Chorea, polycythaemia, and cyanotic heart disease

P. D. EDWARDS, R. PROSSER, AND C. E. C. WELLS

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SYNOPSIS Two cases of polycythaemic chorea are described, both of which were complicated by severe heart disease. The first was a child with patent ductus arteriosus and coarctation of the aorta causing severe cyanosis and secondary polycythaemia. Chorea began intermittently at an early age, becoming continuous by his fifth birthday. The second was a middle-aged male with tight mitral stenosis and a story of paralytic chorea in his teens. Polycythaemia rubra vera was eventually diagnosed two years after mitral valvotomy, some seven years after the onset of chorea.

1. Chorea, polycythaemia, and morbus coeruleus

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Congenital heart disease with a right-to-left shunt is the most common cause of polycythaemia in children. The vital signs are stunted growth, central cyanosis with bloodshot eyes and crimson lips, and clubbing. The number of red blood cells, the haematocrit and haemoglobin concentration are all elevated and the oxygen saturation of arterial blood is decreased. The haemoglobin and haematocrit levels, which may exceed 16 g/dl and 55% respectively, then correspond to a red cell mass greater than 35 ml/kg.

The natural history of polycythaemia is often studded with neurological episodes: transient ischaemic attacks or a frank stroke are typical presenting symptoms. The complications of polycythaemia rubra vera, reviewed in full by d'Eramo and Levi (1972), do not differ from those of polycythaemia which is secondary to hypoxia or to increased production of erythropoietin. Chorea is a rare manifestation. In the past 65 years it has been recorded in little more than a score of cases, mostly middle-aged or elderly females (Table 1), although the incidence of polycythaemia vera is greater in men (Wintrobe, 1967).

Chorea associated with secondary polycythaemia has been reported in a child. Polani and MacKeith (1954) described a 4 year old girl with

