Ebola virus: A global public health frightening

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Abstract

Ebola virus causes disease, previously known as the hemorrhagic fever, is a menace routinely lethal sickness triggered by infection per the Filoviridae family’s Ebola virus. The Ebola virus (EBOV) is feast by commerce by means of the blood or bodily fluids of someone who has contracted or died from EVD, as well as contaminated things like as needles, animals that was infected, otherwise bush meat. EVD takes two days to three weeks’ incubation period then an rushed inception devoid of the need for a carrier status. A flu-like syndrome is characterized by a rapid onset of high fever, chills, and myalgia, among other clinical signs. It’s generally identified by a combination of clinical symptoms, including start of disease, high fevers lasting fewer than 3 weeks, and as a minimum 2 hemorrhagic symptoms in spite of no predisposing causes. This is usually enough evidence for physicians to suspect EHF and start supportive therapy until test findings confirm the virus. Patients first show with nonspecific influenza similar symptoms before succumbing to shock and several organ failures. Severe bleeding problems and several organ failures are the leading causes of death. For the reason that there is presently not any conventional rehabilitation for EVD, this one is perilous to minimize contagion and the virus’s transmission. Supportive treatment for patients includes maintaining fluid balance, electrolyte balance, hypertension, and oximetry, beside that treatment consequences arise after secondary infections. This review gives a comprehensive summary of the Ebola virus.

Keywords: Ebola virus disease; Ebola hemorrhagic fever; diagnosis; treatment.

INTRODUCTION

A severe infectious outbreak characterized by acute viral hemorrhagic fever first emerged in 1976. Because this outbreak in Yambuku had 318 cases and an 88 % mortality rate (280 deaths), it was called Ebola then subsequently Ebola River, which was discovered wherever the Democratic Republic of Congo (DRC) [1]. Besides the genus Marburg virus, the Ebola virus belonging to family Filoviridae. The envelope of this extremely pathogenic virus has a filamentous or branching convoluted structure and contains a linear RNA genome non-segmented single-stranded negative-sense [2]. This virus comes in five distinct species, each with its own set of biological properties and pathogenicity [2, 3], as shown in figure (1).

The original source of the Ebola virus is unknown to this day, but this is thought to be animal-borne because infected animals transmit the virus directly to other animals, such as monkeys, chimps, and apes, and even humans, leading to Ebola virus distributed throughout the human species via human-to-human disease transmission [4]. Fruit bats, which belong to the Pteropodidae family, seem assumed to be the likely source for Ebola viruses (EBOV), with humans and the rest mammals reflect as occasional accidental hosts [5].

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The outbreak of Ebola originated inside West Africa in a rural place in south-eastern of Guinea between 2014 and 2016. In the interior weeks, this one be obligatory extent to metropolitan areas and across borders, and within months, the aforementioned must come to be an international pandemic [4]. This epidemic was the start of the West African Ebola pandemic, and it was also the biggest outbreak of Ebola virus that recorded [6]. From 2014 to 2016, the West African (EVD) outbreak caused at least 28,616 cases and 11,310 mortalities [7]. In previous Ebola virus epidemics, case treatment has mostly centered proceeding by segregating patients infected with EVD, avoidance infection and governor protocols, besides varying levels of rudimentary loyal upkeep [8]. EVD was managed with increasingly supplementary unconventional supportive cure aimed at patients throughout the West African pandemic [9,10,11]. Mortality was over 74 percent at the start of the outbreak [12], although it later dropped to 31–37 percent [13, 14], possibly owing to better prompt case revealing and supportive dealing. In West Africa, specific medication against Ebola virus be situated merely accessible in rare cases. Patients who were evacuated to European as well as American hospitals and got intensive care plus anti-Ebola virus therapies, on the other hand, had a fatality rate of 18.5% [9, 15,16,17,18,19].

**Clinical manifestations**

EVD is a kind of intense hemorrhagic fever with 2 to 21 days for symptoms to appear (mean 4-10) with that are similar to the respiratory tract infection, such as high temperature, chills, malaise, and muscle aches. The hot condition possibly will be modest for primary infection stages; on the other hand, it can quickly worsen with chills in addition to rigors [20]. Fever with just a skin rash mostly on face, neck, chest, and artilleries, generally developing afterwards 5-7 days of sickness, lethargy, vomiting, diarrhea, as well as anorexia are the most often document symptoms in the existing outbreak in Africa. [21,22].

**Diagnosis**

At the time of clinical manifestation, leukopenia with lymphocytes and a higher proportion of granulocytes typically frequent laboratory results. Leukocytosis can develop as the illness advances, also an upsurge in undeveloped granulocytes. A long-term condition that lasts till death is Thrombocytopenia. Alanine as well as aspartate aminotransferase (ALT, AST) serum levels are commonly found to be somewhat increased [20]. In the early stages of the disease, blood total bilirubin levels are either normal or increased, and jaundice is uncommon. Severe fluid loss through diarrhea and vomiting beside that lack of sufficient water replacement results in elevated serum urea nitrogen and creatinine levels [23].

To confirm infection, enzyme-linked immunosorbent assay (ELISA), as well as molecular test such polymerase chain reaction (PCR), and viral isolation techniques are utilized [20].

**Treatment**

Symptomatic as well as helpful therapy as electrolyte standby, dietary maintenance, preserving oxygen prominence and blood pressure upkeep, and handling superfluous infections, remain the chief measures ordered to infected individuals. In attendance is definitely not authorized EVD vaccination or conventional therapy that has been tested in people [24, 25]. ZMapp is a pharmacological drug for EVD that is still in the early phases of consideration. It includes three "humanized" monoclonal antibodies (mAbs) against the EBOV GP protein. Nicotiana, in particular, is a kind of plant benthamiana, generate these antibodies [26]. This medicine was initially tested in people in an experimental setting for the duration of the 2014 West African EVD pandemic, nonetheless never randomized organized study has been conducted to assess if it works or if it is secure enough to have already commercialized [27, 28].

Inmazeb (REGN-EB3), unlike ZMapp, would be the first FDA-approved therapy for children and adults with EVD [29]. Atoltivimab, maffitivimab, and odesivimab are the three completely monoclonal antibodies (mAbs) for human that make up REGN-EB3. These three mAbs in the REGN-EB3 combination won't be competing for binding to the glycoprotein (GP) of Ebola virus, but instead attach to the GP at separate sites at the same time, resulting in strong viral neutralization [30,31]. The envelope protein GP of the Zaire ebolavirus facilitates virus attachment and membrane fusion with cell membranes of host cell. The antibodies will attack the GP protein expressed on the virus-infected host cell and destroy the virus by antibody-dependent cellular cytotoxicity. [31]. Ebanga would be monoclonal antibody for human that
is used to cure the ebola virus from Zaire. Fever, tachycardia, diarrhea, vomiting, hypotension, tachypnea, and chills are the most prevalent side effects linked with Ebanga, compared to serious side effects related with ZMapp as well as Inmazeb [32].

**Conclusion**

EVD is a highly contagious illness that has caused a number of epidemics, mostly in Africa. The specific origins of the Ebola virus remains unknown to this day, however it is thought to be animal-borne since sick animals transmit the virus directly to other animals. For EVD, there are currently just a few medicines and therapy options. In addition to supportive care, the most regularly utilized medications include ZMapp, Inmazeb, and Ebanga. The mortality rate in patients treated with Inmazeb, is lowered by 17%. Inmazeb is the preferable medication in terms of efficacy. Ebanga is an optimal in patients suffer cardiovascular complications.

**References**

32. US Food and Drug Administration. FDA Approves Treatment for Ebola Virus