OBSERVATIONS ON THE HAEMORRHAGIC DIATHESIS IN MULTIPLE MYELOMA

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ABNORMAL bleeding has been recognized as a presenting symptom in multiple myeloma (Wintrobe, 1961) though less frequently encountered than pain, tumours, deformity, pathological fractures and neuralgic or neurological symptoms. Bernard, Inceman, Zara and Christol (1952) comment that hæmorrhage is found more frequently as a secondary or terminal feature than as a presenting symptom, but concede that it is common. In their series of 100 cases hæmorrhage was present in 37, in 18 cases moderate and in 19 cases severe. Of 13 cases described by Rosenthal and Vogel (1937) hæmorrhage occurred in five. While Lichtenstein and Jaffé (1947) do not include abnormal bleeding as a common presenting symptom, they refer to a purpuric tendency and call attention to a disorder of the clotting mechanism, characterized by poor clot retraction, which may occur. Osserman comments on bleeding into the gastro-intestinal tract as a feature of the disease (1959).

It is the purpose of this paper to report a case of multiple myeloma in which the presenting symptoms were those of hæmorrhage, together with an investigation into the nature of the abnormal bleeding.

Case Report

A female, aged 40, was first seen in Moorfields Eye Hospital five days after she had noticed a 'reddish blur' over the left eye. On examination of the fundus the retinal veins were seen to be turgid and there were retinal hæmorrhages. Increased viscosity of the blood was noted and it was thought this might be causing retinal vein obstruction. Hess's test was positive. She was referred for a further opinion and admitted to a general hospital two days later.

She complained of tiredness and flagging energy for several months and of severe headache. She was unable to read due to blurring of vision in the left eye. Her last three periods had been excessively heavy, the menses soaking rapidly through sanitary towels and on occasion soiling her outer garments. During her last period, a fortnight before admission, she had passed several large clots. After her periods she felt exhausted. In the past six months she had had frequent epistaxes. For a few weeks she had noticed shortness of breath. For as long as she could remember she had always bled for several hours following tooth extraction. Since the birth of her child 11 years before (F.T.N.D.) she had suffered from bleeding piles. There was no history of haemoptysis, hematemesis, melena, hemorrhia or purpura. There was no family history of anaemia or abnormal bleeding.

On examination she was a pale, ill-looking woman with a muddy complexion. There was marked pallor of the mucous membranes, but no jaundice. There was no purpura and Hess's test was negative. The temperature and pulse rate were normal. The liver and the spleen were each enlarged two fingers' breadth. There were no other physical signs except a grade two systolic murmur best heard at the apex and conducted up into the neck.

Investigations. ESR (Westergren) 140 mm./hr., Hb 33% (4.8 g.), R.B.C. 1,240,000. PCV 13%, MCHC 36%, MCV 104 cu. mm. Serum calcium: 9.2 mg./100 ml. Urinary urobilinogen was not increased. Blood ura. 30 mg./100 ml. Albumin was present in the urine, but no Bence Jones protein was detected. No casts or casts were seen in the centrifuged deposit. Total serum protein: 14.0 g./100 ml. Albumin 2.1, globulin 11.9, A/G ratio 0.2 : 1. The electrophoretic pattern showed a gross excess of gamma globulin. Serum calcium: 9.2 mg./100 ml.

In view of the disorder of the clotting mechanism, blood was cross-matched before the sternal marrow was aspirated. This marrow showed the presence of numerous plasma cells (approximately 25% of the total nucleated cells) and also partially megakaryoblastic erythropoiesis. The appearances were compatible with myelomatosis.

A chest X-ray showed some cardiac enlargement with enlargement of the pulmonary conus and of the main pulmonary vessels. The appearances were compatible with a hyperdynamic circulation. No lung lesion was seen. X-ray of the lumbar spine and pelvis showed the bones to be slightly osteoporotic. There was also some suggestion of multiple translucencies in the ischial bones. These appearances were thought to be the early changes of multiple myeloma. The skull X-ray was within normal limits.

