THE DIAGNOSIS AND MANAGEMENT OF PULMONARY EMBOLISM

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Embolic of the pulmonary arteries presents in one of three main ways. Most dramatic is acute failure of the circulation. A large clot obstructs more than half of the cross-section of the pulmonary tree at the bifurcation of the main stem or the hila of both lungs. More commonly it is the development of an infarct at the base of one or both lungs that gives evidence of an embolism which has lodged some hours before. Rarely, repeated small emboli, which do not separately give rise to symptoms or signs, may lead to mounting pulmonary hypertension and chronic right heart failure.

The Attendant Circumstances

Pulmonary embolism is distinctly uncommon in patients who have previously been healthy and fully mobile but attention was drawn to this possibility by Homan's (1943) report of 11 such cases. Of the 120 cases of pulmonary embolism reported by Short (1952), three were well and fully mobile at the time of their first symptom.

Any patient confined to bed for a few days may develop the disease. Especially at risk are those ill with heart disease (Carlotti, Hardy, Linton and White, 1947; Short, 1952; Byrne, 1955), neoplasms (Trousseau, 1877; Sproul 1938), injuries to the bones of the legs (Sevitt and Gallagher, 1959), recent operations (Barker Nygaard, Walters and Priestley, 1940) or thrombophlebitis (Byrne, 1955). Although almost every patient with pulmonary embolism or pulmonary infarction has a source of thrombus formation in the venous system of the abdomen or legs, the condition will be diagnosed on the evidence in the heart or lungs. In two necropsy studies, McLachlin and Paterson (1951) found venous thrombosis in all of 19 cases of pulmonary embolism and in the series of Sevitt and Gallagher (1959) 32 of 38 patients with venous thrombosis had pulmonary embolism. Clinical methods of detecting venous thrombosis are, however, imperfect and in the absence of signs in the legs the absence of extensive thrombosis cannot be assumed. Thus, in Sevitt’s patients, 9 of 17 who died of pulmonary embolism were without clinical evidence of thrombosis.

In a clinical study of 72 patients with pulmonary embolism reported by Barritt and Jordan (1961), 32 were found to have tenderness of the calf on squeezing, 20 oedema of the leg, 10 superficial thrombophlebitis, 17 varicose veins and 16 no abnormal signs. Clinical evidence of a possible source in the leg may, therefore, be taken as a confirmatory sign of pulmonary embolism but in its absence no conclusions can be drawn.

Evidence of Obstruction to the Pulmonary Tree

Massive embolism of the pulmonary artery may be followed by immediate loss of consciousness and death within minutes. The patient is usually cyanosed and the pulse rapid and of very low volume. If the patient survives, evidence of severe circulatory obstruction will persist and the following are the principal clinical features:

Breathlessness and Restlessness. Anxiety accompanies breathlessness which may show itself with the least movement about the bed. This immediate breathlessness is not due to pleural pain but indicates a profound disturbance of ventilation. Orthopnea is unusual.

Faintness. With low levels of blood pressure there may be feelings of faintness or loss of consciousness on sitting up. Faintness in the early period of getting out of bed after operations should be viewed with particular suspicion.

Central Chest Pain. Discomfort in the centre of the chest is indistinguishable from the pain of myocardial infarction. Radiation to the arms or jaw is uncommon but the pain may be severe and occasionally may last for hours.

Tachycardia. A rise in heart rate is a very frequent accompaniment of pulmonary embolism. In cases where the first symptoms are those of the development of an infarct, pulse charts frequently show a sharp rise in heart rate some hours before the onset of pleural pain, and this serves to date the arrival of the embolus. Paroxysmal arrhythmias also occur. Short (1952) considered that five of his 120 patients developed atrial fibrillation as a result of embolism and one other had ventricular tachycardia. Barritt and Jordan (1961) confirmed the onset of atrial fibrillation in four
of their 72 patients and paroxysmal supraventricular tachycardia in a fifth. The onset of atrial fibrillation during the course of heart disease should, therefore, lead to thought being given to the possibility that pulmonary embolism has occurred.

