Excess of Pappenheimer bodies (siderocytes) has been found in two splenectomized siblings with congenital dyserythropoietic anemia - type II (CDA-II) and iron overload.

Pappenheimer bodies were first time described in 1945 by Alwin M. Pappenheimer in three patients after splenectomy because of hemolytic anemia of undetermined cause (12). Pappenheimer stressed that these bodies appeared only after splenectomy although morphological characteristics suggested a relationship to certain intraerythrocytic parasites. The author has not obtained any conclusive proof of their parasitic nature. The inclusions gave positive Perl’s reaction. He considered the possibility that these iron containing bodies might be identical with the iron containing bodies described in 1941 by Grüneberg (3). According to our knowledge it is the first description of Pappenheimer bodies in CDA-II.

Patient Reports

In two siblings at the age of 20 (K. J.) and 5 years (K. K.) diagnosis of CDA-II has been made 30 years ago (7).

The diagnosis of CDA-II was established:

- By the demonstration of erythroid hyperplasia, karyorrhexis and erythroblastic bi- and multinuclearity in the bone marrow.
- By positive results of the acidified serum tests but negative results with patient’s own serum.
- By demonstration of typical ultrastructural features of erythroblasts and some erythrocytes showing the “double membrane phenomenon”.

The study of survival of erythrocytes with $^{51}$Cr revieled shortened survival with destruction in the spleen. The liver biopsy performed during operation demonstrated iron overload. Both siblings were splenectomized at the age of 23 (K. J.) and 15 years (K. K.) respectively to diminish the iron overload due to hemolysis in addition to the ineffective erythropoiesis (7). Iron overload is a frequent complication in CDA-II (4, 6). Splenectomy led to moderate increase in hemoglobin, increase or nearly normalization of the red cell survival but did not prevent further iron loading. In 2004 the serum ferritin reached in K. J. 1450,4 µg/l and in K. K. 1131,7 µg/l (normal value 30–350 µg/l), liver biopsy revieled excessive iron overload in both patients with che...

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Fig. 1: Pappenheimer bodies in patient K. J. In one erythrocyte Pappenheimer body and Howell-Jolly body are present.
mically determined liver iron concentration 14843 µg/g and 15415 µg/g respectively (normal value 70–1400 µg/g).

The blood films reported at that time in K. J. and K. K. have shown 46.4 % and 15.9 % of Pappenheimer bodies respectively (Fig. 1). The iron stain confirmed that the Pappenheimer bodies were siderocytes.

Discussion

Pappenheimer bodies are siderosomes (iron-containing granules with positive Perl’s stain). Electron microscopy of these bodies shows that the iron is often contained within a lysosome as confirmed by the presence of acid phosphate. Siderosomes may also contain degenerating mitochondria, ribosomes, and other cellular remnants. Using iron stain we observed that siderocytes bodies were formed by 1 to 3, occasionally even by more particles. Basophilic stippling represents dispersed blue granulation. Howell-Jolly bodies are spherical in shape, usually not larger than 0.5 µm in diameter and give a positive Feulgen reaction for DNA.

Pappenheimer bodies can be mistaken for Babesia (2). Human babesiosis is an acute febrile illness sometimes complicated by hemolytic anemia common in North America. The illness is caused by intraerythrocytic protozoal parasites of the genus Babesia. Babesia organism measures 0.9–2 µm, Pappenheimer bodies are up to 2 µm in size.

Spuriously elevated platelet counts counted by an electro-optical counter due to Pappenheimer bodies have been reported too (11,13).

Kent has shown that siderocytes are not present in normal erythropoiesis with intact spleen. In hematologic disturbances with reticulocytosis and intact spleen the observed counts were 0.10–1.11 %, in splenectomized persons because of traumatic rupture of the spleen 0.20–0.40 %. In splenectomized patients with persistent hematologic disturbance the counts varied from 0.9 % to 57 % in a case of hemoglobin C disease with splenectomy (9).

Pappenheimer bodies were also observed in congenital sideroblastic anemias (5,14). Iwama et al. (8) reported a case of macrocytic anemia in a 75-year-old alcohol abuser who suffered vitamin B6 deficiency. Abstention of alcohol led to the disappearance of Pappenheimer bodies commonly observed in the red blood cells of drinkers. Mende and Fülle (10) mentioned a 36-year-old alcohol abuser with hemolytic anemia (Zieve-syndrom) with 25 % of Pappenheimer bodies in admission with their complete disappearance after 20 days of abstention of alcohol.

In our two patients the presence of Pappenheimer bodies was due to the CDA-II and the splenectomy. How much the iron overload contributed will be possible to be assessed after normalization of serum ferritin levels.

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