ANAEMIA AND POLYCYTHAEMIA WITH RENAL DISEASE

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Anaemia has long been recognized as a feature of renal failure but the relation between renal function and the bone marrow has been slow to unravel. Much has been learned in this field in recent years, but a great deal remains unexplained, and the anaemia continues to defy all treatment short of transfusion.

Red Cell Morphology in Renal Anaemia

In early studies by Ashe (1929) and Parsons and Ekola-Strolberg (1933) the anaemia was found to be associated with erythrocytes normal both in size and haemoglobin content. A striking feature was the absence of any disturbance of white cells or platelets. Studies of bone marrow morphology by Dameshek (1935) and Nordenson (1938) revealed the same pattern. Erythropoiesis appeared qualitatively normal, but red cell precursors were markedly reduced in numbers. Nordenson recorded the interesting and apparently paradoxical finding that the erythroid precursors showed a normal incidence of mitoses despite the reduced number of these cells. This observation takes on a new significance in the light of recent studies of the causes of this bone marrow depression. Gingold, Comsa and Roman-Crivat (1938) found evidence of increasing aplasia of the marrow as renal failure progressed but the haemoglobin level correlated poorly with urea retention. Some investigators, on the other hand, found no indications of defective erythropoiesis, assessed by marrow biopsy, an actual increase in number of normoblasts being noted on occasions (Scott, 1939; Callen and Limarzi, 1950), which would suggest either arrested maturation without morphological abnormality, or decreased life span of the red cell as the cause of the anaemia.

Haemolysis and Red Cell Life Span in Renal Anaemia

In patients with acute renal failure and less commonly in terminal uraemia, a fall in haemoglobin level may occur which is too rapid to be attributed to defective red cell production. Uraemia may be associated with a marked bleeding tendency, and concealed haemorrhage may contribute to the anemia in some cases. However, there is now clear evidence that red cell survival may be reduced in some cases of renal anaemia in the absence of detectable bleeding, the usual inference being that haemolysis occurs. The occasional appearance of gross morphological changes in red cells in acute uraemia, the 'burr cells' of Schwartz and Motto (1949), supports this view, but the hallmarks of haemolysis—hyperbilirubinaemia, increased urobilinogen excretion, or reticulocytosis—are not found (Loge, Lange and Moore, 1958). Red cell life span has been estimated in renal anaemia using the Ashby cross-transfusion technique (Emerson and Burrows, 1949; Chaplin and Mollison, 1953; Loge, Lange and Moore, 1950 and 1958) and using a radioactive chromium label for the patients' own red cells (Sutherland, McCall, Jones and Muirhead, 1955; Joske, McAlister and Prankerd, 1956; Rees, Scheitlin, Pond, McManus, Guild and Merrill, 1957; Verel, Turnbull, Tudhope and Ross, 1959).

In each of these studies shortening of red cell survival was demonstrated in some patients although many of the studies are open to criticism on technical grounds as care was not taken to exclude artifacts due to changing plasma volume or red cell mass during the period of observation. (The former is likely to be of considerable importance in acute or terminal renal failure, leading to dilution of labelled cells per unit volume of blood; the criticism applies to both the radiochromium and Ashby methods as usually carried out.) Shortening of erythrocyte life span in renal anaemia may, in fact, be considerably less common than reported in the earlier studies. Verel and others (1959) used the radiochromium technique with repeated estimation of red cell volume (employing a second isotope, radioactive phosphorus) and found a shortened red cell life span in only five of 40 patients with anaemia associated with nephritis, one being a patient with coexistent Hodgkins disease, which may have been the cause of the haemolysis. Of the five patients with shortened red cell survival, three showed...
malignant hypertension with papilledema, and a fourth was in terminal uremia. Whether the shortened life span was due to intravascular haemolysis or could have been attributed to extravasation of red cells into tissues (consequent on widespread vasculitis) could not be determined. It is of considerable interest, however, that in one patient with malignant essential hypertension, anaemia was associated with shortened red cell survival, despite a normal blood urea; effective treatment of the hypertension was followed by a return of the haemoglobin level and the red cell survival to normal.