**Hypotension.** An abrupt fall in blood pressure can be taken to indicate a reduced cardiac output. Occasionally, the diagnosis is first suggested when routine records of blood pressure show a fall in systolic pressure to less than 100 mm. Hg. As in other low output states an obvious fall in systolic pressure during inspiration may be a feature but pulsus paradoxus on palpation is rare. With the low blood pressure and poor volume pulse the skin is cold and clammy.

**Cyanosis.** Arterial oxygen desaturation is very difficult to detect in the patient with a cold skin, but cyanosis of the tongue and mucous membranes is occasionally obvious and arterial puncture will then confirm the desaturation. An increase in the dead space within the lung and disturbance of ventilation perfusion ratio, together with diminished compliance are likely to be the causative mechanisms.

**Raised right atrial pressure.** Obstruction to the outflow from the right ventricle leads to
a rise in right atrial pressure that may be obvious in the neck. In others, breathlessness is accompanied by such large variations in intra-thoracic pressure, that the mean right atrial pressure cannot be estimated clinically.

Auscultation of the Heart. Auscultation of the heart gives little information. In the presence of tachycardia gallop rhythm may be heard. In a small number of patients, friction sounds are present for short periods at the base of the heart. Abrupt distension of the pulmonary artery or of the right ventricle are probably responsible.

The Electrocardiogram

Characteristic changes can be expected in the electrocardiogram of patients with massive embolism of the pulmonary arteries. These consist of a shift to the right of the electrical axis of the heart with clockwise rotation and T-wave changes in right ventricular leads. Cutforth and Oram (1958) analysed records from 50 cases of pulmonary embolism, of which 28 showed evidence of a fall in blood pressure or a rise in venous pressure (severe cases) and 22 neither feature (mild cases). These authors considered the following three electrocardiographic patterns to be diagnostic of pulmonary embolism:

1. $S_1$, $Q_3$, $T_3$ plus right ventricular T-wave inversion.
2. $S_1$, $T_3$ or $T_3$ plus right ventricular T-wave inversion.
3. $S_1$, $Q_3$, $T_3$ plus right bundle branch block.

Of their 28 severe cases, 18 showed these patterns and nine of the remaining ten showed changes considered to be suggestive of pulmonary embolism. Of their 22 mild cases, there was diagnostic evidence of pulmonary embolism in the electrocardiogram in half. Barritt and Jordan (1961) noted T-wave inversion in $V_1$ and $V_2$ in 14 of their patients, of whom 11 also showed T-wave inversion in leads II or III or aVf. Transient right bundle branch block was present in a further four patients. Of these 18, there was a fall in blood pressure and other evidence of massive embolism in 16. Changes in the electrical position of the heart were present in 13 others whilst of 12 patients with normal ECGs none had a fall in blood pressure. Obvious changes in the electrocardiogram, therefore, are likely in severe cases and suggestive evidence may strengthen the diagnosis in those with less circulatory disturbance.

In a small number of acutely ill patients with low blood pressure and high venous pressure, the electrocardiogram may be less helpful. Medd and McBrien (1962) cite patients with massive pulmonary embolism with S-T depression or T-wave changes in left precordial leads.
which suggest the presence of coronary artery disease. Figure 2 shows the electrocardiogram in a woman of 40 years whose first symptom was a brief attack of effort syncope whilst on holiday. Very high venous pressure with tachycardia, small volume pulse and pulsus paradoxus suggested the diagnosis of cardiac tamponade. Shortly after attempted aspiration of the pericardium she died and at necropsy massive, repeated embolism of the pulmonary arteries was found.

