Acute pulmonary embolism

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Pulmonary embolism, despite being common, often remains elusive as a diagnosis, and clinical suspicion needs to remain high when seeing a patient with cardiopulmonary symptoms. Once suspected, diagnosis is usually straightforward; however, optimal treatment can be difficult. Risk stratification with clinical scores, biomarkers and imaging helps to refine the best treatment strategy, but the position of thrombolysis in intermediate risk (submassive) pulmonary embolism remains a grey area. Pulmonary embolism response teams are on the increase to provide advice in such cases. Direct oral anticoagulants have been a major advance in treatment this decade, but are not appropriate for all patients. Follow-up of patients with pulmonary embolism should be mandatory to determine duration of anticoagulation and to assess for serious long-term complications.

KEYWORDS: Pulmonary embolism, thrombolysis

Introduction

Pulmonary embolism (PE) is a common presenting diagnosis in an emergency department. It may present with classical features such as breathlessness and pleuritic chest pain, but also less characteristically, for example insidious onset breathlessness over days-to-weeks or syncope with relatively few respiratory symptoms. Therefore, clinicians need to have a high degree of suspicion for PE in patients presenting with potential cardiopulmonary symptoms, since the consequences of missing or delaying the diagnosis of PE can be severe. As with most other areas of medicine, PE diagnosis and management has become heavily protocolised, but as will be discussed, there are still many grey areas in decision making in PE, necessitating experienced senior clinical decision making.

Diagnosis

Diagnostic algorithms and techniques have remained relatively unchanged over the past 10 years, with computed tomography (CT) pulmonary angiography (PA) being the principal tool. Since it involves ionising radiation, is not appropriate to use CTPA in all cases of suspected PE, hence clinical probability scores and d-dimer testing are used to filter out those with low probability of PE. While there are several clinical probability scores, the Wells score remains the predominant score in international guideline algorithms. When clinical probability of PE is low, a normal d-dimer has a high negative predictive value for excluding PE, however where the d-dimer is elevated or the clinical probability of PE is high, diagnostic imaging should be performed.

ABSTRACT

Pulmonary embolism (PE) is a common presenting diagnosis in an emergency department. It may present with classical features such as breathlessness and pleuritic chest pain, but also less characteristically, for example insidious onset breathlessness over days-to-weeks or syncope with relatively few respiratory symptoms. Therefore, clinicians need to have a high degree of suspicion for PE in patients presenting with potential cardiopulmonary symptoms, since the consequences of missing or delaying the diagnosis of PE can be severe. As with most other areas of medicine, PE diagnosis and management has become heavily protocolised, but as will be discussed, there are still many grey areas in decision making in PE, necessitating experienced senior clinical decision making.

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Interest is increasing in using age-adjusted d-dimer, however, evidence has not yet reached significance to be considered part of routine practice. In certain circumstances, in particular pregnancy, d-dimer, inflammation and cancer cannot be used to triage patients for imaging. An alternative strategy to CT is to consider ultrasound Doppler of the leg veins where it is important to avoid ionising radiation, such as in the pregnant patient, but this strategy in the non-pregnant patient is not of sufficient yield to justify its use, resulting in a positive find in approximately 1 in 10 cases.1 Ventilation-perfusion (VQ) scanning can be used where available, in the presence of a normal chest radiograph. Performing a half-dose perfusion scan is also an option in the pregnant patient. With advancing technology, CTPA is able to detect smaller filling defects in the pulmonary circulation. Where it is certain that these filling defects are genuine pulmonary emboli, it is important for the clinician to assess whether such small volume data account for the presentation, for example, is an isolated subsegmental PE sufficient to account for patient presenting with syncope? Conversely, where there has been poor contrast opacification or breathing artefact, scans may be misinterpreted as showing small peripheral embol and there is poor inter-observer agreement where emboli are <6 mm or subsegmental.1 Given the important consequences of diagnosing PE, such as the need for lifelong anticoagulation, it is important in these circumstances to review scans carefully and sometimes consider repeating or opting for a different modality.

Risk stratification

Risk stratification plays an important role in management of the suspected and confirmed PE (Fig 1). Systemic lysis is indicated for the high-risk (previously known as massive) PE or in the context of cardiac arrest due to suspected PE. Where lysis is contraindicated, surgical embolectomy should be considered. This group represents a relatively small proportion of patients presenting with PE (<5%) and the greater challenge is to identify patients in whom thrombolysis/reperfusion therapy should be considered from the non-high-risk group.

