A CASE OF OSTEOPETROSIS (ALBERS-SCHÖNBERG) WITH INTERCURRENT PNEUMOCYSTIS PNEUMONITIS

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Albers-Schönberg (1904, 1907) defined the clinical and radiological features of a sclerosing disease of bone for which Karshner (1926) suggested the term osteopetrosis. As published cases of the disease accumulated, two types came to be distinguished (McCune and Bradley, 1934; Harnapp, 1937; Nussey, 1938; Zetterström, 1958). The first, or clinically 'malignant' type may be characterized by a refractory and often leucoerythroblastic anaemia, enlargement of liver, spleen and lymph nodes, a liability to multiple fractures, and death in infancy or early childhood. In the second or clinically 'benign' type, more commonly seen in adults, anaemia is mild and the osteosclerosis, though often severe, may yet be symptomless.

In the more severe forms of osteopetrosis, intercurrent infection, particularly of the respiratory tract, is a serious hazard. The case to be reported here, an example of the 'malignant' type, succumbed to an interstitial 'plasma cell' pneumonitis.

Though this unusual pneumonitis has been recognized for many years in central Europe its aetiology had been obscure until 1951. In that year Vaněk, in Czechoslovakia, identified Pneumocystis carinii in the lungs of all of 16 fatal cases of this disease. In a series of papers (Vaněk, 1951; Vaněk and Jirovec, 1952; Vaněk, Jirovec and Lukeš, 1953) evidence was presented that Pneumocystis carinii was the causal agent of interstitial plasma cell pneumonitis. Nevertheless, Deamer and Zollinger (1953), who first drew American attention to the disease, concluded that it was probably a viral one; they dismissed a fungal or protozoal agent as unlikely. Baar (1955) reported the first case to be observed in England and subscribed to the hypothesis that Pneumocystis carinii was the pathogen. The subject is comprehensively reviewed by Gajdusek (1957).

Pneumocystis carinii pneumonitis has been found associated with congenital agammaglobulinaemia (Hutchison, 1955; Bird and Thomson, 1957; Russell, 1959), cytomegalic inclusion disease (Baar, 1955; Hamperl, 1956), and neoplastic disease of the lympho-reticular tissue in both children and adults (Vaněk, 1952; Jirovec and Vaněk, 1954; Le Tán Vinh, 1954; Hamperl, 1956). Association with Albers-Schönberg's osteopetrosis has apparently not been reported before.

Case Report

W.T. (Reg. No. R.I. 250922), was a male infant, the first-born and only child of healthy unrelated parents. Pregnancy and labour had been uneventful. In the first 24 hours bruises were noticed on his upper lip and on several toes. These bruises disappeared within a few days. On the second day he regurgitated blood-flecked milk. Nothing further happened until the 13th day when another bruise appeared on his right leg and a small haematoma-like swelling was found on his umbilicus. As these subsided others appeared near the anus and over the chest wall. He became progressively paler.

On admission to hospital at 3 weeks of age, he was pale but reasonably vigorous and not acutely ill. He weighed 5 lb. 14 oz. There were bruises on the left chest, right thigh, right groin and umbilicus. A small abscess cavity was found close to the anus posteriorly. The liver was palpated about a finger breadth below the costal margin, and though the spleen was not palpable at first it could be tipped after four days and readily palpated within a week. Apart from these there were no unusual clinical findings.

Laboratory Investigations on Admission. Hb 45% (6-6 g./100 ml.); erythrocytes showed some anisocytosis and poikilocytosis; there were no Heinz bodies. Leucocytes 34,000/c.mm. (blast cells 340, promyelocytes 700, myelocytes 1,400, band forms 2,700, segmented neutrophils 12,000, eosinophils 700, monocytes 3,400, lymphocytes 13,000); reticulocytes 1-0%; normoblasts (occasionally in mitosis) 5/100 leucocytes. There was, therefore, a severe leucoerythroblastic anaemia (Fig. 1). In addition, the platelet count (Lempert) gave 21,000/c.mm. (a proportion of which were abnormal giant forms) and the bleeding time (Ivy) was prolonged to 13 minutes. Clotting time (Dale and Laidlaw) was normal at 1 min. 50 sec. and the.
Fig. 1.—Peripheral blood film showing immature erythroid and myeloid cells. May-Gruenwald-Giemsa × 584.

tourniquet test was negative. The blood group was O Rh positive. The direct Coombs test was negative and the mother’s serum was found to be compatible with the patient’s erythrocytes. No platelet agglutinins could be demonstrated in the mother’s serum. The toxoplasma dye test dilution was 1/6 and the toxoplasma complement fixation test was negative. Wassermann and Kahn reactions were negative in both mother and infant. Puncture of the tibial marrow was attempted unsuccessfully.