The studies of a number of investigators have indicated that the shortening of red cell survival is due to extracorpuscular factors rather than to a red cell defect. Chaplin and Mollison (1953) and Loge and others (1958) observed that whilst the survival of erythrocytes from normal donors was shortened in patients with renal anaemia, red cells from the uraemic subjects survived normally in healthy recipients. Rees and others (1957) studied the effect of artificial dialysis on red cell life span in chronic renal failure and found that the rate of destruction of erythrocytes returned towards normal four to six days after dialysis. Muirhead, Stirman and Jones (1960) reported that bilaterally nephrectomized dogs maintained by peritoneal lavage showed a marked shortened red cell survival, but that this was corrected by explantation of renal medullary tissue to the peritoneal cavity. These remarkable observations are difficult to evaluate; from the studies in human subjects it would appear that the defect is correctable by dialysis but it remains unknown whether a toxic factor operates directly on the circulating erythrocyte, or whether the process is indirect, damaging vascular endothelium with consequent extravasation of red cells into the tissues. Possibly both processes play a part under certain conditions.

Haemolysis may clearly contribute to the anaemia of renal disease, but it has been found principally in patients with acute or terminal uraemia, or with malignant hypertension. It is unlikely that it plays a major role in the anaemia associated with other stages of chronic renal disease, where defective red cell production appears to be the primary disorder.

**Isotopic and Chemical Studies of Erythropoiesis**

The use of the radioactive iron isotope $^{59}$Fe has largely confirmed the earlier morphological studies of erythropoiesis. Thus Finch, Gibson, Peacock and Fluharty (1949), Loge and others (1950 and 1958), and Joske and others (1956) all demonstrated markedly decreased red cell production in chronic renal failure.

Evidence of iron deficiency is seldom found in such patients (Cartwright, Huguley, Ashenbrucker, Fay and Wintrobe, 1948; Loge and others, 1958; Verel and others 1959) and when present is likely to be concidental. Cartwright and others (1948) found elevation of free erythrocyte protoporphyrin in six of seven patients with uremic anaemia, a finding similar to that in other anaemias with defective erythropoiesis, but changes in plasma iron, iron-binding capacity or copper levels typical of toxic marrow depression as seen in infection or heavy metal intoxication were not observed. Loge and others (1948) could detect significant elevation of free erythrocyte protoporphyrin in only eight of 21 patients with renal anaemia, and such elevation could not be correlated with the serum iron level, the degree of azotaemia, or the severity of the anaemia. In the light of this evidence it would be difficult to attribute the undoubted defect in red cell production in uremia to simple toxic depression of erythropoiesis. Recent experimental studies have, however, implicated the kidney in the physiological regulation of erythropoiesis, and have opened a new approach to the problem of anaemia in renal failure.

**Relation of the Kidney to Erythropoiesis**

**The Humoral Hypothesis**

Our understanding of the mechanisms by which the body finely regulates the production of red cells to meet its requirements for oxygen transport is still very inadequate. A great impetus to this study was provided by Erslev (1953) and Borsook, Graybiel, Keighley and Windsor (1954), who were able to demonstrate that the plasma of anaemic animals contained a humoral factor capable of stimulating erythropoiesis in normal recipients. This humoral factor, now usually known as 'erythropoietin', has, to date, eluded chemical isolation, but a considerable body of evidence has accumulated to indicate that it is a specific entity, chemically a mucoprotein, stimulating erythropoiesis in experimental animals when administered in extremely small quantities (Goldwasser, 1962). It affects the bone marrow principally by inducing differentiation of very immature marrow cells, possibly multipotential haemocytoblasts, to subsequent maturation through the erythroid series. It does not constitute a stimulus, at least primarily, to mitosis or to haemoglobin synthesis, these functions being governed, apparently, by processes within the normoblast itself, or by other factors originating within the marrow.

The extent to which erythropoietin regulates the day-to-day production of red cells in health is unknown. The hormone has been demonstrated
in normal rat (Reichlin and Harrington, 1960) and human plasma (Penington, 1962), but following transfusion of a normal human subject, slowing of erythropoiesis was found to precede the disappearance of erythropoietin from plasma, suggesting that factors other than the hormone must play some part in controlling normal bone marrow activity.