The European Society of Cardiology (ESC) 2014 guidelines pioneered the concept of risk-based management (Fig 2).1 Many clinical risk scores exist, but perhaps the one that has gained the most traction is the Pulmonary Embolism Severity Index (PESI) and subsequently its simplified version (sPESI, Table 2). These scores can be made at the bedside using clinical history and simple observations. Intermediate risk patients can be considered those in PESI classes III–V or sPESI >0. This represents a large cohort of patients and most will not benefit from thrombolysis and therefore further risk stratification is required.

The ESC guidelines suggest that intermediate risk patients who also have an elevated cardiac biomarker, such as troponin or N-terminal pro-brain natriuretic peptide, and have evidence of right ventricular dysfunction on CT or echocardiography, should be labelled as intermediate-high risk and be considered for thrombolysis/reperfusion. However, the Pulmonary Embolism International Thrombolysis Trial (PEITHO), a large study of over 1,000 patients with this risk profile, demonstrated no mortality benefit at 7 or 30 days with thrombolysis, but did demonstrate a significant increase in adverse events related to bleeding.4 In the extension study, long-term outcomes including incidence of chronic thromboembolic pulmonary hypertension were no different.5

When comparing against the expected mortality from the PESI scoring system (Table 2), the overall risk of mortality was low in the short-term PEITHO study, approximately 2%, which may reflect either better overall patient management as part of a clinical trial, such as closer observation, or a bias towards excluding enrolment of higher-risk patients according to clinician judgement. Therefore, there does remain a need to identify a further subgroup of intermediate-high risk patients who might benefit from more aggressive therapy. One study has suggested that patients with a combination of a sPESI greater than zero, positive brain natriuretic peptide and troponin, and finding of proximal deep vein thrombosis have the highest risk of complicated outcomes, with positive cardiac biomarkers outperforming, and therefore negating...
analysis has suggested that catheter directed lysis and half-dose lysis may provide optimal outcomes in terms of reduced mortality and bleeding, but the supporting data are relatively weak. Most recently, the EKOS® system, which co-ministers local thrombolysis with ultrasound to increase fibre in separation via pulmonary artery catheter, has been shown to improve right ventricular to left ventricular diameter ratios with low doses of tissue plasminogen activator. The role for this particular technique and half-dose thrombolysis are yet to find their place in guideline algorithms, and their use remains at the discretion of senior clinicians / PERTs.

Patients with intermediate-low risk PE should usually be admitted to hospital for a short inpatient stay for anticoagulation treatment. There are no clear guidelines on how long patients should be admitted, but 48 hours seems appropriate. Patients with low risk PE can now be considered for full outpatient management following the British Thoracic Society guidelines (Figure 3). Patients in PESI I/II or sPESI 0 may be sent home provided they do not have any other social or medical reasons for admission (Box 1). Alternatively another set of criteria may be used, known as the Hestia criteria, which have been specifically validated for outpatient selection. Clinicians may wish to use a negative cardiac biomarker to ‘override’ the finding of right ventricular dilatation the need for, right ventricular imaging on multivariate analysis. A post-hoc analysis of the PEITHO study (in those already in the intermediate-high risk group) suggested that two or more of the following: systolic blood pressure ≤ 110 mmHg, respiratory rate > 20 breaths per minute, history of heart failure and active cancer were associated with the highest risk of major adverse outcomes. Thus, there appears to be a number of risk factors associated with unfavourable outcomes but as yet no clear guidance can be given on when to administer thrombolysis.

Balancing the risk of bleeding against risk of PE-related mortality requires shared decision making between senior clinicians and patient, including informed consent. In recent years, there has been a rise of pulmonary embolism response teams (PERTs), particularly in the USA but more recently in the UK, to provide expert advice in such scenarios. The need for PERTs has been increased with the advent of more complex therapies such as catheter directed thrombolysis, which uses very low doses of lytics, as well as the potential use of half-dose systemic lysis. A recent network meta-analysis has suggested that catheter directed lysis and half-dose lysis may provide optimal outcomes in terms of reduced mortality and bleeding, but the supporting data are relatively weak. Most recently, the EKOS® system, which co-ministers local thrombolysis with ultrasound to increase fibre in separation via pulmonary artery catheter, has been shown to improve right ventricular to left ventricular diameter ratios with low doses of tissue plasminogen activator. The role for this particular technique and half-dose thrombolysis are yet to find their place in guideline algorithms, and their use remains at the discretion of senior clinicians / PERTs.