(Accepted 26 February 1975.)
TABLE 1

CASES OF POLYCYTHEAEMIA AND CHOREA PREVIOUSLY REPORTED IN JOURNALS CIRCULATING IN GREAT BRITAIN

<table>
<thead>
<tr>
<th>Date</th>
<th>Authors</th>
<th>Sex Age (yr)</th>
<th>History, treatment, progress</th>
<th>F.U.</th>
</tr>
</thead>
<tbody>
<tr>
<td>1909</td>
<td>Bardachti (case 1)</td>
<td>F 59</td>
<td>Abrupt onset of chorea, beginning in R hand, chiefly of face, tongue, limbs, with dysarthria and ataxia. Severe headache. Splenic infarct, skin haemorrhages, boils, carbuncle of sacrum. Hb 135%, RBC 14-10^12, WBC 27-10^9/l, splenomegaly. Remission of chorea, clearing of sepsis after sodium iodide therapy. Hb 115%, RBC 8.56-10^12/l, splenomegaly.</td>
<td>5 m</td>
</tr>
<tr>
<td>1909</td>
<td>Umney</td>
<td>F 34</td>
<td>For 5 yr increasingly livid complexion; then abortion, menorrhagia, persistent albuminuria. Hb 138%, RBC 11.5-10^12, WBC 17-10^9/l, splenomegaly. After 2 yr speech indistinct, onset of chorea and thrombosis of left innominate vein. Chorea subsided with phenazone therapy. Within 3 m generalized venous thrombosis, death in coma. No necropsy</td>
<td>3 m</td>
</tr>
<tr>
<td>1919</td>
<td>Naville and Brütsch (case 1)</td>
<td>F 33</td>
<td>Relapsing chorea aged 19-22 yr. Abrupt loss of consciousness for 3 d and R hemiplegia at 23 yr. Onset of epilepsy at 31 yr. Falling vision with papilloedema, consecutive optic atrophy at 33 yr. Hb 100%, RBC 6.758-10^12, WBC 16.43-10^9/l splenomegaly. Laparotomy for obstruction: splenic 'tumor' not excised.</td>
<td>14 yr</td>
</tr>
<tr>
<td>1922</td>
<td>Pollock</td>
<td>F 38</td>
<td>For 6 m attacks of headache, dizziness, cyanosis, dyspnoea; relieved by radiotherapy. 3 w, speech indistinct, grimacing, involuntary movements of jaws, tongue, limbs. Hb 115%, RBC 8.1-10^12, WBC 8.5-10^9/l, splenomegaly. Remission of chorea after venesection and radiotherapy. RBC 6.4-10^9/l, splenomegaly.</td>
<td>3 m</td>
</tr>
<tr>
<td>1933</td>
<td>Schiff and Simon</td>
<td></td>
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</tr>
<tr>
<td>1936</td>
<td>Schiff et al.</td>
<td>F 78</td>
<td>Transient flaccid tetraparesis at 73 yr. Over next 5 yr increasing lividity. Hb 90%, RBC 6.8-10^12, WBC 16-10^9/l, 6 w later, sudden onset of chorea of face, trunk, limbs, with confusion, disorientation, drowsiness. Ataxia and scanning speech. Xanthochromic CSF. Death within months P.M.: congestion of cerebral veins with some thromboses, demyelination of anterior globus pallidus, degeneration of hypothalamic nuclei, and loss of Purkinje cells from cerebellar cortex.</td>
<td>2 m</td>
</tr>
<tr>
<td>1940</td>
<td>Dameshek and Henstell (case 7)</td>
<td>M 39</td>
<td>For 1 yr inappropriate sleep, loss of concentration, irritability. Recent dizziness, nausea. Involuntary movements of arms, legs. Hb 150%, RBC 8.7-10^12, WBC 12-10^9, platelets 800-10^9/l, PCV 80%, splenomegaly. Symptomatic relief with low iron diet.</td>
<td></td>
</tr>
<tr>
<td>1940</td>
<td>Dameshek and Henstell (case 8)</td>
<td>F 49</td>
<td>Twitching of fingers, orofacial dyskinesia began 2 yr after complaint of headaches, hot flushes, vertigo, tinnitus. Superficial venous thrombosis. Hb 118%, RBC 10.2-10^12, WBC 10.8-10^9, platelets 2.5-10^12/l, splenomegaly. Symptoms subsided after venesection and treatment with arsenic, phenylhydrazine, low iron diet.</td>
<td></td>
</tr>
<tr>
<td>1942</td>
<td>Kotner and Tritt</td>
<td>F 64</td>
<td>For 5 yr symptomless polycythaemia: Hb 130%, RBC 9.4-10^12, WBC 12.7-10^9/l. Uler of mouth after dental extraction and chorea involving face, mouth, trunk, limbs. Explosive speech. Hb 130%, RBC 9.5-10^12, WBC 16.5-10^9/l, PCV 76%, splenomegaly. Xanthochromic CSF. Despite venesection, rapid deterioration and death on 9th day P.M.: pulmonary, mesenteric thrombi; congestion of all cerebral veins with multiple thrombi, choroidal haemorrhage of 4th ventricle; periventricular demyelination. No areas of necrosis or cell loss.</td>
<td>7 w</td>
</tr>
<tr>
<td>1946</td>
<td>Reinhard et al. (case 19)</td>
<td>F 56</td>
<td>Nervous for many years, dizziness and nausea for 1 yr, chorea and dysarthria for 1 m. Mother had polycythaemia. Chorea improved after venesection and before radiophosphorus therapy.</td>
<td>1 m</td>
</tr>
<tr>
<td>1950</td>
<td>Harvier et al.</td>
<td>M 49</td>
<td>For 7 yr 'writer's cramp', mild dysarthria, Bleeding gums. 6 yr, joint pains followed by onset of chorea beginning in lips and tongue. Hb 128%, RBC 7.0-10^12/l splenomegaly. Therapy with Boudin's solution. Chorea fluctuated, tending to worsen. Hb 150%, RBC 8.3-10^12, WBC 28.4-10^9/l. Further therapy with Boudin's solution, phenylhydrazine, and radiotherapy. Gradual remission of chorea. Final Hb 137%, RBC 6.2-10^12, WBC 18.2-10^9/l.</td>
<td>8½ yr</td>
</tr>
<tr>
<td>1952</td>
<td>Alajouanine et al. (case 4)</td>
<td>F 65</td>
<td>Livid complexion for 4 yr. Rapid onset of involuntary movements of face, arms, trunk, together with Parkinsonian tremor of hands. RBC 9.9-10^12/l, splenomegaly. Remission followed venesection and radiophosphorus therapy.</td>
<td>3½ yr</td>
</tr>
<tr>
<td>1954</td>
<td>Trotsemburg and Koster</td>
<td>F 63</td>
<td>For 18 m involuntary movements chiefly of right side with orolinguual dyskinesia, dysarthria, and right hemiparesis. Hb 140%, RBC 7.8-10^12, WBC 7.0-10^9/l, platelets 700-10^9/l, PCV 61%, splenomegaly. After radiophosphorus therapy, remission of chorea apparent 1 m before fall of haemoglobin and red cell count</td>
<td>3 yr</td>
</tr>
<tr>
<td>Date</td>
<td>Authors</td>
