Marburg virus disease

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Summary
In the late summer of 1967 an epidemic in thirty-one patients in Germany and Yugoslavia of a disease transmitted from African green monkeys occurred; seven patients died. The incubation period was from 4 to 7 days. The main clinical features were headache, high fever, diarrhoea, a very characteristic rash, severe bleeding tendency and involvement of the central nervous system. Nearly all organs were involved and showed severe cell necroses. The aetiological agent was identified as an RNS-virus and was named Marburg virus. It was detected in the blood, urine, throat-washing and seminal fluid.

In August 1967 a hitherto unknown severe infectious disease with features of haemorrhagic fever was observed simultaneously in Germany and Yugoslavia which could be traced to imported monkeys (Cercopithecus aethiops) from Uganda.

Until then only single cases of diseases transmitted by monkeys were known (e.g. herpes simiae, rabies, hepatitis). The new disease was the first to occur as an epidemic.

Incidence
Altogether thirty-one persons (twenty males, eleven females) contracted the disease: at Marburg twenty-three, at Frankfurt six, and at Belgrade two persons. Out of these thirty-one patients seven died.

The epidemic appeared in August and ended in November 1967. Before and afterwards no more cases were seen.

Clinical picture (Figs. 1 and 2)
The incubation period could be determined in some patients who had only once been exposed to the infectious material; it varied from 3 to 9 days.

The prodromal phase was very short. The disease began rather abruptly with extreme malaise, pains in the limbs, and severe frontal and temporal headache.

Within a few hours the fever increased to 39°C without rigors and reached a maximum on the 3rd and 4th days; the temperature fell gradually afterwards with a second peak between the 12th and 20th days.

Relative bradycardia occurred, especially in the early days, whereas tachycardia was only found in the fatal cases. Many patients complained from the beginning about nausea and suffered from frequent, and occasionally uncontrollable, vomiting. Watery diarrhoea with blood or mucus occurred up to ten times daily with symptoms and signs of severe dehydration and acute renal failure.

All patients developed a characteristic maculopapular rash on the 5th to 8th days. It began on the face and on the buttocks, then progressed to the trunk and extremities. Most characteristic were dark red pinhead papules round the hair follicles. After 1 or 2 days it developed into a sharply delineated maculopapular lesion which coalesced into a more diffuse and dark red erythema. Cutaneous purpura was rare. The rash was in many patients accompanied by scrotal dermatitis or erythema of the greater labia. About the end of the second week all patients peeled, especially on the palms and soles.

The external rash was in most cases accompanied by an enanthem with dark red colouring of the soft palate and tapioca-like transparent lesions.

A conjunctivitis and photophobia developed in half of the patients. Swelling of the lymph nodes was observed in the nuchal, cervical, and axillary regions between the 3rd and 6th days of illness. The lymph nodes were soft and sensitive to pressure. The spleen was palpable in only one patient, the liver was tender but not enlarged. The central nervous system was affected in different ways. Nearly all patients showed a sullen, slightly aggressive, and negativistic behaviour. Those who died became restless, confused, unconscious, and died in deep coma. Two had severe convulsions. Some patients complained of paraesthesia and restless legs. One patient developed a severe post-infectious myelitis.
with a Landry-type of paralysis with albumino-cellular dissociation in the cerebrospinal fluid (Guillain–Barré syndrome). One patient became psychotic several weeks after the acute illness.

About half of the patients exhibited marked haemorrhagic diathesis with spontaneous bleeding from the nose, gingiva and gastro-intestinal tract, and haematuria. The women had genital bleeding. At the sites of injections and needle punctures severe bleeding occurred.

The pancreas was affected in several patients as shown by the increased serum amylase values.
Nearly all patients had signs of severe hepatocellular damage with an extreme increase of the transaminase levels. Neither jaundice nor true hepatic failure developed. In the electrocardiogram alterations compatible with diffuse myocarditis and disturbances of cardiac rhythm were seen. In many patients cardiac failure developed. Nearly all of the patients who died had severe kidney involvement with acute renal failure. Those who survived had longer periods of polyuria. The haemopoietic system was severely damaged in nearly all patients as could be recognized by marked leukopenia and severe thrombocytopenia.

Course of the disease

The illness lasted 15–20 days. In some cases there was a relapse. Five patients showed a further increase of serum transaminase at this time with an increased temperature. The convalescent period was very drawn out. There were disturbances of the autonomic nervous system with exhaustion and increased perspiration. Some of the patients had a considerable loss of hair and complained of tenderness of the liver and alcohol intolerance. Unilateral testicular atrophy was observed in five patients. Some had reduced libido and potency and oligosperma; the ketosteroid levels were normal. Three women were delivered of four healthy babies with no signs of disease.

