. NiV genome organization comprises six major genes present in all Paramyxovirus. Fruit bats (Macrophirotida) of the family Pteropodidae particularly species belonging to the Pteropus genus- are the natural hosts for Nipah virus.

Epidemiology

It presents the variable picture. In Singapore it was associated with importing and slaughter of imported pigs from Malaysia were as in Malaysia the diseases progressed from the natural host “Fruit Bats” to amplification host (livestock) and finally to humanswhere as in Bangladesh it was due to the consumption of fruits and fruits products (raw date palm sap) contaminated with urine or saliva from infected fruit bat. In India, some cases human to human transmission has been reported. In most Nipah outbreaks in South Asia, the virus has displayed a “stuttering chain of transmission”, meaning that once the virus spreads from fruit bats to humans, it moves mainly to people in close contact with patients, such as family, hospital staff and caregivers. In this picture, nursing staff of Kozhikode medical college wear masks.

Clinical features

The Nipah virus infection causes influence like symptoms of high fever, headache, myalgia, sore throat, weakness along with nausea and vomiting. The infection has incubation period of around 2 weeks to 4 weeks. Occasional case have been reported of having a typical pneumonia and acute respiratory distress syndrome. Autopsy finding are predominantly involving the CNS (Figure 2).

The important features of Nipah virus genome include

I. Viral proteins: fusion protein (F) and attachment glycoprotein protein (G)

II. Phosphoprotein (P): it role as a polymerase cofactor, enhancing polymerase processivity and allowing the encapsidation of the newly synthesized viral genomes and antigens.
III. Phosphorotein of Nipah virus has an additional role in immunosuppression: blocking interferon signaling by binding host STAT-1.

IV. Nipah virus (NiV) is a RNA virus belongs to family Paramyxoviridae and genus Henipavirus. Size: 40-600nm, Shape: pleuromorphic, Envelope: present

Figure 2 Genome of Nipah Virus.

Diagnosis: RT-PCR is the main mode of diagnosis serum Neutralisation (SN) tests can also be done.

Treatment: Symptomatic treatment is mainly done as currently no antiviral drugs or vaccines available to treat Nipah virus infection for either people or animals

Recent events in India

The outbreak of Nipah virus in Kerala presents to us several lessons in infectious diseases. Importantly the detection of disease depends on old fashioned clinical skills. The three patients from the same family were admitted to the baby memorial hospital in Calicut. The family had recently changed residence to a new house where there was an abandoned well. Hoping to clear the well for use, two boys of the house descended but were scared away by a bat colony residing there. Subsequently the boys and their father fell ill and were taken to the local hospital in Perambra where they were attended to by nurses of whom three were infected and one nurse died. The patients were evaluated at BMH and right away zoonotic viral encephalitis was suspected. CSF and Blood sampled were sent to the Institute of Virology Alleppey and also to Manipal where Nipah was first suspected and confirmed by the Pune Virology Institute.

a. Prompt and extraordinary response of the local population and local health authorities to meet the challenge. This was prompted by high literacy rate among the local population, better health infrastructure, adequate resources allocation coupled with mass awareness campaign etc launched by the stake authorities helped in “containing” the epidemic at a relatively earlier stage, unlike the once which had happened earlier in North Bengal and Bangladesh.

b. High index of suspicion regarding the possibility of Nipah virus among the local doctors along with the prompt support and correct guidance provided by the Manipal Center of Virus Research enabled the medical fraternity to reach an earlier scientific breakthrough.

c. India has a poor record of outbreak investigations. About 10,000 people develop encephalitis-like symptoms each year but never get a diagnosis. Some regions, such as Uttar Pradesh’s Gorakhpur and Bihar’s Muzaffarpur, saw thousands of deaths in repeated annual outbreaks before the causes were established. Against this background, the discovery of an exotic pathogen in the very second patient hit by an outbreak, as was the case in Kozhikode, has few precedents.

Prevention of emerging infectious diseases: It consists of the following measures, namely, Surveillance and Response, Applied Research, Infrastructure and Training, Prevention and control.13

Conclusion

Whole genome sequencing of Nipah virus along with the establishment of BSL-4 laboratory is essential for managing future epidemics and also providing rapid and effective confirmative diagnosis. Commitment to technology transfer and global collaboration is essential if we are to have the agility required to keep pace with emerging infectious diseases. Pathogen surveillance and discovery can promote global interaction via collaborations on matters that know no national or political boundaries but simply reflect our common goals.14

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None.

Conflicts of interest

There is no conflicts of interest among the authors.

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