<td>Sex</td>
<td>Age (yr)</td>
<td>History, treatment, progress</td>
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<tr>
<td>1958</td>
<td>Paliard et al. (case 1)</td>
<td>M</td>
<td>73</td>
<td>After disappointment 9 m before, rapid onset of abnormal movements of arms and of slurred speech. RBC 7.8·10¹²/l, splenomegaly. Remission within few months of radiophosphorus therapy. Relapse after 4 yr, involving all four limbs, face, tongue, with rhythmic myoclonus of pharynx and larynx. RBC 4.8·10¹²/l, and no splenomegaly. Remission with chlorpromazine. Severe relapse after 5 m; Hb 140% RBC 7.45·10¹²/l, PCV 77%. Gradual remission after radiophosphorus. Final Hb 80%, RBC 4.0·10¹²/l, slight splenomegaly</td>
</tr>
<tr>
<td>1959</td>
<td>Calabresi and Meyer</td>
<td></td>
<td></td>
<td>'One case had choreoathetoid movements with speech impairment...'</td>
</tr>
<tr>
<td>1962</td>
<td>Bogolepov</td>
<td>M</td>
<td>70</td>
<td>Polycythaemic for 10 yr. 4 yr ago nocturnal epileptic fit followed by chorea with orofacial predominance (illustrated). Remission after venesection and radiophosphorus therapy, 3 yr later, further relapse, trophic lesions of several fingers. Further remission with therapy. Final Hb 4.62·10¹², platelets 254·10⁹/l</td>
</tr>
<tr>
<td>1963</td>
<td>Bakke and Stavem</td>
<td>F</td>
<td>74</td>
<td>After complaint of dizziness, developed involuntary movements of hand, arm, becoming generalized within 3 m. Staccato speech and abnormal gait. Hb 18.4%, RBC 7.7·10¹², WBC 18.4·10⁹, platelets 250·10⁹/l, splenomegaly. Little change after venesection</td>
</tr>
<tr>
<td>1965</td>
<td>Friedemann et al.</td>
<td>F</td>
<td>62</td>
<td>One year after chance finding of raised haemoglobin, onset of malaise, pain in mouth and tongue, and of involuntary movements beginning in tongue and rapidly becoming generalized. Hb 18.5%, RBC 6.5·10¹²/l, WBC and platelets not increased. Total red cell volume 43.6 ml/kg. Within 2 m of venesection, radiophosphorus therapy, chorea had disappeared, remission preceding significant fall of blood count</td>
</tr>
<tr>
<td>1967</td>
<td>Gautier-Smith and Pranker (case 1)</td>
<td>F</td>
<td>57</td>
<td>Sydenham's chorea at age 12 yr. Onset of headaches, drop attacks, vertigo, amблиопіа at 53 yr followed after 4 yr by increasing fidgetiness. Chorea chiefly of face, tongue, and upper limbs, jerky respirations, grunting speech. Hb 22.2 g/dl, RBC 8.02·10¹², WBC 18.1·10⁹, platelets 450·10⁹/l, PCV 77%, splenomegaly. Red cell mass 81.1 ml/kg. After venesection and radiophosphorus therapy, remission of chorea preceded that of polycythaemia. Temporary relapse of chorea—RBC 8.92·10¹²/l—responding well to further venesection and radiophosphorus. Final Hb 16.0 g/dl, PCV 54%</td>
</tr>
<tr>
<td>1967</td>
<td>Gautier-Smith and Pranker (case 2)</td>
<td>F</td>
<td>74</td>
<td>Sudden onset dysphagia and dysarthria, confusion and chorea, deteriorating after 5 m. Hb 23.0 g/dl, WBC 8.3·10⁹, platelets 650·10⁹/l, PCV 73%. No splenomegaly. Red cell mass 65 ml/kg. Remission of chorea and general improvement after radiophosphorus therapy. Final Hb 14.0 g/dl, WBC 6.5·10⁹, platelets 143·10⁹/l</td>
</tr>
<tr>
<td>1968</td>
<td>Bietti et al.</td>
<td>F</td>
<td>75</td>
<td>Polycythaemia diagnosed at age 56 yr and treated with radiotherapy. Further therapy at 72 yr for sciatica and at 74 yr after haemorrhages into skin. Regular venesection. After a short lapse of therapy, onset of involuntary movements of left hand and episodes of unconsciousness. RBC 8.0·10¹²/l. Progression of chorea to whole of right side and explosive speech. Hb 17.3 g/dl, RBC 7.8·10¹², WBC 14.5·10⁹, platelets 360·10⁹/l. Died 1 m after radiophosphorus therapy. Final RBC 6.64·10¹², platelets 320·10⁹/l. P.M.: distended intracranial veins with thrombi of smaller vessels and haemorrhage of L putamen; areas of perivascular demyelination; large subdural haematoma</td>
</tr>
<tr>
<td>1968</td>
<td>Heathfield</td>
<td>F</td>
<td>77</td>
<td>Six month story of chorea involving face, tongue, limbs. Hb 18.5 g/dl, PCV 66%; 1 m later RBC 8.0·10¹², WBC 8.8·10⁹, platelets 320·10⁹/l, PCV 64%, no splenomegaly. Chorea controlled by venesection and thioridazine therapy</td>
</tr>
<tr>
<td>1969</td>
<td>Nehil et al.</td>
<td>F</td>
<td>70</td>
<td>Onset of generalized chorea involving face, tongue, limbs at age 68 yr. RBC 7.03·10¹²/l, increased WBC, PCV 70%, no splenomegaly. Remission after venesection. Relapse after 2 yr with marked orofacial dyskinesia and dysarthria. Further remission after therapy with cytotoxic drug and haloperidol</td>
</tr>
<tr>
<td>1970</td>
<td>Sangster</td>
<td>F</td>
<td>71</td>
<td>Six weeks after onset of giddy turns, spontaneous jerking movements of face and limbs with dysarthria. Became rapidly helpless. Hb 24.9 g/dl, RBC 8.63·10¹², WBC 15.6·10⁹, platelets 120·10⁹/l. Total red cell volume 68 ml/kg. No splenomegaly. Immediate improvement after venesection and radiophosphorus therapy. Temporary relapse with worry. Final RBC 5.5·10¹², platelets 100·10⁹/l</td>
</tr>
<tr>
<td>1972</td>
<td>Ashenhurst</td>
<td>M</td>
<td>68</td>
<td>Onset of involuntary movements 7 yr after diagnosis of polycythaemia rubra vera (Hb 21.2 g/dl, WBC 6.0·10⁹, platelets 500·10⁹/l, PCV 65.5%,) which was treated by venesection. Recurrence of polycythaemia after 5 yr treated by venesection and radiophosphorus, last dose being given 2 w before onset of chorea when Hb 18.0 g/dl. Chorea settled within 10 d of further venesection. Final Hb 12.7 g/dl, RBC 4.7·10¹²/l</td>
</tr>
</tbody>
</table>

