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**Haemolytic-uraemic syndrome with thrombotic microangiopathy of the retina following cytomegalovirus infection: postmortem findings**

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**Summary:** Haemolytic–uraemic syndrome is characterized by thrombotic microangiopathy of the glomeruli and smaller arterial vessels of the kidney. Extrarenal thrombotic microangiopathy occurs, but ocular involvement is rarely demonstrated microscopically. We describe a 33 year old woman with a 3 week febrile episode and seropositivity for cytomegalovirus infection who developed acute renal failure, blindness and severe encephalopathy which was the cause of death. Thrombotic microangiopathy of retinal vessels may be more common than is expected from the literature. The exclusion of ocular structures in postmortem examinations explains the lack of anatomoclinical correlation.

**Introduction**

The haemolytic–uraemic syndrome is a clinical entity, defined by microangiopathic haemolytic anaemia, thrombocytopenia and acute renal failure, whose underlying histopathological feature, thrombotic microangiopathy, is also found in thrombotic thrombocytopenic purpura. Haemolytic–uraemic syndrome and thrombotic thrombocytopenic purpura are considered by some authors as a different clinical expression of the same disease. Thrombotic microangiopathy always involves the glomeruli and smaller arterial vessels of the kidney, but sometimes escapes the extrarenal vascular beds. Ocular structures affected by the thrombotic microangiopathy of haemolytic–uraemic syndrome have been occasionally described. The present case appears to be the first for haemolytic–uraemic syndrome following cytomegalovirus infection with documented retinal histology at autopsy.

**Case report**

A 33 year old women presented with a 3 week history of fever. The patient’s medical history was unremarkable. On the following days after admission to the hospital, the patient developed peri-orbital oedema, hypoproteinaemia with no evidence of proteinuria and decreased visual acuity which was attributed to an acute exudative chorioretinitis with peripapillary retinal oedema. Blood pressure was normal. Treatment with corticosteroids was unsuccessful and acute renal failure developed. The serum creatinine level was 707 μmol/l (normal <123 μmol/l). Laboratory tests disclosed microangiopathic haemolytic anaemia (haemoglobin 6.5 g/dl) reticulocytes 5.8%, platelet count 106 × 10^9/l, schistocytes + + + +, haptoglobin <0.25 μmol/l (normal 0.27–1.21 μmol/l) and
elevated IgM-specific cytomegalovirus antibody. Cultures of the samples were not done. Lupus anticoagulant was not performed but blood-clotting tests including activated partial thromboplastin time were within normal limits. Serological tests for syphilis (VDRL) and human immunodeficiency virus (HIV) infection were negative. Skin changes attributable to scleroderma were not present and antinuclear antibodies were negative. The patient did not respond to haemodialysis and plasma exchange. The renal biopsy revealed extensive glomerular involvement with over 90% showing thrombotic microangiopathy. Focal IgM, C3 and fibrin deposits were also observed. Some small arteries were completely occluded by thrombosis. Refractory renal failure was followed by severe encephalopathy, generalized seizures and blindness. The patient died of hypoxic–ischaemic encephalopathy a week following the onset of alteration of consciousness. A diagnosis of haemolytic–uraemic syndrome and cytomegalovirus infection was made.

Postmortem examination. Microscopically, glomerular thrombotic microangiopathic features were identified. Apart from the kidneys, intravascular thrombosis was seen in the small feeding arteries emerging from the ganglion cell layer of the retina (Figure 1). Signs of ischaemic necrosis were evident in most ganglion cells (Figure 2). Neither inflammatory infiltration nor nuclear inclusions of cytomegalovirus infection were seen in any of the multiple sections examined by light microscopy. *In situ* hybridization with specific probes for cytomegalovirus (Enzo Biochem) was negative. Cerebral oedema and severe diffuse hypoxic/ischaemic changes were widespread in the central nervous system.

**Figure 1** Thrombosis of a small arterial vessel (arrow) in the ganglion cell layer of the retina (longitudinal section) (haematoxylin and eosin × 25).

**Figure 2** Arterial vessel completely occluded by thrombosis (arrow) (transversal section). Note degenerative changes in ganglion cells (haematoxylin and eosin × 25).

**Discussion**

The haemolytic–uraemic syndrome is a rare condition characteristically occurring in children over 4 years of age. In an autopsy series, Zollinger and Mihatsch\(^2\) estimated an incidence rate of 1/25,000. A sex predominance has not been reported.

Different agents have been implicated in the pathogenesis of haemolytic–uraemic syndrome, particularly infection with verocytotoxin-producing *Escherichia coli* which has been demonstrated in 75% of children with this disease. Shigella and Campylobacter-associated haemolytic–uraemic syndrome can also occur.\(^3,9,10\) Viral infections (mumps, chicken-pox, arbovirus, HIV)\(^10,11\) use of oral contraceptives, pregnancy, immune dysfunction, antineoplastic therapy, lupus erythematosus, scleroderma, malignant hypertension and radiation therapy are some of the possible associated conditions with adult forms of haemolytic–uraemic syndrome.\(^1,10–12\)

The patient in our study presented with a 3-week febrile episode and seropositivity for cytomegalovirus infection. To our knowledge, a post-infection form of haemolytic–uraemic syndrome to cytomegalovirus has not been previously described. There is speculation that vascular endothelial cell injury from viral infections, bacterial toxins or immune complexes is the primary event.\(^3,4,13\) This endothelial cell injury might then release proaggregatory substances and initiate the formation of platelet–fibrin microthrombi.\(^4\) It is not clear why thrombotic microangiopathy always involves the renal, but usually spares the extra-renal, vascular beds.

Although several authors have emphasized the importance of neurological symptoms in haemolytic–uraemic syndrome,\(^9,14,15\) the involvement of the central nervous system in this disorder is less frequent and of lesser intensity than that in throm-
microangiopathy involving patients, sight syndrome. Visual abnormalities include vitreous haemorrhage, retinal detachment, optic nerve atrophy and exudates and inflammatory retinal thrombi found in the central nervous system dysfunction. In the case reported here, platelet–fibrin microthrombi found in the small retinal vessels were the cause of necrosis of the ganglion cells and blindness. We have recently documented another case of haemolytic–uraemic syndrome with loss of visual acuity in a 16-year-old boy with acute monocytic leukaemia as a complication of chemotherapy and irradiation. As fundoscopic examinations were not carried out, the loss of visual acuity was attributed to chorioretinitis due to Toxoplasma as this organism was shown to be the cause of encephalitis in the postmortem examination. Both cases illustrate that the optic nerve and retina need to be included in the list of possible extraregional structures affected by thrombotic microangiopathy in haemolytic–uraemic syndrome.

Acknowledgement

The authors thank Marta Pulido, M.D., for editorial assistance and copy editing.

References