**Progress in Hospital.** Towards the end of the first week in hospital the perineal abscess was evacuated. A scalp vein transfusion of 110 ml. of blood brought the Hb level to 96% (14.2 g./100 ml.). After this there was some clinical improvement, but within 17 days the Hb had fallen again to 58% (8.6 g./100 ml.), i.e. at the rate of about 2.6% (0.3 g./100 ml.) daily although there was no haemorrhage during this period. The fluctuations in the haemoglobin levels are shown in Fig. 2 in relation to blood transfusion and blood loss. Serial leucocyte and platelet levels are shown in the Table.

Further scalp-vein transfusion of 100 ml. of blood restored the Hb to 96% (14.2 g./100 ml.). Five days later he developed epistaxis and began to ooze blood from the gums. About this time it was observed that any skin punctures tended to bleed excessively. During the next three weeks the Hb fell steadily to 46% (6.8 g./100 ml.) necessitating blood transfusion once again. A transfusion of 150 ml. was therefore given by an

![Serial haemoglobin levels](http://adc.bmj.com/content/35/183/495)

**Fig. 2.—Serial haemoglobin levels. Blood transfusions and episodes of bleeding are arrowed.**
antecubital vein. Meanwhile the spleen was noted to be enlarging gradually. Septic skin lesions on the head and on the left hand were treated with oral terramycin.

Towards the end of the second month in hospital it was decided to treat him with prednisolone, at first in a dose of 1 mg. twice daily, increasing over the next four weeks to a maximum of 4 mg. four times a day. With this his general condition improved considerably, although he continued to show purpuric lesions over the skin and abdomen. The platelet count began to rise shortly after administration of prednisolone and reached the normal level of 240,000/c.mm. during the subsequent eight weeks.

Nevertheless, episodes of bleeding continued. Ten days after the start of prednisolone treatment severe epistaxis began again and continued in bouts during the next three weeks. It was eventually controlled by intranasal adrenaline sprays. By this time Hb level had fallen to 48% (7·1 g./100 ml.), now at the rather slower rate of 1·4% daily. A further blood transfusion of 100 ml. was given.

Marrow puncture was attempted again and yielded, with some difficulty, a hypocellular but otherwise normal marrow. As a fortnight passed without blood loss, and he was in good general condition, he was discharged home on 16 mg. of prednisolone daily.

While at home he caught a "cold" and developed a cough. He went off his feeds, which he had been taking well till then. He was therefore re-admitted, now at the age of 4½ months. On examination his temperature was 99°-6° F. and his breathing rate was 64 per minute. His colour was dusky and he looked ill. There was some nasal discharge and reddening of the fauces.

His breathing was grunting in character; the alae nasi were working and there was some intercostal recession. However, the percussion note and the breath sounds were normal and there were no adventitiae. The liver and spleen were still palpable but now smaller. A chest radiograph showed no definite radiological lesion.

Two days later there was a fresh crop of purpuric lesions on his abdomen, although his platelet count was now normal. He maintained a persistent low-grade fever with some cough but no physical signs in the chest. After a further transfusion of 180 ml. blood, which brought his Hb level from 62% (9·1 g./100 ml.) to 95% (14·1 g./100 ml.) his general condition improved though his breathing remained rapid and there was some further bleeding from nose and rectum. At this time the spleen had receded so that it could only be tipped. He was feeding well and seemed happy. He was discharged once again on prednisolone and oral erythromycin.

He was re-admitted five days later because his cough had become worse and he had gone off his feeds again. On examination he was pale and ill and had a weak cry. His breathing was still rapid though his chest appeared clinically and radiologically clear. The liver was palpable two or three finger breadths below the costal margin; the spleen could not be felt. He remained somewhat febrile, breathing at the rate of 60-120/min. On the 10th day of this admission he died suddenly at the age of 6 months.
Morbid Anatomy. At autopsy (P.M. 512/58) the
following were the main macroscopical findings:

**MACROSCOPICAL FINDINGS**

**Lungs.** The lungs failed to retract as the chest was
opened. Both were of a mottled café-au-lait colour.
They presented a curious, vaguely nodular consolidation
which was more marked in the lower lobes. The cut
surfaces were dry and showed small scattered yellowish-
white areas. The tracheobronchial tree was normal.

