A Brief Overview Of Hantavirus Infections

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Abstract
Hantaviruses belong to the Bunyaviridae family as a separate genus. It is currently known that there are over 20 serogenotypes/seroserotypes and that a number of new types are under investigation. The types of Hantaviruses which infect rodents tend to be specific to specific hosts and are primarily found in specific rodent species. As a result of the various types of Hantaviruses and their target organs, Hantaviruses can cause a variety of diseases. There are two major diseases in the world: hemorrhagic fever with renal syndrome (HFRS) and hemorrhagic fever with the syndrome (HPS). A new strain of Hantavirus has been observed in progressively increasing numbers within the world as a result of an accelerating increase in Hantavirus cases throughout the world, which represents an issue of public health of great concern across the globe. Hantavirus disease is a relatively new disease, however, its spectrum continues to expand as the number of recognized virus types continues to rise. It has been reported that Hantavirus causes human disease in the British Isles, but at this point, the disease is largely unknown. Hanta viral infection may be underestimated, especially in developing countries, due to its mild asymptomatic nature and lack of standardized laboratory diagnosis in hospitals. There are no simple standardized laboratory diagnoses in hospitals due to its asymptomatic and non-specific nature. Presented is an overview of what is currently known about hantaviruses and hantavirus infections, including their properties, classification, laboratory diagnostics, treatment, and prevention.

Keywords: hemorrhagic fever, renal syndrome, Hantavirus, rodent

INTRODUCTION:
An enveloped single-stranded negative-sense RNA virus, Hantavirus belongs to the Bunyaviridae family. Rodent excreta are normally inhaled by humans when they are transported between rodents and inhaled by rodents.1 The hantavirus breeds recurrent infections in the rodents of the families Muridae and Cricetidae, which are naturally infected by the virus.2 It causes several diseases in humans, including the Hantavirus (cardio) pulmonary syndrome and hemorrhagic fever with renal syndrome. Each year, there are between 60,000 and 100,000 cases of HFRS reported in China.3

Generally speaking, Seven Hantavirus sero/genotypes have been reported in China, out of which 2 of these viruses are responsible for HFRS, namely the Hantaan virus carried by Apodemusagranarius mice and the Seoul virus transmitted by Rattusnorvegicus rats.4 A serious public health issue continues to plague China with HFRS. The rodents usually become infected with them, but they are not afflicted with any diseases by them. It was first identified in Russian clinical records from far eastern Siberia in 1913 that hemorrhagic fever with renal syndrome (HFRS) was a human viral disease. An account of a similar disease written by Ho Wang Lee (1982) dates back to the year 960. Apodemusagranarius, a murid rodent, has been found to get chronic infections with urinary excretion caused by HFRS as part of its unique renal complications.5

There are many areas in Central and Northern Asia where this mouse is one of the more commonly found wild rodents, and it can sometimes be found in cultivated fields, gardens, haystacks, even in humans' homes. There has been considerable work done to understand how social conflict is associated with HFRS and how it evolved clinically, epidemiologically, and ecologically in recent decades.6 The Hantavirus that has now been found in western Europe and Scandinavia may have caused both Allied and German field nephritis in Flanders during World War I (Bradford 1916; Arnold 1944). As a result of their invasion of Manchuria (Kitano 1944), Japanese military doctors discovered the disease in the mid-1930s, and during World War II, Finnish and German troops contracted the disease (Stuhlfauth 1943; Hortung 1944), and during the Korean Conflicts in 1951 (Smael 1953), UN forces discovered the original and actual hantavirus.7

A LIST OF HANTAVIRUS PROPERTIES IS AS FOLLOWS:
As a structure:
These are negative-sense trisegmented RNA viruses with enveloped structures. RNA-dependent RNA polymerase (L), medium (M) and small (S) segments encode for glycoprotein precursor (GPC) and nucleocapsid (N) proteins, which are respectively processed into the envelope glycoprotein (GnandGc) and the nucleocapsid (N) protein. Protein N of the Hantavirus:
It is the Hantavirus N protein that forms the largest proportion of viral proteins in infected cells and virions. The immune system responds strongly and rapidly to this protein. N proteins contain 100 amino acids at the amino-terminus, making them extremely antigenic. Proteins have B-cell epitopes at their N-termini, while T-cell epitopes are randomly distributed throughout the protein. A given Hantavirus strain of a particular serotype tends to maintain its amino acid sequence in relation to different strains of the same serotype of Hantavirus. Several expression systems can be used to develop recombinant N proteins for commercial use, including Escherichia coli, baculovirus and yeast expression systems. Whenever diagnostic tests are carried out, recombinant proteins are used instead of native proteins.

Glycoprotein of the Hantavirus:
G1 and G2 surface glycoproteins are composed of polyprotein precursors, GPCs, that are degraded by proteases within the cell. In addition, these glycoproteins interact with the surface receptors, β2 integrins and promote entry of hantaviruses.

There are several types of Hantaviruses:
Based on the phylogenetic analysis of their N protein, hantaviruses can be divided into four main groups:
- The following viruses are classified into Group A: Murinae (HFRS-causing species), HTNV, Seoul virus (SEOV), Thailand virus (THAIV), and Dobrava/Belgrade virus (DOBV).
- Group B: Arvicolinae-borne species such as Puumala virus (PUUV), a mild form of HFRS in Europe caused by Nephropathia epidemica (NE).
- Group C: HCPS-causing species in the Americas that are from the Sigmodontinae and Neotominae

The species borne by shrews, moles, and bats are included in Groups D and E.