In patients with anaemia, Van Dyke, Layrisse, Lawrence, Garcia and Pollycove (1961) were able to detect the plasma hormone in iron deficiency when the haemoglobin had fallen below approximately 5 g%/o, raising the possibility that the erythropoietin system represents a ‘feedback’ mechanism brought into operation by profound anaemia rather than a system for the physiological regulation of erythropoiesis. However, employing a more sensitive assay procedure, an increase in titre of the hormone has been demonstrated in the plasma of a normal subject following venesection, when the haemoglobin level was greater than 12 g%/o (Penington, 1961) indicating the participation of the hormone in the red cell regeneration following moderate haemorrhage, and it may be concluded that the absence of the hormone from plasma in anaemic states would be of significance in the aetiology of anaemia. The validity of any such conclusion, however, depends on the limitations of sensitivity of the assay procedure employed. To date, in no form of anaemia has a subnormal titre of the hormone been demonstrated in plasma, and claims which have been made for the role of defective production of the hormone in the aetiology of anaemia have been based on less definitive evidence.

**Physiological Studies of Erythropoietin Production in Animals**

The site of production of the hormone in the body has been the subject of considerable controversy. Contopoulos, Ellis, Simpson, Lawrence and Evans (1954) demonstrated that the pituitary elaborated a factor capable of stimulating erythropoiesis in the hypophysectomized animal. However, hypophysectomized rats were found to respond to bleeding with increased erythropoiesis (Jacobson, Plizak, Fried, and Goldwasser, 1956) and the pituitary factor was subsequently shown to be identical with ACTH (Simpson, Evans and Rosenberg, 1959), operating via the adrenal to increase the rate of metabolism in the hypophysectomized animal, and hence to increase its requirement of red cells for oxygen transport. There is no convincing evidence to indicate that ACTH plays a physiological role in the regulation of erythropoiesis in the intact animal.

Stohlman, Rath and Rose (1954) studied a patient with regional hypoxia of the lower half of the body resulting from a patent ductus arteriosus with pulmonary hypertension and reversal of flow through the shunt. Polycythaemia had developed, despite normal oxygenation of the upper trunk, head and arms, suggesting that some organ or organs in the lower part of the body were capable of secreting the humoral factor leading to polycythaemia. Jacobson, Goldwasser and their collaborators (Jacobson, Goldwasser, Fried and Plizak, 1957; Goldwasser, Fried and Jacobson, 1958; Jacobson, Marks, Gaston and Goldwasser, 1959) found that removal of many of the endocrine glands and viscera in the rat did not deprive the animal of the capacity to produce erythropoietin in response to bleeding, but that, when the kidneys were removed, increased production of the hormone could not be detected with certainty in response to bleeding, hypoxia or cobalt administration. Comparable uraemia, induced by bilateral ligation of the ureters, did not impair hormone production. Mirand and Prentice (1957) failed to confirm that nephrectomy in the rat inhibited production of the hormone, but their assay of erythropoietin was carried out in hypophysectomized animals in which ACTH and corticoids (released in the donor animals in response to haemorrhage or anoxia) might be expected to increase erythropoiesis. Reissman, Nomura, Gunn and Brosius (1960) more recently confirmed that bilateral ligation of the ureters in the rat does not impair the normal erythropoietic response to anaemia, whereas bilateral nephrectomy or tubular necrosis induced with mercuric chloride do abolish this response. Injection of erythropoietin accelerates erythropoiesis in nephrectomized rats but to a lesser extent than in normal animals or in those in which the ureters have been ligated.

Whilst, in the rat, the kidney appears to be essential for the normal increase in hormone production in response to anaemia, there is now mounting evidence of species differences. In the dog, Naets (1958) observed disappearance of normoblasts from the marrow following bilateral nephrectomy with virtual cessation of erythropoiesis, whereas obstructive uraemia had little effect on bone marrow function. In the rabbit, on the other hand, Erslev (1958) found that both bilateral ligation of the ureters and nephrectomy abolished the increase in plasma erythropoietin following haemorrhage, and the uraemic rabbits showed impaired bone marrow response to administered erythropoietin whether or not functioning renal tissue remained. This would suggest that uraemia itself depresses erythropoiesis. Fischer and Friedelici (1961) made the important observation that normoblasts were preserved in the bone marrow of bilaterally nephrectomized rabbits maintained for considerable periods by peritoneal lavage. These findings strongly suggest sites of
production of erythropoietin outside the kidney. Kuratowska, Lewartowski and Michalak (1961) have demonstrated erythropoietin production by the isolated perfused rabbit kidney, but more recently Reissman (1962), also studying the rabbit, was able to demonstrate that both the isolated kidney and liver could produce an erythropoiesis-stimulating factor under conditions of hypoxia, provided careful attention was paid to the maintenance of a physiological pH in the perfusion system. The weight of evidence from such studies would, therefore, favour the view that the hormone may be derived from several organs although of these the kidney may be the most important.