**Liver.** The liver (196 g.) appeared normal in size,
consistency and colour.

**Spleen and lymphoid tissue.** The spleen (16 g.) was
rather firm, but not appreciably enlarged. The cut
surfaces appeared normal. Lymph nodes in general
were inconspicuous. The thymus was inviolated.

**Bones.** The calvarium appeared to be of average
thickness and density. The features of the base of the
skull were not detectably abnormal and the middle ears
were opened without difficulty. Both femora were
shorter than normal and very slightly clubbed. They
proved very difficult to cut because of a dense sclerosis
of the medullae. A vertebral body was similarly
sclerosed. These excised bones and the remainder of
the skeleton were radiographed and showed the following
features: There was an overall increase in the density of
the femur as compared with a normal control (Fig. 3).

Trabecular detail was obscured and marrow spaces
appeared small. Particularly dense transverse bands
occupied the metaphysial growth zone; that in the distal
metaphysis of the femur measured more than 1 cm.
in the axis of the bone. The distal epiphysial centre of
ossification was rather small and dense. Very similar
changes were visible symmetrically in tibiae, fibulae,
humeri, radii and ulnae. Clubbing was more obvious
at the distal ends of the radii and the tibiae than else-
where. Similar but slighter bands of metaphysial
density were visible in metacarpal bones. Pelvic bones,
vertebrae, ribs, clavicles and scapulae showed some
overall increase in density. In carpal, tarsal and skull
bones the changes were too slight for dependable assess-
ment.

**HISTOLOGICAL FINDINGS**

**The Lungs.** Both lungs showed an overall interstitial
fibrosis. This was mainly due to an increase in reticulin
but occasional collagen bundles contributed to it. The
thickened alveolar walls and septa were infiltrated by
mononuclear cells, some of which resembled plasma cells.
Polymorphonuclear leucocytes were not seen in signif-
icaent numbers. The alveoli and alveolar ducts in most
areas contained an eosinophilic foamy material which was
bordered in some alveoli by coalescing macrophages
(Fig. 4). This material gave the tingiolar reactions of a
polysaccharide (periodic acid-Schiff) through which small
chromatin bodies (Giemsa: Feulgen) were dis-
tributed. It did not contain fibrin (Mallory's photo-
tungstic acid-haematoxylin) or mucus (mucicarmine).
Silver impregnation (Hortega) confirmed the presence of
morphologically typical *Pneumocystis carinii* (Fig. 5).
Viral studies were not made. The appearances were
those of an interstitial pneumonitis in which the only
detectable organism was the *Pneumocystis carinii*.

**The Liver.** The parenchymal cells contained rather
large quantities of stainable iron.

**The Spleen.** The sinusoids were prominent and their
littoral cells were plump. Many sinusoids contained
large pale free macrophages. These phagocytes con-
tained nuclei or fragments of nuclear material, a good
deal of iron pigment and an occasional red cell ghost
(Fig. 6). There was some excess of reticulin in the
stroma. No evidence of haemopoiesis was found.

**Lymph Nodes.** These showed a fairly well marked
excess of reticulin. Active haemopoiesis was visible in
some nodes.

**Bones.** The disturbances affected endochondral ossi-
fication only mildly. Those seen in the femur will be
taken as illustrative. *Mutatis mutandis*, similar changes
were seen in other sites. The most striking feature was
a massive persistence of calcified cartilage (the primary
spongiosa) as irregular longitudinal islands, bordered
and occasionally penetrated by immature new bone
(Fig. 7). In the electron microscope the calcification of this
cartilage was seen to be patchy but often unusually
dense. The metaphyses were thus packed with stout
anastomosing trabeculae of unresorbed primary elements
and the enclosed narrow spaces were correspondingly
small. This thicket extended in a modified form to the

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**FIG. 3.—Femur showing osteopetrosis compared with normal femur of similar age.**
diaphysis. Here spongy bone was still excessive but the cartilaginous component was diminished and the marrow spaces were considerably larger. In polarized light much of the new bone bordering the persistent cartilage showed normal lamellation. On the other hand the bone which had formed within vacated cartilage cell capsules showed no orientation of its collagen fibres. Both varieties of new bone were Schiff-positive.

Periosteal ossification was less disturbed but Haversian systems appeared poorly formed in many areas. Occasionally the cortex included some persistent cartilage and immature bone.

**Bone Marrow.** The small marrow spaces in the growth zone contained an excess of reticulin through which ran thick-walled capillaries. Normal bone similarly stained showed more delicate mesenchyme and larger thin-walled vessels in this region. The osteopetrotic metaphysis appeared less vascular than usual and the capillaries morphologically abnormal.