Clinical Course and Treatment. The diagnosis made on these findings was multiple myeloma. The patient was transfused with 4 pt. of whole blood and the haemoglobin rose to 51% (7.5 g.).

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She was started on a course of prednisolone, on which she remained until her death three to four weeks later. A further transfusion of 4 pt. of packed cells was given. Meanwhile the pains in her back and legs gradually grew worse and she required morphine. Before death she was bleeding heavily per vaginam and had a haematoma over the left buttock and a left hemiparesis.

**Subsequent Investigations.** The great viscosity of the blood and poor clot retraction made investigations difficult. Serum was separated off with difficulty even at 37°C. Blood for the last transfusion could be cross-matched only by the Coombs technique owing to marked rouleaux formation.

The total serum protein before death was 16.7 g./100 ml., of which only 1.9 g. was albumin and the rest globulin. The formol gel test was positive and the Kunkel test greater than 10 units. The daily urine output was 1 to 2 l. On admission the average daily output of protein in the urine was 2 to 3 g./l. (Esbach), but this output steadily rose and terminally was variable with a maximum of 10 g./l. and an average of 7 g./24 hr.

Viscosity studies using a capillary frame viscometer showed a plasma viscosity about seven times greater than normal and a whole blood viscosity about two and a half times the normal in the later stages of her illness. The haemoglobin before death was 54% (8.0 g. %). Feupptake test showed that the bone marrow was almost entirely non-functioning. The erythrocyte sedimentation rate remained elevated and was last recorded as a fall of 168 mm./hr. (Westergren). Terminally there was a leucocytosis of 36,000 cells/cu. mm. with 50% plasma cells and the blood urea was 64 mg./%.  

**Investigation of Bleeding Diathesis.** Initially the patient was found to have a prolonged prothrombin time (27 seconds with control of 17 seconds), a normal clotting time and a prolonged bleeding time (30 minutes). The platelet count at that time was 123,000/cu. mm. Clot retraction was noted to be consistently poor.

An injection of vitamin K was given and the patient transfused with 4 pt. of whole blood. Thereafter the bleeding time was 10 minutes and the prothrombin time 31 seconds with control of 17 seconds. The platelet count was 98,000/cu. mm.

The prothrombin time repeated a week later was 25 seconds with a control of 17 seconds. The thromboplastin-generation test showed no evidence of deficiency of antihemophilic globulin, Christmas factor or the St邠er Prower factor. Platelet thromboplastic function was normal and there was no evidence of a circulating anticoagulant. By mutual correction studies using the thromboplastin-generation test with known Factor V, Factor VII and prothrombin-deficient blood samples, and by the two-stage prothrombin test, the prolongation in the prothrombin time was shown to be due to a Factor VII deficiency, with no deficiency of Factor V or prothrombin. The serum fibrinogen was 250 mg./100 ml. (biuret).

** Necropsy Findings.** Death was due to pulmonary edema. There were scanty cutaneous petechial haemorrhages. The principal pathological findings were: pulmonary edema and left ventricular hypertrophy, anemia, petechial haemorrhages in the brain and the heart, and multiple myeloma affecting bone, spleen and probably liver. The marrow of the sternum, ribs, vertebral and right femur was diffusely infiltrated by soft greyish semi-translucent tumour tissue which in places was haemorrhagic. The spleen was grossly enlarged (1,078 g.) and markedly congested with irregular areas (up to 1 cm. diameter) of greyish tissue throughout its substance. The liver was markedly enlarged (2,704 g.), pale and fatty. Histologically, there was multiple myeloma invading bone, liver, spleen and lymph nodes. The kidney was abnormal with deposits of protein. There was no evidence of amyloidosis.