F.U. = follow-up from onset of chorea; yr = year, m = month, w = week
Fallot's tetralogy who became anoxic and hemiplegic during angiocardiography and, eight days later, developed choreoathetosis which persisted over the next two years. The abnormal movements were probably due to infarction of the upper brainstem.

In the following account of a boy with severe cyanotic heart disease and secondary polycythaemia, chorea and cerebral thrombosis were major complications. On two occasions his chorea subsided after substantial venesection but his hemiplegia persisted.

CASE 1

M.P. was born normally at term on 24 November 1965 after an uneventful gestation. He weighed 2867 g. At the end of the first week his left arm, both legs, and the lower part of his body were seen to be deeply cyanosed whenever he cried. A soft systolic murmur was audible over the precordium. The femoral pulses were palpable but were weak. The systolic blood pressure was 130 mmHg in his right arm and 150 mmHg in his left. The clinical diagnosis of preductal coarctation of the aorta and patent ductus arteriosus was supported by catheter studies (Dr. L. G. Davies).

Respiratory infections were frequent and led to many admissions to hospital. Because of the risk of congestive heart failure, digoxin was started. When the child was 3 years old, Dr Davies repeated the catheter studies and confirmed his earlier findings. He also reported severe pulmonary hypertension with a right-to-left shunt through the patent ductus. In the right brachial artery oxygen saturation reached 95% but in the abdominal aorta it fell to 60%. The severity of his pulmonary hypertension excluded him from surgery.

