CHRONIC HYPOPLASTIC ANAEMIA ARISING IN INFANCY

BY


(From the Royal Victoria and West Hants Hospital)

An aplasia of the erythroblastic tissue in the bone marrow, without involvement of either leucocytes or platelets, is referred to by Whitby and Britton (1946) as a medical curiosity. This aplasia may be complete or partial. If complete, there is no formation of normal red-cell precursors. Strictly speaking, the term 'aplastic anaemia' should be reserved for cases of this kind, where there is a pure red-cell anaemia without involvement of leucocytes or platelets. If the aplasia is partial, then the term 'hypoplastic' is used. 'Aleukia haemorrhagica' (Frank, 1915) would be a better term to use for other children, he was born of a trial labour, and weighed 6 lb. 4 oz. at birth. He had been bottle-fed from birth (National Dried Milk), and there was no history of jaundice or other infection before admission.

Examination showed a very small pale child with a slight lemon tint to the skin and moderate ptosis of the right upper eyelid which had been present since birth (fig. 1). The central nervous system was otherwise normal. No abnormality was detected in the cardiovascular or respiratory systems, and the liver and spleen were not palpable. There was no koilonychia. He was afebrile,

Fig. 1.—The patient.

conditions where red cells, leucocytes, and platelets are all reduced in numbers.

With the more general adoption of bone-marrow biopsy as a routine haematological investigation in cases of severe anaemia, a pure red-cell anaemia may be found to be more common than is thought at present. Such cases have been described in infants, children, and adults.

Case Report

D.S., aged 1½ years, an illegitimate male child, was admitted to the Royal Victoria and West Hants Hospital on Sept. 25, 1946, with a history of having been very pale since birth. The mother had no

weighing 17 lb. 5 oz., and was therefore approximately 5 lb. under weight for his age. Mentally he was backward, being unable to say words of more than one syllable. A blood count on admission showed severe anaemia (table 1) and a marrow puncture of two days later showed evidence of aplasia of the red-cell elements. A blood transfusion was given on Oct. 2 with good effect (fig. 2). From that date onward blood transfusions

Fig. 2.—Showing per cent. haemoglobin. On Jan. 31, 1947, proteolysed liver, drachm 1, three times a day, was begun, and there was an apparent response on Feb. 14, the Hb. per cent. having risen from 48 to 60. This good effect was not maintained, however, and on Apr. 1 the Hb. per cent. was down to 36. The estimate of the Hb. per cent. was unfortunately omitted after blood transfusion of June 27, hence the gap in the chart.
proved the only effective treatment. At different times he was given large doses of iron and crude liver preparations, yeast, ascorbic acid, and folic acid. Iron preparations containing traces of cobalt and manganese were tried. All forms of therapy excepting blood transfusion were without much effect.

Radiographs of the skull, chest, and long bones were normal. The Wassermann reaction was negative. Both mother and child are Rh-positive, and there was no Rh-antibody in the maternal serum. Urine examination was normal. Fat analysis of the stools was within normal limits. A fractional test meal in April, 1947, showed a complete achlorhydria, but when repeated after the subcutaneous injection of histamine it showed 18 ml. N/10 acid per cent. at the end of one hour. His weight in October, 1947, was 23½ lb. He was then still five or six pounds under weight, and his height, 31½ inches, was two or three inches below the average for his age. His mental condition had improved considerably.

Repeated blood counts during the year in which he was observed constantly showed a normocytic normochromic anaemia and very low or absent reticulocytes but normal platelets and leucocytes (table 1). The bleeding time, clotting time, and clot retraction were normal and no haemorrhagic manifestations occurred. The serum bilirubin was less than 0.2 mg. per 100 ml. of blood. Marrow puncture was performed six times in all. In September, 1947, a portion of marrow from the right tibia was removed under general anaesthesia and was examined by Dr. R. G. MacFarlane, who reported on the remarkable absence of red-cell precursors and who considered an erythroblastic hypoplasia, almost amounting to an aplasia, to be well established. Details of two myelograms are given in table 2.

During the year the child was under observation he was remarkably free from intercurrent infections. When his haemoglobin was low he became drowsy and irritable and slight pyrexia was evident, but after a transfusion he became active and playful.