CLINICAL SYNDROMES ASSOCIATED WITH HANTAVIRUS:
Hemorrhagic fever associated with renal syndrome
Five stages of clinical manifestation were identified: febrile, hypotensive, oliguric, polyuric, and convalescent incubation period of this virus can last anywhere between 2–4 days, and from there, it can manifest itself into symptoms such as aches and pains in the abdomen, headaches, vomiting, nausea, vomiting, headaches, stomach pains, backaches and vision disturbances that can last up to 3–7 days. There may be conjunctival suffusion as well as petechiae on the palate at the end of this process. For the hypotensive (shock) period, the time range can range from a few hours to two days.

Hantavirus cardiopulmonary syndrome:
The severity of this condition is greater than that of HFRS. Fever, chills, malaise, headaches, gastrointestinal discomfort, vomiting, stomach pain and diarrhea are some of the prodromal symptoms. Cardiopulmonary and convalescent symptoms are described as convalescent.

Diagnosis by laboratory tests:
It is important to understand that HFRS and HCPS29S exhibit the same significant laboratory findings. These include: thrombocytopenia, leukocytosis, elevating hematocrit, haematuria, proteinuria, and serum creatinine. The early clinical signs of Hantavirus infections cannot be diagnosed, since these infections are nonspecific. Serology is the principal method of diagnosing Hantavirus infections. It is almost always the case that anti-hantavirus IgM and/or IgG antibodies are present when symptoms first begin to appear.

Transmission and Epidemiology:
Insectivores (Suncus murinus) and rodents from the subfamilies Murinae, Arvicolinae, and Sigmodontinae all tested positive for Hantaviruses. HNTV, DOBV, SAAV, SEOV and Amur viruses are the most common hantaviruses related to HFRS and non-human diseases. Murinae are the most common rodent species of the Old World. As with Prospect Hill virus and other viruses in the United States, there is a correlation between PUUV and HFRS in Europe, but not for human infections.

There are several viruses from the New World that are transmitted through the species of rats and mice called Sigmodontinae, which are widespread throughout the New World. In each subfamily of rodents, there is a phylogenetically distinct virus, some of which may be human pathogens, while others may not be. There is a significant concordance between the phylogenetic relationships between the viruses and those of their dominant host (with rare exceptions), which shows a close association between the two. The majority of hantaviruses under study have their own
murid rodents host, except for TPMV, along with other newly discovered hantaviruses such as AVG, Ripley virus (RPLV), Tangganya virus (TGNV), and Cao Bang virus (CBNV). Hantaviruses, however, are not just found in insectivores like Suncus murinus; they have also been identified in bats, cats and birds, as well as other species of insectivores.\(^{17}\)

The presence of hantavirus infection in dogs and pigs was also demonstrated by serological tests.\(^{18}\) The presence of these species is unclear and it is unclear whether they are persistently infected or if they are just spilling over infected secondary hosts after contacting the primary host.\(^{19}\)

Transmitting:

Animal excreta, such as saliva, urine, and feces, are believed to be the primary transmission source of Hantavirus.\(^{20}\) It is undoubtedly true that rodents and humans, as well as rodent rodents, are most commonly infected by aerosols. However, virus transmission by bite can also occur among rodents.\(^{21}\)

The epidemiological features are as follows:

Rural areas have a greater prevalence of hantavirus infections, but urban areas have an increased prevalence of SEOV-induced HFRS. Hantavirus infections are underestimated due to asymptomatic or nonspecific mild infections.\(^{22}\) Some Hantaviruses can be found in Europe to have a ratio of 14:1 to 20:1 between subclinical and clinical infection. In Finland, HFRS cases averaged 13% per year in an eight-year study (ranging from 4% to 30% depending on the region).\(^{23}\) According to these findings, as many as 70% of all cases of PUUV infection are underdiagnosed because of mild or subclinical symptoms that are associated with the infection.\(^{24}\)

**Fig No. 1: Rodent borne disease**

Management of clinical cases:

HFRS and HPS do not currently have specific treatments; therefore, supportive measures remain the cornerstone of treatment. Intensive care units that can monitor vital signs such as blood and tissue oxygenation, cardiac output, central blood pressure, and cerebral pressure must be immediately accessed. The balance of fluids must be maintained at all times.\(^{25}\)

The dosage of haemodialysis for HFRS patients normally requires one to two sessions, while for HPS patients, mechanical ventilation is considered to be essential when high pressures are indicated and sufficient ventilation is needed.\(^{26}\) As a rescue therapy for severe HPS, extracorporeal membrane oxygenation has proven effective. In severe cases of HFRS and HPS, corticosteroids were used, although they are not standard treatment for hantavirus infection.\(^{27}\)

Diagnostic testing in laboratories:

Laboratory tests and clinical information are used to diagnose hantavirus infections. Hantavirus infection can be difficult to diagnose in individuals with mild to moderate symptoms.\(^{28}\) In addition to native or purified virus preparations, the detection of hantavirus antibodies has also been accomplished using recombinant proteins produced by bacteria, yeast, and insects. Identifying IgM antibodies or low-avidity IgG antibodies usually involves indirect fluorescent analysis (IFA) and enzyme immunoassays (EIA).\(^{29}\) There are several methods for detecting IgM that can be used to detect acute infections, particularly in endemic areas where IgG is highly prevalent due to previous infections, which is highly useful for diagnosis of acute infections.\(^{29}\)
CONCLUSION

There is no specific treatment for hantavirus infection, and the best way to prevent infection is to avoid contact with rodent excreta and to take precautions when cleaning areas where rodents have been present. The hanta virus infected patients receive medical attention in an intensive care unit early and are diagnosed, we know they will do well. To assist patients undergoing extreme respiratory distress, intensive care facilities use incubators and oxygen therapy.

REFERENCES