Chorea was first seen in the paediatric outpatient clinic in 1970 when he was nearly 5 years old. With prompting, his mother then recalled having noticed occasional spells of involuntary movements since his first birthday. They were present only when he was awake, lasting a few hours at a time, and subsiding as soon as he fell asleep. They had been continuous for two months: his face would screw up, his tongue dart in and out, his grip become weak and clumsy, and he would drop things unexpectedly. More recently, his walking and balance had been affected and during the past fortnight he had been off his feet.

He was small for his age and deeply cyanosed. Finger clubbing, like the degree of cyanosis, was more intense in his left hand than in his right and was marched by similar changes in his feet. Chorea was generalized and his speech was slurred, his words being either explosive or distorted by frequent grimacing. His limbs were hypotonic, the knee jerks pendular and 'hung up', the plantar responses flexor. Although the retinal veins were congested, the optic discs were flat. The cardiovascular signs were unchanged and both liver and spleen were easily palpable, the former extending four fingers' breadth below the costal margin. All peripheral pulses were present. Serial blood counts over the previous four years revealed increasing polycythaemia (Fig. 1) and a skull radiograph showed abnormal density of bone in vault and base (Fig. 2).

After repeated venesection—a total of 820 ml blood being withdrawn in a fortnight—he was able to walk again and his chorea subsided. The haemoglobin and haematocrit had then fallen to levels just above the normal range (Fig. 1). Within a month of
leaving hospital he was readmitted with another chest infection, during the course of which he developed right-sided weakness. Despite anticoagulant therapy, this progressed over the next four days to become a dense hemiplegia. His blood count had not altered. During the next two years further respiratory infections, sometimes associated with cardiac failure, responded to conventional therapy with digoxin, frusemide, and antibiotics. Anticoagulant therapy with warfarin was continued. At the beginning of 1973, when he was aged 7 years, his chorea relapsed but again subsided after a second series of venesections. The initial haematocrit reading of 66% fell to normal levels within a month of withdrawing 750 ml blood (Fig. 1). His hemiplegia, however, remained dense.

2. Chorea, polycythaemia, and mitral stenosis

C. E. C. Wells

Discussing the differences between polycythaemia rubra vera (erythraemia), a disease of unknown aetiology, and polycythaemia in response to a known stimulus (erythrocytosis), Wintrobe (1967) admitted that the distinction between the two was not always clear and that the expected splenomegaly, leucocytosis, and thrombocytosis of polycythaemia vera might be missing from an otherwise typical case. Glass and Wasserman (1972) listed 13 criteria by which they classified primary, secondary, and spurious types of polycythaemia but their table revealed the same overlap which Wintrobe (1967) had found.

The following case history describes a middle-aged man who presented with chorea and was found to have tight mitral stenosis. Respiratory function was also impaired. A mild degree of polycythaemia, affecting only the red cell series, was for many years attributed to his considerable hypoxia. When polycythaemia vera was diagnosed and treated accordingly, control of his chorea became simpler but complete remission was not achieved. Permanent structural injury may have resulted from the combined effects of venous congestion and long-continued therapy with phenothiazines grafted onto a striatum already weakened by Sydenham's chorea in adolescence.

CASE 2

In October 1959 a 45 year old man was brought to casualty with a crush injury of his left forearm. The skin and soft tissue were intact, the circulation normal, and no signs of bone injury were found. He soon returned to his work as a machine operator but became depressed and continued to complain about his arm. Perphenazine was prescribed and he stayed at work with occasional breaks for the next two years. In November 1961 a series of bereavements led to relapse of his depression and to the onset of chorea. He was admitted to a psychiatric hospital where change of therapy first to chlorpromazine and then to trifluoperazine, with benzhexol, seemed to control the movements and the depression. Later, Huntington's chorea was suspected and he was referred for neurological opinion.

His wife reaffirmed his story but quickly dispelled doubts about an inherited disorder. She had known his parents and grandparents as well as his three sisters and none had had chorea or been demented. His eldest sister had had rheumatic fever in her teens and was subject to fainting turns. Of their own three children, the two boys were well but their daughter had had varicella encephalitis, diagnosed by an experienced paediatric neurologist, and was left with a speech defect. She attended a school for handicapped children and had been examined in the neurological clinic when the family first moved to Wales from London.