**Discussion.**

The aetiology of the abnormal bleeding in multiple myeloma has not been clearly elucidated although it has occasioned much discussion (André, Dreyfus, Jacob and Ley, 1952). It has been explained as secondary to the low platelet count and prolonged bleeding time (Wintrobe, 1961), but the latter may be normal or increased without relation to the platelet count. Moreover, the platelets may be within normal limits. Defective platelet function (thrombocytopenia) has been noted and may contribute to bleeding when the platelet count is normal. In one case giant platelets measuring 5 μ diameter were observed (Czernobilsky and Alexander, 1961). Capillary damage from impaired blood flow due to the often greatly increased blood viscosity has also been considered as a factor in the development of the haemorrhagic diathesis (de Gruchy, 1958).

Since bleeding may occur in other disorders characterized by abnormality of the plasma proteins (Owen and Got, 1960), these may play a more direct aetioloigical role: possible modes of action of abnormal protein include infiltration of the vascular wall and interference with the mechanism of coagulation (de Gruchy, 1958).

A disorder of the clotting mechanism proper has been observed although the essential nature of the disturbance is obscure. Wintrobe (1961) comments that cutaneous haemorrhagic manifestations such as purpura and massive subcutaneous hæmorrhages may develop as the result of this disturbance. Failure of clot retraction and subsequent difficulty in obtaining adequate samples of serum for pathological investigations seem to be encountered relatively frequently (Stewart and Weber, 1938). The observation was first recorded by Runeberg in 1883 (Frick, 1955). Lichtenstein and Jaffé (1947) commented on failure of clot retraction, and of the five haemorrhagic cases reported by Rosenthal and Vogel (1937) one showed failure of clot retraction in addition to thrombocytopenia. Four of their 13 cases had platelet counts below 100,000/cu.mm., but the one mentioned above was the only thrombocytopenic case in which there was abnormal bleeding. Osserman (1959) states that in a small number of cases there is a defect in the clotting mechanism, the nature of which he does not specify. He considers that thrombocytopenia, present in about a third of cases, may contribute to bleeding.

The prothrombin time, thrombin time and conversion of fibrinogen to fibrin may all be abnormal. In some cases an anticoagulant (anti-
thrombin V) has been demonstrated. Increase in the one-stage prothrombin time (Quick) may be due to the antithrombin, but Verstraete and Vermeylen (1959) thought that there was no relationship between the presence of the antithrombin and the haemorrhagic phenomena which occur in some cases of myelomatosis. Craddock, Adams and Figueroa (1953) confirmed the observations of Lüscher and Labhart (1949) and Uehlinger (1949) that the conversion of fibrinogen to fibrin is prevented by an abnormal plasma protein, and showed that this defect could be largely overcome by the addition of an excess of calcium ions. They offered the conjectural explanation that impairment of fibrinogen conversion may lie partly in deficient availability of ionic calcium due to interference by the abnormal protein, which may be a reflection of its high calcium binding power. Frick (1955), in a study of 45 patients with myeloma, observed abnormal conversion of fibrinogen to fibrin by a circulating anticoagulant in 12 patients. In two, who were studied in detail, the anticoagulant occurred in the beta and gamma globulin fractions. From the available data he concluded that only a severe impairment of fibrinogen conversion, with clot retraction virtually absent, causes haemorrhagic symptoms per se, but suggested that lesser degrees of this abnormality combined with mild thrombocytopenia induce bleeding. He recalled that uremia, amyloidosis (Magnus-Levy, 1933) and cryoglobulinemia (Wintrobe and Buell, 1933) may all contribute to the bleeding.

Bernard and others (1952) considered that the causation of the bleeding is multifactorial, but principally falling into two categories: in the first, bleeding secondary to the low platelet count and etiologically resembling the haemorrhages of the leukemic states; in the second, unrelated to the platelet count. As examples of the second type they cited two cases in which they regarded the cause of the bleeding as the result of a toxic effect of the dysglobulinemia on the capillaries. They commented that clot retraction is usually normal, prothrombin normal or slightly depressed, the coagulation time usually normal and fibrinogenopenia not found although a qualitative alteration is possible. They placed little emphasis on the disorder of clotting which is sometimes encountered.

It seems likely that any or all of the factors mentioned above may operate to bring about abnormal bleeding in myelomatosis, the interesting feature in the etiology being the multiplicity of factors which may be contributory.