Laboratory findings

The erythrocyte sedimentation rate rarely exceeded the upper normal range. The haematologic changes apart from leukopenia and thrombocytopenia were characterised by a shift to the left in the granulocytes with up to 40% stab cells, promyelo-, metamyelo- and myelocytes. This shift to the left was accompanied by the appearance of so-called degenerative granulocytes and Pelger-like cells. There was a marked increase of atypical lymphocytes, plasma cells, immunoblasts and so-called pyroninophil blast cells, up to 15% of the total count. Bone marrow puncture showed an elevated number of immature megakaryocytes. All patients developed severe thrombocytopenia with sometimes less than 10,000/mm³. At the same time the thrombin time was prolonged, indicating consumptive coagulopathy. In none of ten patients in whom the prothrombin time, partial thromboplastin time and fibrinogen were measured did the alterations and plasma coagulation factors explain the severe haemorrhagic diathesis.

The extreme increase of serum transaminases and other liver enzymes reflected the severe hepatocellular damage. In four of the five patients who died, the transaminases increased up to 3000–6000 U/l. The relation between SGOT and SGPT was 7 : 1. The maximum increase was between the 7th and 8th days, with a second peak around the 35th day in three patients. The bilirubin was only slightly increased in some patients with anuria. The creatinine and urea were elevated in those patients who died with renal failure. The total serum protein decreased markedly in several patients.

The other complications included bronchopneumonia (five cases), leg oedema (five cases), orchitis (three cases), post-infectious myelitis (one case), and psychosis (one case).

Pathology

In nearly all organs areas of focal necrosis without much inflammatory reaction were seen. They were particularly obvious in the liver, in the lymphatic system, in the spleen, in the testicles and ovaries, and in the pancreas. In addition the lymphatic tissue showed plasmacellular, monocytoidal transformation. Together with the areas of necrosis so-called basophilic bodies were noticed near the necrotic cells or as inclusion bodies in parenchymal cells. Those patients who died showed glial nodules, encephalitis in all areas of the brain and serious parenchymal damage of the kidneys with definite signs of tubular insufficiency and haemorrhagic diathesis in all organs.

Diagnosis

Laboratory methods included the direct isolation of the agent in the blood, organ tissues, throat washings, urine and semen (Fig. 3). Virus antigen was made visible by means of immuno-fluorescence. Direct electron microscopic demonstration of the virus was possible after centrifugation of serum on to carrier films.

The morphology of the Marburg virus is most unusual. It differs in size and shape from most known...
Marburg virus disease

![Graph showing onset of Marburg virus disease in thirty-one patients.](Image)

**Fig. 3.** Onset of Marburg virus disease in thirty-one patients. (Courtesy of Professor Siegert, Marburg, and Springer Verlag, Heidelberg.)

Viruses. The mean length is about 665 nm. It shares certain similarities with other viruses (plant viruses, Rabies, Egtrle virus of rainbow trout).

A complement fixing antibody test using a cell culture antigen from infected Vero carrier cells is highly specific.

**Epidemiology**

There were four groups of infection (Fig. 4). The first and largest group consisted of persons who had direct contact with blood and organs of monkeys imported from Uganda. The second group consisted of laboratory workers who had contact with simian organs or cell culture material. The third group contracted the infection from other patients’ blood.

Exposure in the household and family surroundings remained without consequences. In one woman a venereal way of transmission was the most probable, since her husband was found to excrete the Marburg virus in his semen.

The simultaneous outbreak of the disease in Marburg, Frankfurt and Belgrade was traced to common imported monkeys of the species *Cercopithecus aethiops* from Uganda. These monkeys were flown because of the Near East crisis, to Germany and Yugoslavia via London, where they were kept at the airport together with other animals. Seroepidemiologic studies using a complement-fixing antibody test seemed to suggest a high percentage of positive reactions in monkeys from Uganda and from other areas. These results could not be confirmed by other authors using a more specific method. The reservoir and method of distribution of the Marburg virus are still unknown. Intrathoracic multiplication of the virus was possible in *Aedes aegypti*.

**Treatment**

The therapy was symptomatic. Antibiotics were used prophylactically against secondary bacterial infections. Acute renal failure was treated according to standard rules including peritoneal dialysis. Low sodium human albumin was infused in patients with hypoproteinaemia, and electrolytes and fluid supplemented. The most difficult problem was the treatment of the haemorrhagic diathesis. Fresh blood thrombocyte concentrates, fibrinogen, e-amino caproic acid, vitamin K and the French preparation PPSB (Prothrombin, Proconvertin, Stuart-factor and antihaemophilicglobin B) were given. The latter proved to be the most efficient.
Course and prognosis

The disease varied greatly in severity. There were some mild cases but most of the patients were critically ill. The fatality rate was 22%. Complete recovery was the rule, but most patients even now after 5 years complain about easy fatigability, inability to concentrate, loss of libido, etc. It is very difficult to decide what complaints have a real background or which symptoms are due to the unique situation, as all infections were acknowledged as occupational hazards and the pension scheme makes it necessary that follow-up studies are required every year. So a kind of post-Marburg virus disease neurosis might play a role.

References