Additional points from the past history were confirmed by the family doctor's records. Paralytic chorea of his right leg had been diagnosed in November 1932 and between that date and July 1933, when he had resumed work as a baker's roundsman, involuntary movements of his face and limbs had been recorded. During war service in West Africa he had had malaria but had otherwise remained well until his accident in 1959 to which all his present ills were attributed. From 1948 onwards he had frequented the surgery with minor complaints and at the time of a febrile illness in 1955 he had complained of pain below the left costal margin. His spleen had not been palpable. In January, February, and November 1961 he had had nocturnal epileptic fits and had also had a number of minor episodes in which he became confused, sweated profusely, and afterwards mentioned
TABLE 2
CASE 2. REPRESENTATIVE HAEMATOLOGICAL VALUES 1962–73

<table>
<thead>
<tr>
<th>Date</th>
<th>Hb (g/dl)</th>
<th>RBC (10⁹/l)</th>
<th>WBC (10⁹/l)</th>
<th>Platelets (10⁹/l)</th>
<th>PCV %</th>
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<tr>
<td>6.</td>
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FIG. 3 Case 2. Chart showing relationship of chorea to severity of polycythaemia, as indicated by the haemoglobin level, and beneficial effects of blood loss at operation and of subsequent venesection and of subsequent venesection and of subsequent venesection.

a feeling of being 'muddled up' and of experiencing vividly events long past.

On examination he was a tall thin man with a lugubrious and unchanging expression. His speech was slurred. His cheeks, lips, and tongue were crimson, his fingers and toes brick red. He had signs of mitral stenosis with normal rhythm and was not in congestive failure. He was moderately dyspnoeic and disliked lying flat. Liver and spleen were not palpable. Despite the improvement recorded in his psychiatric notes, he showed typical chorea of face, tongue, and extremities and his gait was interrupted by bizarre movements. His limbs were hypotonic and knee jerks pendular. His IQ was 108 on the Wechsler scale with no evidence of dementia. A sharp wave focus was recorded from the left temporal region in the EEG, which was otherwise normal. The spinal fluid was normal and the Wassermann reaction negative in blood and fluid. Serum copper oxidase activity was normal. Cardioscopy and electrocardiography (ECG) (Dr A. J. Thomas) confirmed tight mitral stenosis and, after a short period of observation, valvotomy was advised but refused. His haemoglobin level was slightly elevated but the other blood elements were in the expected range (Table 2).

His subsequent progress is recorded graphically in Fig. 3 and essential haematological data in Table 2. Little change was observed over the next two years apart from slowly progressive dyspnoea but fixity of expression and bradykinesia without tremor or
rigidity were frequently mentioned. These signs were attributed to the trifluoperazine without which his chorea became troublesome. At a visit in January 1964 he complained of increasing shortness of breath and his spleen was palpable. Deterioration over the following year led to his readmission in the autumn of 1965 when the cause of his obvious polycythaemia was again questioned. Mild dementia was confirmed by psychometry but the severity of his hypoxia was still held responsible for his polycythaemia. Eventually he agreed to operation and valvotomy was performed on 15 June 1966. Re- mission of chorea was immediate but it relapsed later in the year. The effect on his cardiorespiratory function was less satisfactory and his shortness of breath was not relieved.

Polycythaemia rubra vera was not established until April 1968 when Professor Allan Jacobs demonstrated a red cell volume of 52 ml/kg (normal range 26–33 ml/kg) and a total blood volume of 91 ml/kg (normal range 60–80 ml/kg). Repeated venesection improved his chorea but sustained control of the polycythaemia proved difficult and radioactive phosphorus was given in October 1969, July 1970, and August 1971. With the improved blood picture, his chorea again subsided but never disappeared and mild relapse often signalled, or preceded, a significant rise in the haemoglobin level.