In our case there was no uremia, cryoglobulins were not detected and there was no histological evidence of amyloidosis. It seemed unlikely that the bleeding could be attributed solely to thrombocytopenia since the lowest recorded platelet count was 98,000/cu.mm. Hess's test, however, was found positive on one occasion soon after the observation of the retinal haemorrhages. Qualitative deficiency of the platelets was ruled out by demonstration of a normal thromboplastic activity. The blood was extremely viscous, particularly in the late stages of the illness, so it is possible that capillary anoxaemia mediated by poor blood flow, or a direct toxic action on vessel walls by abnormal protein, might be an associated cause of haemorrhage in this case. There was little evidence to support this since there was no external purpura until just before death, and Hess's test, on all but one occasion, was negative. At necropsy, however, there were petechial haemorrhages in the heart and the brain.

The failure of clot to retract, the prolongation of the bleeding time in the absence of thrombocytopenia and the prolongation of the prothrombin time led us to suppose that we were dealing with a simple disorder of the clotting mechanism which might explain, either wholly or in part, the haemorrhagic manifestations. A Factor VII deficiency was demonstrated.

Although the patient had volunteered that from early childhood she had bled for hours following tooth extraction it seemed unlikely that the Factor VII deficiency was congenital in this case. There was no family history of anaemia or bleeding tendencies and there were no other symptoms in early life. Congenital deficiency of Factor VII is a rare disease, the symptoms usually being severe and manifest in childhood.

The deficiency of Factor VII was probably related to poor liver function rather than to malabsorption of vitamin K since vitamin K administered parenterally had no effect on the prothrombin time which remained prolonged. During life the liver was palpable and the Kunkel test over ten units indicating derangement of liver function. At necropsy the liver appeared to be affected by myelomatosis and this was confirmed histologically. It seemed, therefore, that the defect was acquired and probably secondary to liver disease. Another explanation of the Factor VII deficiency might be a qualitative alteration in the clotting factor. Such an effect might be produced as a consequence of the abnormal protein synthesis which characterizes the disease, the clotting factors themselves being involved in the abnormal and excessive synthesis of protein. Alternatively, deficiency of the factor might have been secondary to the over-elaboration of abnormal protein replacing normal synthesis of the clotting factors.

Although in this case as in other reported cases
there was no single factor which could be considered as solely responsible for the hemorrhagic state, there were a number of additive factors, namely, moderate reduction in Factor VII, greatly increased plasma and whole blood viscosity, abnormal proteins with a marked increase in gamma globulin, and a slightly reduced platelet count. In addition there was a very poor clot retraction, which may have been due to poor conversion of fibrinogen to fibrin (Craddock and others, 1953).

Since it is recognized that increased bleeding with trauma may be induced by a simple failure of the coagulation mechanism alone (Stefanini and Dameshek, 1955), it seemed probable that in our case a moderate deficiency in Factor VII, together with the other factors mentioned above, would suffice cumulatively to explain the clinical state and the abnormal laboratory findings.

Unfortunately the patient died before an attempt could be made to control the bleeding diathesis apart from administration of vitamin K. The last blood transfusion she received consisted of packed cells.

The part played by specific defects in the clotting mechanism in multiple myeloma, even where these are of moderate severity, may be of importance clinically, since such defects may be remediable. Even in the presence of thrombocytopenia or other explanation of hæmorrhage it would seem worth while to investigate the clotting mechanism, since this may influence the decision to give transfusions of packed cells or of whole blood, fresh or stored.

Summary

A case of multiple myeloma is described in which the presenting symptom was retinal hæmorrhage accompanied by epistaxes and recent excessive menstrual bleeding. A disorder of clotting was discovered and a deficiency of Factor VII demonstrated. It is suggested that the clotting mechanism should be investigated in all cases of multiple myeloma in which there is abnormal bleeding, as replacement therapy may be indicated and could, if the defect were severe, ameliorate the course of the disease.

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