Erythropoietic Activity of Tissue Extracts

Early attempts to demonstrate erythropoietic activity in extracts from normal tissues met with no success (Gordon, Pliero and Tannenbaum, 1955; Erslev, 1958; Rambach, Alt and Cooper, 1957). Naets (1960) found that homogenates of anemic dog kidney significantly stimulated erythropoiesis in a rat in which erythropoiesis was slowed by starvation, but as erythropoietin is excreted in the urine when the plasma titre is high, and may be reabsorbed by the tubules, such a demonstration of the hormone in the anemic kidney is of doubtful significance. Slight stimulation of erythropoiesis by homogenates of liver and spleen were attributed to the supply of nutrients with the material assayed. Rambach, Alt and Cooper (1961) recently found that injection of washed homogenates obtained from a wide variety of normal rat tissues led to stimulation of erythropoiesis in their assay animals (in this instance, rats with relative polycythemia induced by dehydration); it should be noted that injection of these insoluble tissue-homogenates produced considerable local inflammatory reaction in the assay animals, and the mechanism by which erythropoiesis was stimulated is far from clear. Recently, however, soluble protein preparations have been obtained from liver, spleen and kidney of normal rats and dogs, and from the kidneys of slaughterhouse pigs, which stimulate erythropoiesis in an assay animal in which red cell production has been slowed by transfusion (Penington, unpublished), and it appears likely that many tissues are capable either of elaborating erythropoietin, or stimulating its elaboration elsewhere within the body. Whether this apparently widespread tissue factor and the mucoprotein hormone of anemic plasma are identical has not been established; it must be borne in mind that the demonstration of erythropoietic activity in tissue extracts, particularly if associated with insoluble particulate matter as found by Rambach, cannot be accepted as evidence that each of these tissues is capable of influencing red cell production under physiological conditions. The tissue factor, however, whether the same or different from that of anemic plasma, may have great relevance to the occasional occurrence of polycythemia with primary tumours of a number of organs—kidney, ovary, uterus, liver, cerebellum and mesencephalic region. Further chemical studies of this tissue factor will be required to establish its relation to the physiological hormone.

Site of Production of Erythropoietin in Man

Studies of erythropoiesis in a human subject maintained by artificial dialysis following surgical extirpation of a sole remaining kidney showed that erythropoiesis may continue at something approaching a normal rate (Rees, Scheitlin, Giordano, Guild and Merrill, 1960). In this respect, therefore, man resembles the rabbit more closely than the dog or the rat, the kidney not being essential for the maintenance of normal erythroid activity. It would be of great interest to know whether the plasma of such a patient contains the physiological erythropoietic hormone, and whether such a patient could respond to a physiological stimulus to erythropoiesis. This would establish whether extrarenal sites of production of the hormone exist in man, or whether the erythroid marrow is capable of some degree of autonomous activity. Fortunately, opportunities for such studies are rare, but in so far as it is ever justifiable to infer patterns of human physiology from studies in experimental animals it appears likely that whilst a number of organs have the capacity to elaborate an erythropoiesis-stimulating hormone, under conditions of anaemia the kidney appears to be the most sensitive to the reduced oxygen carrying capacity of the blood. The reason for this primary role of the kidney lies, almost certainly, in the pattern of the renal vasculature and its response to anemia. Both Bradley and Bradley (1947) and Baldini and Lamperi (1956) found that in chronic anaemia the renal plasma flow is maintained at a normal rate, the total renal blood flow falling with the haematocrit, further reducing the number of red cells reaching the renal tissue.

The kidney would appear, therefore, to be the organ with the highest threshold to defective oxygen supply and its vasculature is such that the stimulus of anaemia is 'amplified' as the haemoglobin level falls, leading to the greatly increased outpuring of erythropoietin found at low haemoglobin levels.