Evidence of Pulmonary Infarction
When thrombus becomes tightly packed in a branch of the pulmonary artery, symptoms and signs of pulmonary infarction usually follow. This is more likely to occur when the embolus passes beyond the bifurcation into one or both branches towards the periphery of the lung. Only the lower lobes are usually affected and one may assume that this is because blood flow to the upper lobe is extremely small at rest in the upright position (Dollery, Hugh-Jones, Matthews, 1962).

Pleural Pain—Friction and Effusion
Pleural pain is the commonest symptom to draw attention to pulmonary embolism. It may be slight or severe and on one or both sides. A friction rub may or may not accompany it. In patients whose embolism is dated by the onset of faintness, breathlessness or tachycardia, pleural pain may follow some hours later. A pleural effusion may develop. The fluid is usually bloodstained.

Cough and Haemoptysis
Cough, with the production of a small quantity of sputum, is frequently present. Haemoptysis, which varied in amount from a speck of blood on one occasion only, to heavy staining of the sputum for many days, was present in 31 of 72 of Barritt and Jordan’s patients.

Breathlessness and Cyanosis
In the presence of pulmonary infarction, dyspnoea is most commonly the result of pleural pain. The pain is avoided by taking shallow, rapid breaths. More severe breathlessness points to a high degree of circulatory obstruction. Cyanosis is usually inconspicuous except in the presence of extensive bilateral infarction.

Physical Signs in the Chest
Breathing is usually quiet but increased respiratory variation in venous pressure observed at the root of the neck may point to diminished lung compliance.

Percussion of the chest usually confirms the presence of a lung lesion because resonance is impaired at the affected base. The loss of resonance rarely amounts to stony dullness. In contrast to the findings with consolidation of lobar pneumonia, breath sounds and voice sounds are diminished over an infarct. Bronchial breathing is a rare finding. The most frequent auscultatory sign is the presence of rales. Rales are rarely absent in pulmonary infarction, and are present at both bases in about half the cases, thus pointing to the presence of clots in both lungs.

Fever and Jaundice
Fever is an almost invariable accompaniment of pulmonary infarction. Characteristically, fever is present at the time of onset of pleural pain or haemoptysis. With effective anticoagulant therapy and without the addition of antibiotic agents, fever soon disappears. Jaundice is much less frequent, especially in the absence of congestive heart failure.

The Chest Radiograph
Although the patient cannot be taken to the X-ray Department, and despite breathlessness and pain at the time of the examination, changes in the chest radiograph will usually be found at the first examination in the ward. Short (1951) described such changes in 85 of 94 patients examined. The abnormalities were infarct shadows in 88 per cent, pleurisy in 56 per cent, elevation of the diaphragm in 39 per cent, and multiple infarcts in 43 per cent. These positive radiological changes were usually present within 12 hours of infarction. A negative chest X-ray should not weigh heavily against the diagnosis of pulmonary embolism, particularly if the film is of poor quality. Embolism without infarction may be present and here the radiological changes may be much less conspicuous. Shapiro and Rigler (1948) considered the characteristic abnormalities in embolism without infarction to be increased radio-translucency and ischaemia of the involved pulmonary segment, with abrupt termination of the embolised pulmonary artery.

Pulmonary Function Tests
Obstruction of a major branch of the pulmonary artery must necessarily cause an important disturbance of pulmonary function. Perfusion of the affected lung segment will cease, ventilation of the unperfused lung will continue and when infarction occurs the inflammatory process will disturb the lung further. Robin, Forkner, Bromberg, Croteau and Travis (1960) discussed the alveolar gas exchange in pulmonary embolism. They list three major components of disturbance: arterial oxygen desaturation, hyper-ventilation, and the
Fig. 3.—Successive tracings with an interval of three hours in an elderly patient with extremely high venous pressure but no orthopnea. Pulmonary embolism was suspected and consideration was given to surgical exploration. Progressive changes of left bundle branch block suggested myocardial infarction and necropsy confirmed the diagnosis.