3. DISCUSSION

Chorea is a rare complication of polycythaemia. Since 1909 when Bardachzi and Umney published their reports within a few days of each other only 22 cases have appeared in journals circulating in Britain (Table 1). This contrasts with the frequency of strokes and other neurological disasters in the literature of polycythaemia. Several of the early accounts, beginning with Vaquez’s classic case of vertigo (Vaquez, 1892), mentioned the nervous system (Cabot, 1899; Osler, 1903; Hutchison and Miller, 1906) and a neurological syndrome soon became commonplace in the natural history of the disorder (Jacobs, 1912; Lucas, 1912; Christian, 1917; Weber, 1921; Brockbank, 1929; Sloan, 1933; de Secondi, 1940). In recent years, the predominance of cerebral symptoms has emerged from many reviews and the neurological complications of polycythaemia, both central and peripheral, are now regarded as a major source of morbidity and mortality (Tinney et al., 1943; Videbaek, 1950; Johnson and Chalgren, 1951; Lawrence et al., 1953; Calabresi and Meyer, 1959; Croizat et al., 1960; Silverstein et al., 1962; Campbell et al., 1970).

The chorea of polycythaemia is usually a lone disorder, although it may sometimes be associated with other cerebral and peripheral complications. It resembles hereditary, rheumatic, and senile chorea and, particularly in its orofacial-lingual emphasis, the tardive dyskinesia of prolonged phenothiazine treatment (Hunter et al., 1964; Crane, 1968) and the chorea induced by the oestrogens of the contraceptive pill (Fernando, 1966; Lewis and Harrison, 1969) or by the dopamine of levodopa therapy (Cotzias, 1969; Godwin-Austen et al., 1969; Yahr et al., 1969). It should be distinguished from choreoathetosis due to infarction of basal nuclei and their associated pathways (Martin, 1957) and from the many types of metabolic disturbance including the acute lesions of hypoxia (Foley, 1954; Cree, 1969), hypernatraemia (Mann, 1969), and withdrawal of alcohol (Mullin et al., 1970) and such chronic disorders as hepatolenticular degeneration (Wilson, 1912) and hepatic cirrhosis (Victor et al., 1965). It is also distinct from the transient involuntary movements which may precede a cerebral haemorrhage (Cabot, 1899).

Crosetti (1929) mentioned polycythaemia in a case of Huntington’s chorea and Doll and Rothschild (1922) described an extraordinary family afflicted with the two disorders. In none of the reported cases of polycythaemic chorea (Table 1) nor in the two cases described above was a family history of dementia or of chorea obtained.

The first of the two patients reported by Gautier-Smith and Prankerd (1967) had had Sydenham’s chorea at puberty and the young woman described by Naville and Brütsch (1919) had had three attacks of chorea in the four years before the onset of catastrophic symptoms more certainly attributable to polycythaemia. Paralytic hemichorea followed by more generalized symptoms which persisted for six months was diagnosed in our second case 29 years before his presentation with polycythaemic chorea.

The occasional involvement of basal structures during the course of polycythaemia is indicated by the reports of tremor (Lucas, 1912; Dameshek and Henstell, 1940; Kramer, 1961), dystonia resembling Wilson’s disease (Brockbank, 1929), narcolepsy with chills, tremors, and cataplexy (Lhermitte and Peyre, 1930), and essential
tremor with additional extrapyramidal signs (Liessens, 1951).

Epilepsy, another uncommon feature of our second case, has previously been reported by Naville and Brütsch (1919), Tinney et al. (1943), Alajouanine et al. (1952), and Forsberg (1962).

The extent of the pathological lesion underlying chorea is unknown (Yahr, 1972). Denny-Brown (1962) suggested that degeneration of basal ganglia was matched by a spectrum of clinical phenomena: as the disorder advanced the shifting postures of chorea and athetosis changed into the fixed attitudes of dystonic rigidity. As chorea is a symptom of early disease the visible changes might well be slight.

Three cases of polycythaemic chorea have been studied at necropsy (Schiff et al., 1936; Kotner and Tritt, 1942; Bietti et al., 1968). Congestion of cerebral and meningeal veins, multiple small thrombi, and scattered zones of periventricular demyelination were findings common to all three; in addition, subdural and putaminal haemorrhages were described in the Italian case (Bietti et al., 1968). The basal ganglia shared in the generalized venous engorgement but local lesions were reported only by Schiff et al. (1936) who found intense demyelination of the anterior pallidum on both sides. The benign consequences of long-continued venous engorgement—at least in terms of light microscopy—were emphasized by Courville (1958). In the brain of a polycythaemic respiratory cripple, also an alcoholic, whose last months had been complicated by signs of dementia and Parkinsonism, he found an extraordinary degree of venous engorgement but a dearth of thrombi and more or less intact arteries and capillaries. Despite the presumed severity of carbon dioxide retention, the signs of tissue injury were slight and almost wholly attributable to agonal changes.