Haemopoietic tissue occupied the marrow spaces in the remainder of the bone. Erythropoiesis and leucopoiesis were both in progress and occasional mega-karyocytes were seen. There appeared to be a mild hyperplasia of myelocytes, and colonies of these cells were especially noted along the adventitia of small
arterioles and along the borders of many trabeculae. Fat cells were very scanty. This marrow differed from normal marrow, similarly stained, in being more densely cellular and in not showing the usual prominence of marrow sinusoids. Where sinusoids could be discovered they contained both erythropoietic and leucopoietic cells, an appearance which suggested intravascular haemopoiesis.

Other Organs. Apart from some foci of calcium deposition in the medulla of the kidneys other organs were unremarkable.

Discussion

Generalized osteosclerosis in infancy may be secondary to congenital syphilis, heavy metal poisoning, excessive dosage with vitamin D and perhaps leukaemia. Of these diseases only leukaemia merits consideration in the present case. However, leukaemia typically causes osteolysis and on the rare occasions when it causes sclerosis this is found to be due to direct metaplasia of reticular or collagenous structures into woven bone (Windholz and Foster, 1949). Therefore this disease also may be ruled out.

Albers-Schönberg’s osteopetrosis, on the other hand, accounts satisfactorily for both the clinical and the pathological findings. The clinical features closely resemble those of cases reported by Pines and Lederer (1947) and Turano, Fagan and Corbo (1954). The pathological structure of the bone is typical of osteopetrosis. As in previously reported cases the most obvious and probably the primary abnormality is a persistence of the calcified cartilaginous scaffolding upon which new bone is built. Layers of new Schiff-positive bone are seen on the surfaces of the cartilage masses, and tinctorially similar bone appears in the spaces from which cartilage cells have vanished. Normally a provisional design of this sort undergoes considerable modification. Most spongy bone and all remnants of cartilage are resorbed from the marrow cavity and increasingly lamellated bone is incorporated peripherally into the cortex. The appearances in the present case may be interpreted as the effect of a retarded remodelling process. Relatively little lamellar bone is present. Islands of immature bone, of which the collagen is unorientated, form a mosaic with the persistent irregularly calcified cartilage. The tensile strength of such tissue must be well below normal in spite of its apparent solidity.

Either the cartilage matrix is unusually resistant to remodelling or the mechanism of remodelling is defective. Of course, both factors may cooperate, but there is no evidence at present of abnormal cartilage matrix. On the other hand the vascular metaphyseal mesenchyme, which plays a major part in resorptive modelling, may well be at fault. Evidence in favour of this thesis may be found in the experimental work of Trueba and Amato (1960). They found that interruption of the blood supply to the metaphysis of growing bone inhibited the resorption of chondro-osseous tissue and delayed the formation of final bone. Large zones of calcified cartilage and preliminary bone remained at a level in the metaphysis where, normally, remodelling is taking place.

Many authors (McCune and Bradley, 1934; Lamb and Jackson, 1938; Clifton, Frank and Freeman, 1938; Pines and Lederer, 1947) have adopted Klemperer’s (1931) suggestion that the fundamental disturbance in Albers-Schönberg’s disease may lie in the undifferentiated mesenchymal anlage, the common progenitor of bone and marrow. Osteogenesis and haemopoiesis, according to this hypothesis, are both blighted by a prior defect involving mainly the former process in benign cases and the latter in malignant. But in the present case histological study did not suggest an intrinsic defect of osteogenesis. Similarly, Zetterström’s (1958) biophysical studies led him to the conclusion that osteoprotic bone comprises all those types of skeletal tissue which appear during the normal development of bone, the abnormality being a failure of normal removal. Furthermore, Pines and Lederer (1947) noted that the skeleton in general was poorly vascularized but that bone and marrow approached normal at sites of enhanced vascularity. In the present case also an impression of poor skeletal vascularity and of abnormal vessel structure was gained. The hypothesis of mesenchymal perversion might therefore be restated in a form which postulates abnormal vasoformation rather than abnormal osteogenesis. This view would suppose a mesenchymal defect which may involve predominantly the vascular remodelling tissue in benign osteopetrosis or the closely allied haemopoietic tissue in malignant osteopetrosis.

As in the cases of Lamb and Jackson (1938), Kramer and Halpert (1939), and others, the refractory anaemia could certainly not have been due to a simple crowding out of otherwise healthy haemopoietic tissue, as is commonly supposed to happen. In the present case the volume of the marrow cavity was estimated and found to be fully 70% of normal. Moreover, the marrow tissue was a good deal more cellular than normal. Therefore a merely mechanical theory of dyshaemoipoiesis is plainly inadequate.