Differential Diagnosis of Massive Pulmonary Infarction

When the physician is faced with a cardiovascular emergency in a patient with very low cardiac output, high venous pressure and central chest discomfort, the usual differential diagnosis is between massive pulmonary embolism and myocardial infarction. The distinction is usually not difficult for evidence of incipient pulmonary oedema together with the onset of an injury current in the electrocardiogram make the diagnosis of myocardial infarction. In the absence of these two features...
in the early hours of the emergency the circumstances of the attack may weight the scales in one direction or another. Clear evidence of venous disease of the legs suggests pulmonary embolism, and a fleck of blood stained sputum in the absence of wet lungs is confirmatory. Again, electrocardiography will usually bring confidence to the diagnosis of embolism. If the patient’s condition is not deteriorating, reliance should be placed on serial electrocardiograms at frequent intervals (Fig. 3). An early rise in serum glutamic oxalacetic transaminase favours muscle necrosis from myocardial infarction.

It may be felt that the differential diagnosis is barely of practical importance as treatment of these two conditions is so similar. There will, however, be occasions when consideration is given to the idea of surgical removal of an embolus and, under these circumstances, diagnosis must be certain. There may be a place here for more adequate X-ray visualisation of the disturbance in the pulmonary vascular tree. High quality plain films of the hila of the lungs will repay close study. The possibility of the bedside use of contrast radiography needs to be further explored. Robb and Steinberg (1939) claimed that accurate studies could be made of arterial and venous patterns of the lung by injection of contrast medium into an arm vein with the patient sitting before a cassette. Williams and Wilcox (1963) have recently made a study using this technique in 50 patients with pulmonary embolism. All the patients tolerated the procedure well but few details are given of the severity of their illness or the type of apparatus used. Mention is made of the possibility of this approach to the problem of the patient in whom surgical exploration is considered. Proof of major obstruction to vessels and localisation of the main areas of embolism might then be considered a necessary preliminary to operation.

When the emergency presents with paroxysmal arrhythmia differential diagnosis is likely to be more difficult. Treatment of the arrhythmia will take priority. The possibility of pulmonary embolism should always be considered when arrhythmia occurs in the course of heart disease of all types. The distinction from cardiac tamponade will not arise frequently. Where doubt exists a diagnostic aspiration is essential but one hopes that exploration of the praecordium with ultrasound recording may display or exclude a pericardial effusion without the use of a needle.

Pulmonary infarction must usually be distinguished from pneumonia especially in the post-operative period. No single feature is in itself decisive with the exception of the diagnostic electrocardiographic change which will be present in the minority of cases when a lung lesion is prominent. Physical signs of venous thrombosis will be present in the majority, and it is the combination of venous thrombosis and an appropriate pulmonary lesion that points to the diagnosis most frequently. Difficulties arise when there are no abnormal signs in the legs. Pneumonia will then often be diagnosed and antibiotics prescribed. The majority of patients will recover with these measures, but there will often be surprises as the late development of leg signs or recurrence of chest troubles, or even fatal embolism correct the diagnosis.

Where doubt exists response to anticoagulant therapy will sometimes give diagnostic help. With the infusion of heparin, fever may vanish within hours and not recur.

Repeated Minor Emboli leading to Progressive Pulmonary Hypertension

With the widespread recognition of the value of anticoagulant therapy in curing venous thrombo-embolism, repeated major embolisation leading to progressive right heart failure is a rare condition. A small group of patients exist, however, who exhibit progressive pulmonary artery obstruction who have minimal clinical evidence or no clinical evidence at all of pulmonary embolism or infarction. This condition appears in previously healthy people who have not been confined to bed. Goodwin, Harrison and Wilcken (1963) described 11 such patients and Wilhelmsen, Selander, Soderholm, Paulin, Varnauskas and Werko (1963) and a further 10. There is little except evidence of major filling defects in the pulmonary arteriogram, to distinguish this condition from so-called primary pulmonary hypertension. Thus, the patients present with breathlessness of effort, effort syncope, chest pain and haeomptysis. The electrocardiogram shows right ventricular dominance and catheterisation confirms the pulmonary hypertension. Goodwin lays great stress on the necessity for early recognition of this condition as successful arrest of the condition with anticoagulant treatment can be hoped for only before pulmonary hypertension becomes established. Slight effort dyspnæa with hyperventilation at rest is the clue. Under these circumstances arterial-end tidal CO₂
gradients may well be demonstrated and arterial oxygen desaturation on effort is likely.