At the molecular level, information has come principally from cases of Huntington's chorea but also from chorea induced by phenothiazines, butyrophenones, levodopa, and oestrogens (Klawans et al., 1970). The normal concentration of catecholamines in the striatum suggests that dopamine is more active than acetylcholine in the conditions which cause chorea and that this functional derangement is eventually accompanied by loss of type II Golgi cells, particularly in the caudate nuclei (Barbeau, 1973). The release which follows of the globus pallidus and substantia nigra from the inhibitory caudato-fugal fibres is the probable basis of chorea. As the neurotransmitter of these fibres is gamma-amino butyric acid (GABA), support for this hypothesis has come from the studies of Perry et al. (1973) and of Bird et al. (1973) who found reduced levels of GABA and its biosynthetic enzyme glutamic-acid decarboxylase in the brains of patients dying of chorea. The latter authors (Bird et al., 1973) also found reduced activity of choline-acetyltransferase, further evidence of depressed cholinergic transmission.

Although the fluctuations of polycythaemic chorea suggest a reversible biochemical lesion nothing is yet known of catecholamine and acetylcholine activity. Stagnation of the capillary and venous circulation is almost certainly the immediate precipitant of striatal dysfunction but the rarity of the syndrome suggests another and more subtle defect. Riddoch et al. (1971) in their discussion of oestrogen-induced chorea came to a similar conclusion. The slowed circulation of polycythaemia was attributed by Millikan et al. (1960) to increased viscosity of the blood with resultant rise in cerebrovascular resistance and fall in blood flow (Nelson and Fazekeas, 1956). Abnormal stickiness of platelets (Shield and Pearn, 1969) adds to the stagnation. Multiple small thrombi form when the fragile platelets disintegrate en masse (Fiehler, 1950) but they readily break up (Rosenthal, 1949) and allow the circulation to return promptly to the obstructed vessels; hence the lack of visible tissue injury (Courville, 1938).

4. Conclusion

The importance of chorea as a warning sign of polycythaemia far outweighs its rarity as, in the majority of cases, diagnosis leads to effective therapy and to avoidance of a catastrophic illness (Table 1). In the child reported above (case 1), venesection was at first performed reluctantly as the polycythaemia was compensatory but the result fully justified the decision. The subsequent hemiplegia should not be regarded as a total failure of treatment: the early literature of polycythaemia vera abounds with examples of generalized thrombosis in untreated patients, such as the cases of Hutchison and Miller (1906),
Umney (1909) and Naville and Brütsch (1919). If this child's polycythaemia had been left uncorrected, the thrombosis might well have extended to other vessels.

The adult patient (case 2) is a lesson in the risks of delayed diagnosis. Had polycythaemia rubra vera been diagnosed in 1962 and treated appropriately, years of needless therapy with phenothiazines would have been avoided and valvotomy might have been postponed indefinitely. It is very probable that the drug which alone controlled his chorea, trifluoperazine, so damaged his striatal neurones that full remission of his chorea became impossible. In an environment of transmitter imbalance, perhaps aggravated by carbon dioxide retention, a dwindling population of dopaminergic receptor neurones, damaged first by adolescent chorea and secondly by chronic phenothiazine blockade, may have become supersensitive to dopamine, thus releasing lower centres from inhibitory control. His epilepsy may also indicate deficiency of gamma-aminobutyric acid. The persistence of his chorea despite a normal blood count and withdrawal of trifluoperazine suggests, like the slowly progressive dementia, Parkinsonian mask, and bradykinesia, that an initially reversible disorder has now passed beyond the point of recovery.

ADDENDUM

Aminoff et al. (1974) have recently reported an abnormal uptake of dopamine and of 5-hydroxytryptamine by the platelets of patients with Huntington's chorea. This suggests a mechanism in polycythaemia, whereby an excess of dopamine might be concentrated near receptors by increased numbers of normally loaded platelets circulating sluggishly through the basal ganglia. The chorea thus induced would be theoretically reversible by receptor blockade—for example, by phenothiazines—or by reducing platelet numbers by venesection. This mechanism would also account for the familial cases of polycythaemia with Huntington's chorea.

We are grateful to Dr L. G. Davies and to Dr A. J. Thomas for their cardiological advice in cases 1 and 2 and to Professor A. Jacobs for the haematological investigation of case 2.

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