Erythropoietin in Renal Anaemia

Assay methods used in the study of erythropoietin in anaemia have been extremely varied. The intact animal shows little increase in erythro-
poiesis following the administration of small doses of ‘anaemic’ plasma, and reticulocytosis, a sensitive but capricious index of erythropoiesis, must be used to detect the changes. Results obtained using this method have differed from those of other techniques (Linman and Bethell, 1956; Osnes, 1959; Clotten and Clotten, 1959), and judgment on such observations should be reserved until the variety of factors which appear to influence reticulocytosis are better understood. In many instances conclusions have been drawn from very slight elevation in reticulocyte counts, and much must depend upon the impartiality of the observer.

Techniques which lessen the endogenous stimulus to erythropoiesis in animals have been employed by many investigators to improve the sensitivity to administered erythropoietin. Here again, however, errors have been introduced. The most widely used procedures have been hypophysectomy or acute starvation, both of which markedly slow erythropoiesis, presumably reflecting reduced tissue requirements for oxygen (Fried, Pflzak, Jacobson and Goldwasser, 1957) but in each instance, hormones other than erythropoietin may well influence the assay findings. A more specific, and more sensitive method has been that in which animals—rats or mice—have been rendered polycythaemic by transfusion (Fried and others, 1957; Gallagher, McCarthy and Lange, 1960; Penington, 1961) or by exposure to low oxygen tension (Cotes and Bangham, 1961) following which the plethoric animals lack any physiological stimulus to erythropoiesis. Radioactive iron has provided the most widely used method of measuring the rate of erythropoiesis in the animals following one or two injections of the material for assay. Quantitative assessment of the hormone in plasma remains crude, however, and conclusions concerning defective production of the hormone in disease have been based on a comparison of findings in anaemia due to different causes. There is now a growing body of evidence to indicate that in the anaemia of renal failure the hormone titre is elevated to an extent considerably less than that observed in other types of anaemia of comparable severity (Gurney, 1957; Gallagher and others, 1960; Penington, 1961). Whilst it would be tempting to conclude that lack of erythropoietin is the sole cause of defective erythropoiesis in renal failure, further studies do not confirm this view. With chemical concentration of the hormone from plasma, some increase in titre is found in renal anaemia when compared with similar concentrates of normal plasma (Penington, 1961).

It may be concluded, therefore, that in chronic renal disease several factors, including hemolysis, and most probably toxic marrow depression conspire to produce anaemia, but that the normal increase in erythropoietin titre in response to anaemia is lacking. The relative importance of these various factors in the aetiology of the anaemia must await the availability of the hormone for clinical trials.

Nature of the Tissue Receptor for Erythropoietin Release—A Hypothesis

The arterial tree contains specialized tissues such as the carotid body sensitive to anoxia of arterial blood, but secretion of erythropoietin occurs with a fall in oxygen content of blood, for in anaemia the arterial oxygen tension is normal. Whilst reduced oxygen tension in arterial blood also constitutes a stimulus to erythropoietin secretion, the common factor to anaemia and hypoxia would appear to be reduced oxygen supply to some sensitive tissue, the final common pathway being lowered tissue oxygen tension.

Osnes (1958), from studies in mice, suggested that renal juxtaglomerular granules might represent the site of production of the hormone, these granules increasing in number with haemorrhage and disappearing following transfusion. However, these experimental procedures would be expected to cause great alterations in juxtaglomerular granulation accompanying altered secretion of renin (known to be derived from this tissue). Furthermore, these cells, situated as they are in the afferent and efferent arterioles of the glomerulus, are bathed by arterial blood, in which oxygen tension is normal in anaemia. It would appear unlikely, therefore, that they would be capable of ‘recognizing’ anaemia in the arterial blood.

The vascular structure of the nephron does, however, provide a mechanism whereby the body can readily translate ‘oxygen content of blood’ to ‘altered tissue tension’. After leaving the glomerulus, blood passes along the course of the proximal convoluted tubule and loop of Henle to the distal tubule, oxygen being extracted to an extent dependent on the rate of metabolism of the tubular cells. If, when blood reaches the distal tubule, the oxygen content has been depleted below a critical level, production or release of the hormone would be initiated. The latent interval of some three to six hours which has been observed between the exposure of an animal to hypoxia and the appearance of the erythropoietin in plasma would suggest that the response to the stimulus is not simply one of release of preformed hormone. The steps involved in the release of the hormone are of considerable interest, but remain to be clarified. Papain, a plant enzyme similar in substrate specificity to several tissue hydrolytic enzymes (cathepsins) and like them activated by