**Management of Venous Thrombo-Embolism**

It would be inappropriate to discuss the treatment of pulmonary embolism without considering prophylaxis first. Attempts to diminish the incidence of pulmonary embolism in hospital by systematically allowing patients to get out of bed at an earlier stage of their illness have met with little success. This is probably because at an early stage of recovery from illness or operations the level of activity is of necessity very low and nothing is gained by exchanging bed rest for chair rest.

This has led to a study of the use of anticoagulant drugs given prophylactically to patients who are especially at risk. Sevitt and Gallagher (1959) in their classical study compared 150 elderly patients with fractured hips (control series) with a similar number who were given Phenindione routinely soon after admission to hospital. This prophylactic use of Phenindione resulted in complete freedom from pulmonary embolism in the period of treatment, whereas in the control series 18 per cent of patients developed pulmonary embolism and 10 per cent died of it. There was also a great saving in venous thrombosis in the legs. In a somewhat similar study, Chalmers, Marks, Bottomley and Lloyd (1960) tested the use of prophylactic anticoagulant therapy in selected groups of gynaecological and obstetric patients. In all they tested 1877 patients in a five year period and made a comparison with a preceding five year period when prophylactic anticoagulants had not been used. They too found an impressive reduction in venous thrombosis and pulmonary embolism in treated patients. A strong case can, therefore, be made for the use of anticoagulants given prophylactically to patients who are at special risk.

A different approach to the problem has been suggested by Sharnoff, Kass and Mistica (1962) on the basis of a state of increased blood coagulability as demonstrated by Lee-White clotting time values. They administered Heparin subcutaneously before operation and before mobilising the patient in the post-operative period. Their recommendations were not supported by a sufficiently extensive clinical trial to prove the value of this type of therapy.

**Treatment of Massive Embolism with Circulatory Failure**

As in other circulatory or respiratory emergencies, reassurance of the patient is of prime importance. The relief of anxiety by the confidence of the physician, aided where necessary by pain relieving drugs, will do much to economise respiratory effort and diminish heart rate.

When the pulmonary artery becomes nearly completely obstructed by a large clot, immediate therapy can only be directed to sustaining the contractile force of the right ventricle. So long as sufficient blood can be forced past the obstruction to maintain an impoverished circulation it can be expected that the emboli will tend to be packed into smaller branches, thus allowing full perfusion of some areas of the lung. In the critical period, maintenance of arterial oxygen saturation and perfusion pressure in the coronary circulation, and control of the heart rate are likely to assist the ventricle.

Arterial oxygen saturation will certainly fall under these circumstances and high concentrations of oxygen should be given by face mask.

With low central aortic pressure, coronary blood flow will fall. Mephenitine sulphate, metaraminol or infusions of angiotension or noradrenaline should be used if the systolic blood pressure falls below 90 mmHg. In the presence of signs of tachycardia or paroxysmal arrhythmia rapid digitalisation is indicated.

Heparin should be given at once to discourage the formation of fresh thrombus at sites where blood flow is obstructed by the embolism.

If the patient survives long enough for these measures to be put into operation then he can expect to recover with continued medical therapy. Of the 53 patients in Barritt and Jordan’s (1960) treated series, none died of pulmonary embolism who survived long enough to receive the first injection of Heparin.