**Discussion**

The number of reported cases of aplasia or hypoplasia of the erythroblastic tissues alone is

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**TABLE 1**

**BLOOD EXAMINATIONS**

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**TABLE 2**

**DETAILS OF TWO MYELOGRAMS**

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very few. Lescher and Hubble (1932), in an
investigation on aplastic anaemia in general, could
find only three recorded adult cases. Each of these
cases survived only a short time after their diagnosis,
the condition appearing to have been relatively
acute. Mills, however, in 1931, recorded the case of
a man aged 50 years in whom the disease ran
an acute course, blood transfusion being ineffective
and death occurring within four months. Bone-
marrow biopsy was not performed. MacFarlane
and Currie (1943) described the condition in a
woman aged 22 years who died after a short illness,
the immediate cause of death being fatal pyrexia
following the second blood transfusion. The
diagnosis in this case was confirmed by marrow
biopsy, which showed aplasia of the red-cell elements.
Kark (1937) described two very interesting cases,
the first a man of 30 years who had two hundred
and ninety blood transfusions in nine years, event-
ually dying from a transfusion reaction. At one
period this patient developed severe agranulocytosis
and eventually haemochromatosis, possibly due to
a failure to excrete unutilized iron. Bone-marrow
biopsy was reported as showing deficient haemo-
poietic tissue. Kark’s second case was a girl aged
21, who had sixty-one transfusions during the three
years she was under observation. She finally
developed thrombocytopenia and died from uncon-
trollable bleeding. Leslie (1945) described the case
of a child who developed an apparently pure
red-cell anaemia six days after birth. This child
recovered after two and a half months, following
blood transfusions. Marrow biopsy was not
performed.

Blackfan and Diamond (1944) described five cases
which seem without doubt to be true examples of
pure red-cell anaemia. Of their patients, all
children, two eventually recovered completely
following blood transfusions, one died of pneu-
monic septicaemia, and two were still under treat-
ment at the time of publication of the paper, the
eldest, aged 11, fully developed and normal in
every respect, but requiring regular transfusion.

Their description of chronic hypoplastic anaemia
arising in infancy, and presumably congenital in
origin, is identical with that of the case recorded
here. In all their cases severe normocytic normo-
chromic anaemia developed within the first three
months of life. There was no familial incidence,
and no history of infection either in the patient or
in the mother during gestation. There was no
evidence of endocrine disturbance and little inter-
ference with growth and development; and with
the exception of the one who died from pneumo-
coccal septicaemia the children were remarkably
free from infection. Blackfan and Diamond’s
statement that after many years’ transfusions
pigmentation suggestive of mild haemochromatosis
develops, corresponds with Kark’s experience as
mentioned above.

Though in their cases the liver and spleen were
not at first palpable, after some years these organs
began to enlarge. The marrow showed aplasia of
the red-cell elements only. Leucocytes and
platelets were always normal. The bleeding times,
clotting times, and clot retractions were normal.
The only treatment they found effective was blood
transfusion at regular intervals of six to eight weeks.
The case here recorded closely fits the above
description, and is of added interest in view of the
associated congenital ptosis. The ultimate outcome
is uncertain, though in view of Blackfan and
Diamond’s report of recovery following repeated
transfusions in two of their cases it appears justified
to continue transfusions in the case described.

Summary

1. A case of chronic hypoplastic anaemia
arising in early infancy is described.

2. From a survey of the literature it would
appear that this disease can occur in infants,
children, and adults, and may be acute or chronic,
presumably depending upon the degree of aplasia
of the red-cell elements in the bone
marrow.

3. The cases arising in early infancy are probably
congenital in origin, but as stated by Wintrobe
(1946), there is no information about the cause of
this disorder.

Our thanks are due to Dr. Facey for the numerous
blood investigations, and to Dr. R. G. MacFarlane
(Radcliffe Infirmary, Oxford), who examined the
bone-marrow sections.

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1, 425.


Wintrobe, M. M. (1946). Clinical Hematology. Lon-
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Chronic Hypoplastic Anaemia arising in Infancy

Thomas Robson and Patrick J. Sweeney

Arch Dis Child 1948 23: 294-296
doi: 10.1136/adc.23.116.294

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