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reducing conditions, is known to stimulate erythropoiesis when injected into rats (Yen, Cheng-Chun Lee and Flemming, 1960) although it destroys erythropoietin in vitro. The step linking tissue hypoxia with liberation of the hormone may well be provided by such an enzyme, activated by reducing conditions in the tubular cells, converting a hormone precursor to an active form. Preliminary studies using tissue preparations suggest that such a step is involved. The localization of this humoral function in the nephron remains unknown. It may be a property of the tubule as a whole, or of a specialized region of the distal tubule such as the macula densa, or the granular ‘intercalated cells’ proximal to the collecting tubule. Perhaps the latter would be the most favourably situated to function as a regulator of erythropoiesis.

The hypothesis outlined above has a number of interesting corollaries. Removal of oxygen from blood perfusing the nephron will be governed by the metabolic requirements of the tubular cells. Increased oxygen extraction by stimulation of cell metabolism should constitute a stimulus to erythropoietin secretion, as shown by Fried and others (1957) using dinitrophenol, an agent known to uncouple oxidative phosphorylation in the tissues. Reduction of metabolic rate should be accompanied by reduced hormone secretion and it has been suggested by Jacobson and others (1959) that the reduction in erythropoiesis following acute starvation or in myxœdema reflects such a change. The anaemia of myxœdema in man, however, represents a more complex disturbance (Penington, unpublished).

Factors influencing the rate of perfusion of the nephron should greatly affect erythropoietin secretion. Thus, as nephrons are obliterated by disease, an increased rate of perfusion of remaining nephrons would lead to a constant ‘overestimation’ of the oxygen content of the blood, and a ‘resetting’ of the homeostatic level for erythropoiesis at a point well below the normal haemoglobin concentration. The erythropoietic defect in chronic renal disease might, therefore, be one of an altered threshold for erythropoietin secretion, rather than an inability of the kidney to elaborate the hormone. That such is the case is suggested by the observation of Verel and others (1959) that following haemorrhage in a patient with a chronic renal anaemia, regeneration of the red cell mass occurred at a normal rate, but ceased on reaching the original anaemic level.

A further consequence of the hypothesis would be that reduced perfusion of nephrons would constitute a stimulus to erythropoietin production. A fall in blood flow due to increased intrarenal tension in hydronephrosis or in the presence of cysts could account for the unusual phenomenon of pure red cell polycythaemia which may occur with these conditions. The reduced rate of perfusion would lead to a constant underestimation of the oxygen content of the blood, hence constituting a stimulus to erythropoietin release as in anaemia.

In polycythaemia vera, where the marrow proliferation is primary, subnormal secretion of the hormone would be anticipated. This has now been observed in some instances (Penington, unpublished) but in others, normal or slightly elevated erythropoietin titres may occur when the hematocrit is particularly high. This presumably reflects inadequate perfusion of nephrons consequent on greatly increased viscosity of the blood. Lesions such as hydronephrosis or cysts might aggravate the situation in polycythaemia thus establishing a vicious circle of events, increased viscosity of blood providing a stimulus to further erythropoiesis. It is of interest that in a case of polycythaemia associated with polycystic kidneys, Gurney (1960) detected erythropoietin in fluid from one of the cysts. The polycythaemia improved following nephrectomy, but subsequently relapsed, at which time splenomegaly and elevated leucocyte and platelet counts—hallmarks of polycythaemia vera—were noted. In a number of such cases of polycythaemia attributed to renal lesions, splenomegaly or other features of polycythaemia vera have been recorded (Lawrence and Donald, 1959), and it would appear likely that in certain cases the renal lesion aggravates coexistent primary polycythaemia, rather than itself initiating marrow proliferation. In other instances, however, the renal lesion does appear to be the primary cause, as permanent remission has followed surgical removal of the kidney. The reader will find the subject reviewed elsewhere (Gardner and Freyman, 1958; Lawrence and Donald, 1959; Gurney, 1960; Jones, Payne, Hyde and Price, 1960).