This favourable outcome cannot, however, be expected in every case and some patients may survive for a number of hours with a falling blood pressure before they die. In these exceptional circumstances the question of surgical removal of the emboli may now be rationally considered. Reports of successful removal of clot have been given by Hampson, Milne and Small (1961) who operated at normal temperature without perfusion, by Alison, Dunnill and Marshall (1960) operating after surface cooling of their patient to 29° C and by Cooley, Beall and Alexander (1961) using cardiopulmonary bypass.

A planned operation of this type should
certainly only be carried out if deterioration continues in spite of intensive medical treatment.

In patients who die within minutes of their embolism, attempts to restore the circulation by continued external cardiac compression and artificial ventilation may succeed from time to time. Such methods are well directed to the younger patient stricken unexpectedly by a fatal circulatory disturbance.

Management of Pulmonary Infarction

The question of more prolonged therapy of pulmonary embolism can be considered under this heading. Additional features which call for further mention are pleural pain and fever. Pethidine is a suitable drug for the relief of the pain of pulmonary infarction and 50-100 mg. by intramuscular injection can be given at once and repeated if need be.

Unless there are strong contraindications, anticoagulant treatment will always be given. Recent operations are rarely a great hazard. In patients with a history suggesting active peptic ulceration it may be preferable to use Heparin as the only anticoagulant drug throughout the course of treatment. Heparin may be given by intermittent intravenous injection or by continuous infusion for many days. The latter method has the advantage that the normal clotting mechanism is restored very rapidly if the infusion is terminated. Usually, an oral anticoagulant will be administered at the same time as Heparin is started so that Heparin is discontinued after 36-48 hours as the oral anticoagulant takes over. Phenindione has been used most extensively in Britain in the last decade but occasional reports of serious toxic reactions affecting kidney, liver and bone marrow have led some authors to question its suitability (Holman). High levels of dosage are called for and one stage Quick times should be prolonged to 2½-3 times control times.

Antibiotic therapy is usually unnecessary for fever which will fall soon after the infarct begins to heal, in the same way as in myocardial infarction. Pulmonary infarcts can, however, become infected with organisms from the upper respiratory tract and this possibility should be borne in mind. Anticoagulant therapy should be continued until clinical evidence of infarction has been cleared and evidence of active disease in the leg has subsided. By this time, clearing of the changes on X-ray of the chest will usually be evident and in most cases a three weeks' course of treatment is adequate. If there are still signs of infarction, either on clinical or radiological grounds at the end of three weeks, then therapy should be prolonged. Occasionally, there will be recurrence of signs in the chest or leg and then a second course of treatment will be called for and it should be prolonged for four to six weeks. Occasionally, continued treatment for 6-12 months may be necessitated by recurring evidence of thrombo-embolism. Ligation of veins will very rarely be indicated.

Thrombolytic Therapy

Fibrinolytic enzymes may find a place in the treatment of pulmonary embolism. It is clear from the natural history of the condition that large clots are spontaneously lysed in the pulmonary artery and it is logical, therefore, to attempt to assist this process. To this end, activators of the Plasminogen-Plasmin reaction, particularly Streptokinase, have been used. Preparations of Streptokinase carry a risk of febrile reactions and in the critical state of the circulation after massive embolism this hazard contra-indicates the use of this type of therapy at the present time. Results with Heparin are so good that new remedies should not be used until more evidence is available of their superiority. Caution concerning the use of streptokinase has been expressed by McNicol, Douglas and Bayley (1962) who made a detailed study of Streptokinase infusions in patients with vascular occlusions.

Repeated Embolism with Pulmonary Hypertension

This uncommon condition calls for the institution of anticoagulant treatment at the earliest possible moment and the treatment should be pressed for months, or years, until all clinical and laboratory evidence of the disease has disappeared.

In the presence of heart failure, digitalis should be added but if this becomes necessary, arrest of the changes in the pulmonary artery cannot be expected.

REFERENCES

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