Though one may hypothetically attribute the leucoerythroblastic anaemia to a dysplasia of
marrow tissue other mechanisms must also be evaluated. Skin sepsis and episodic haemorrhage must have played their parts, though the haemoglobin level continued to fall on occasions when neither was operative. Red cell destruction probably played a major part. Zetterström (1958), using a radioactive chromium technique, found a mean red cell survival time of 35-40 days in one of his cases, a patient aged 10 years. With colleagues (Engfeldt, Karlberg and Zetterström, 1955) he found a shortened survival time of both native and donor cells in another patient aged 6 years, though this observation was not expressed quantitatively. He therefore suggested that the anaemia may be due to hypersplenism. Loeb, Moore and Dubach (1953) have found this to be true of certain cases of myelosclerosis; in these, splenectomy restored a shortened red cell survival time to normal. Sjölin (1959) reported four cases of malignant osteopetrosis in each of which accelerated red cell destruction was proved isotopeically. Splenectomy improved the haemoglobin level in all cases, though the red cell survival time was not always restored permanently to normal. The platelet count, which had been low in all four cases, returned to normal and reticulocytosis diminished. The author concluded that hypersplenism was an important feature of malignant osteopetrosis.

Though no direct estimate of red cell survival was made in the present case the concept of hypersplenism is apposite. Admittedly a cellular marrow is not necessarily a productive one but the abundance of maturing cells, together with the presence of normoblasts in the peripheral blood and a continuous reticulocytosis, suggest that the anaemia was not wholly due to a failure of production. Furthermore, the steep fall of haemoglobin level (2.6% daily before administration of prednisolone), the marked haemosiderosis of liver and spleen (partly due to transfusion, no doubt) and the apparently enhanced activity of splenic phagocytes suggest excessive destruction.

The leukaemoid reaction is commonly supposed to be an expression of extramedullary haemopoiesis, at least in part; immature cells escape into the blood stream more easily from ectopic sites in soft organs. But the extent of haemopoiesis in the tissues at autopsy was surprisingly slight, as in many previously reported cases (Reiche, 1929, quoted by McCune and Bradley, 1931; Pease, DeSanctis and Alter, 1931; Lamb and Jackson, 1938). An alternative cause may lie again in dysfunction of the bone marrow itself, for which the observed myelocytic hyperplasia and the apparently intrasinusoidal haemopoiesis afford some evidence.

The tendency to haemorrhage is not completely explicable by thrombocytopenia and may have had a thrombathenic component. Siegel, Friedman and Schwartz (1957) reported a patient with osteogenesis imperfecta tarda and an associated haemorrhagic diathesis due to increased capillary fragility and abnormal platelet function. These authors believed that their case represented a congenital defect involving two tissues of mesenchymal origin, the osseous system and the platelets. In the present case morphologically abnormal (giant) platelets were present and the haemorrhagic tendency persisted after the total platelet count had returned to normal.

Treatment with prednisolone brought about some improvement in the child's general condition and appeared both to decelerate the fall of haemoglobin level and to correct the leukaemoid state. The sclerosing process in the skeleton, on the other hand, was not abated. This is surprising, in view of the experimental findings of Baker and Ingle (1948) that 11-oxy steroids inhibited the proliferation of cartilage cells and brought about a profound and general atrophy of osteoblasts, so that the resorption of metaphyseal trabeculae occurred more rapidly than their formation. One might have expected prednisolone to be the rational treatment of osteopetrosis.

Prednisolone may have predisposed to infection with Pneumocystis carinii by further depression of a naturally low resistance. The result was the development of an interstitial pneumonitis of a type usually found in premature weakly infants or complicating diseases of the lymphoreticular system. It is of interest, furthermore, that prior and current treatment with prednisolone did not inhibit the development of a marked interstitial pulmonary fibrosis.

Summary

A case of 'malignant' osteopetrosis is reported. The patient presented in the neonatal period suffering from a leuco-erythroblastic anaemia complicated by a haemorrhagic tendency, and died at the age of 6 months primarily of an interstitial/pneumonitis due to Pneumocystis carinii. An attempt is made to illustrate some of the problematic features of osteopetrosis. Future research might be directed towards studying the vascularity of the bone, evaluating the respective roles of dyshaemopoiesis and increased red cell destruction, and determining the adequacy of platelet function.

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