**Studies on Polycythaemia Accompanying Renal Lesions**

Through the kindness of a number of associates, a group of patients has been studied with renal lesions accompanied by polycythaemia. The details of the assay procedure, using rats with sustained transfusion polycythaemia, will be found elsewhere (Penington, 1961). The utilization of radioactive iron at 24 hours was 2.8% of the injected dose in control animals (mean of 25 values with a standard deviation of ± 0.84). The result of each assay is expressed in terms of multiples of the standard deviation between the mean control value and the mean iron utilization of the assay animals. Plasma samples were concentrated by a procedure previously reported (Penington, 1961),
when available in sufficient quantity, and the extract from 12 ml. of plasma was injected per assay animal. Alternatively, a total of 4 ml. of unmodified plasma was assayed per rat. Normal human plasma or plasma extract under these conditions does not significantly stimulate iron utilization in the assay animals. The statistical significance of each assay result is given in terms of the 'P' value calculated by the method for small samples.

**Case 1—Renal Carcinoma and Red Cell Polycythæmia**

Male, aged 44. Complained of headache and palpitation. Found to have mass in the left loin presumed to be the spleen, a haemoglobin level of 17.5 g. %, haematocrit 51.5%, and WBC 11,200/cu. mm. A diagnosis of polycythæmia vera was made. Twelve months later the patient was referred to Queen Mary's Hospital, Roehampton, where the polycythæmia was found to be associated with a large calcifying carcinoma of the left kidney. At operation the tumour was removed, and the spleen was noted to be normal in size. The patient subsequently became anaemic and deteriorated steadily. Evidence of pulmonary metastases was found and the patient died six weeks after surgery.

**Erythropoietin Studies**

Plasma prior to operation (unconcentrated) + 8.5

(P < 0.01).

Tumour extract (extract from 8 g. per rat) + 6.8

(P < 0.01).

**Comment**

Both the patient's plasma and the tumour contained highly significant erythropoietic activity. The tumour was heat extracted after homogenization with acetate buffer, pH 5.5, followed by removal of lipids with ether, dialysis and lyophilization. The active material from the tumour is concentrated chemically by the same procedures as the plasma hormone. It is likely that the tumour was itself secreting erythropoietin, which was responsible for the patient's polycythæmia.

**Case 2—Hydronephrosis and Red Cell Polycythæmia**

Male, aged 53, was found to have a haemoglobin of 21 g. % on routine examination. When seen at the Radcliffe Infirmary, Oxford, he had an enormous mass occupying the left side of the abdomen. The white cell and platelet counts were normal. At operation a very large hydronephrotic kidney was removed and found to be caused by a calculus impacted in the ureter. Three months after operation there was no evidence of recurrence of the polycythæmia.

**Erythropoietin Studies**

Plasma—pre-operative (total dose

2.5 ml.) . . . . . + 2.6 (P < 0.02)

post-operative (total dose

4 ml.) . . . . . + 1.9 (P = 0.05)

Cyst fluid (total dose 6 ml.) . . . + 0.5

**Comment**

Despite the small quantity of plasma available for assay prior to operation, significant erythropoietic activity could be detected. A second sample obtained three months after surgery showed a probable slight in-

crease in hormone titre above normal; the haemoglobin had fallen to 11 g. % at this time, and such activity of the plasma would be anticipated, in response to this mild anæmia.

**Case 3—Renal Cyst and Red Cell Polycythæmia**

Male, aged 56, admitted to the Edinburgh Royal Infirmary with headache and ataxia, and was found to have a mild polycythæmia confined to the red cell series. The red cell mass was found to be 37.5 ml./Kg. body weight. The spleen was not palpable. An intravenous pyelogram demonstrated a large swelling within the left kidney which at operation proved to be a cyst. The day following operation, signs of vascular damage to the brain stem progressed rapidly, and the patient died. Samples of plasma were obtained from a peripheral vein the evening before surgery, and from the left renal vein during the operation.

**Erythropoietin Studies**

Mixed venous blood (extract from

12 ml. . . . . . + 4.1 (P < 0.01)

Renal vein blood (extract from

12 ml.) . . . . . + 2.5 (P < 0.05)

**Comment**

This patient showed erythrocytosis without other features of polycythæmia vera. Plasma obtained before operation showed significant erythropoietic activity, considerably greater than has been found in any subject with polycythæmia vera. Renal vein blood was less active on assay but as it was obtained whilst the patient was under general anaesthesia the two samples are not strictly comparable. The likelihood is that in such cases the hormone is elaborated in increased quantities by the cystic kidney.

**Case 4—Polycystic Kidneys and Papilloma of the Cæcum with Red Cell Polycythæmia**

Male, aged 53. Attended the Postgraduate Medical School, Hammersmith, complaining of recurrent abdominal pain of three years' duration. Noted to be pale but to have a normal blood pressure. The haemoglobin was 18.0 g. %, haematocrit 51.5% and white cell and platelet counts normal. An intravenous pyelogram was carried out because of the erythrocytosis, and polycystic kidneys were demonstrated. The abdominal pain remained unexplained.

Two years later the patient again attended hospital with abdominal pain. The haematocrit was now 56%; white cells and platelets again normal. The red cell mass was found to be 38.2 ml./Kg. body weight and plasma iron turnover was found to be accelerated. The patient was venesected to a total of 1,120 ml. over four days and erythropoietic activity of these samples was determined.

Further investigation revealed that the patient was suffering from a papilliferous tumour of the cæcum which was not evident from previous radiological studies. The tumour was successfully removed and eight months after operation the peripheral blood picture was found to have remained normal in all respects.

**Erythropoietin Studies**

Plasma (unconcentrated)—initial

phlebotomy . . . . . + 3.8 (P < 0.01)

48 hours after 500 ml. phlebotomy . . . . . + 3.8 (P < 0.01)

48 hours after 250 ml. phlebotomy . . . . . + 1.9 (P > 0.05)
Comment

Significantly elevated plasma erythropoietic activity was found prior to phlebotomy but once the haemoglobin level had been restored to normal, the activity appeared to lessen. The significance of this change is difficult to evaluate. In a normal subject the plasma erythropoietin titre rises after venesection and we have raised it in patients with this disease in the absence of renal lesions. The fall in hormone titre following venesection is an evidence of the relatively lower erythropoietin level being produced to maintain the normal haemoglobin level and hence perpetuate the polycythaemia.

Case 5—Hydronephrosis with Polycythemia Vera

Male, aged 36. Two years before admission developed a painful and discoloured small toe. For six months, troubled by shortness of breath and palpitations on exertion and for two months by a nausea and dull frontal headaches. When admitted to the Radcliffe Infirmary, Oxford, a blood pressure of 150/115 mm. Hg. was noted, together with discoloration of two toes and a palpable spleen. The haemoglobin was 22.0 g. %, w.b.c. 10,400 cu. mm., and platelets 600,000 and 1,000,000 on two estimations. An intravenous pyelogram was carried out which showed the right kidney to be enlarged by a space-occupying lesion at the upper pole. This was subsequently demonstrated to be a renal cyst, fluid being aspirated by needle puncture and replaced by radio-opaque dye. The patient was venesection repeatedly until his haemoglobin and haematocrit were restored to normal. However, four months later the polycythaemia had recurred and treatment with radioactive phosphorus was required.

Erythropoietin Studies

Plasma prior to venesection (dose 4 ml.) . . . . + 1.7 (P = 0.05)

Plasma following venesection (dose 4 ml.) . . . . + 1.2 (P > 0.05)

Cyst fluid (total dose 8 ml.) . . . . + 2.0 (P < 0.05)

Comment

The plasma erythropoietic activity prior to phlebotomy did just reach a detectable level, but following restoration of the peripheral blood picture to normal, the increase in activity could not be detected. The cyst fluid also showed a barely detectable elevation of erythropoietic activity. The patient was clearly suffering from polycythaemia vera, and similar slight elevation of humoral activity in the plasma has now been detected in several patients with this disease in the absence of renal lesions. The fall in hormone titre following venesection is an evidence of the lower erythropoietin level being produced to maintain the normal haemoglobin level and hence perpetuate the polycythaemia.

Summary

The anaemia of renal failure cannot be ascribed to a single cause. Shortened red cell life span is found under some circumstances and defective red cell production, the principal factor, may itself be due to toxic depression of erythropoiesis, in addition to inadequate humoral stimulation of the bone marrow. The kidney plays the major role in elaboration of erythropoietin in anemia, although it is not the only site of production of the hormone in the body. A mechanism by which erythropoietin secretion may be related in the kidney to the circulating haemoglobin level has been outlined and the consequences of this hypothesis in the interpretation of renal polycythaemia has been discussed.

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