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<table>
<thead>
<tr>
<th>PESI risk class</th>
<th>Total PESI points</th>
<th>PESI 30-day mortality</th>
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<tbody>
<tr>
<td>I: very low</td>
<td>≤65</td>
<td>0–1.6%</td>
</tr>
<tr>
<td>II: low</td>
<td>66–85</td>
<td>1.7–3.5%</td>
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<tr>
<td>III: intermediate</td>
<td>86–105</td>
<td>3.2–7.1%</td>
</tr>
<tr>
<td>IV: high</td>
<td>106–125</td>
<td>4.0–11.4%</td>
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<tr>
<td>V: very high</td>
<td>≥126</td>
<td>10.0–24.5%</td>
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</tbody>
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<table>
<thead>
<tr>
<th>sPESI risk class</th>
<th>Total sPESI points</th>
<th>sPESI 30-day mortality</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low</td>
<td>0</td>
<td>1.0%; 95% CI 0.0–2.1%</td>
</tr>
<tr>
<td>High</td>
<td>&gt;0</td>
<td>10.9%; 95% CI 8.5–13.2%</td>
</tr>
</tbody>
</table>

CI = confidence interval; PESI = Pulmonary Embolism Severity Index; RR = respiratory rate; SaO₂ = arterial oxygen saturations; SBP = systolic blood pressure; sPESI = simplified Pulmonary Embolism Severity Index.
### Anticoagulation

The mainstay of treatment of PE is anticoagulation. Until recently, the standard of care was low molecular weight heparin (LMWH) followed by warfarin, but in recent years this has been replaced by the direct oral anticoagulants (DOACs). Apixaban, dabigatran, edoxaban and rivaroxaban have all been licensed for the treatment of venous thromboembolism. Apixaban and edoxaban both require a minimum 5-day lead-in period with LMWH, whereas apixaban and rivaroxaban may be administered as soon as PE is confirmed with an initial high-dose regime tapering down at 7 and 21 days, respectively. Overall, the DOACs as a class are non-inferior to LMWH. In two recent studies with edoxaban and rivaroxaban have shown improved efficacy over LMWH in CAT, but increased bleeding. The International Society of Thrombosis and Haemostasis, have produced guidance to state that these two DOACs may be considered in place of LMWH where bleeding risk is low.

PE may be the presenting feature of underlying malignancy, but screening for cancer remains controversial. At present the National Institute for Health and Care Excellence (NICE) guidelines (CG144) still recommend offering patients with first unprovoked PE over the age of 40 investigation for cancer including physical examination, blood tests (including full blood count, calcium and liver function tests), urinalysis, abdomen-pelvic CT and mammogram in women.

### Special circumstances

#### Pregnancy

PE in pregnancy should be managed in the same way as in patients who are not pregnant but with several caveats: diagnosis is discussed above; the PESI scoring system cannot be applied and thus a decision as to whether to manage a patient as an outpatient can only be made after discussion between a senior physician and obstetrician; patients should be anticoagulated with LMWH, not DOACs; and pregnancy should be considered a relative contraindication to thrombolysis.

#### Cancer

Guidelines presently recommend the use of LMWH as the first line anticoagulant in cancer-associated thrombosis (CAT), however, two recent studies with edoxaban and rivaroxaban have shown improved efficacy over LMWH in CAT, but increased bleeding. The International Society of Thrombosis and Haemostasis, have produced guidance to state that these two DOACs may be considered in place of LMWH where bleeding risk is low.

#### Right heart thrombus

The presence of right heart thrombus increases the risk of mortality due to the risk of pulmonary outflow tract obstruction and increased right ventricle afterload. Free-floating clot in the right-sided cardiac chambers should be considered an indication for systemic thrombolysis, however if the clot is straddling a patent foramen ovale, then surgical embolectomy may be appropriate. Anticoagulation is preferred for patients with a clot adherent to the endocardium unless it is mobile and risks valvular obstruction, in which case surgery may also be considered.

#### Follow-up

Detailed discussion regarding follow-up of patients with PE is beyond the scope of this review, however it is critical that patients should be fed in to a clinic to review the outcome of treatment, in particular to assess for the possibility of chronic thromboembolic pulmonary hypertension and determine duration of anticoagulation. Where investigations for possible underlying malignancy have been performed, this also provides an additional safety net. There is also a high burden of psychological issues following PE, which can largely be addressed by a physician, but also referred on for psychological support where necessary.

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**Box 1. British Thoracic Society exclusion criteria for outpatient management**

- Haemodynamic instability (HR >110 bpm, SBP <100 mmHg, requirement for inotropes or critical care, requirement for thrombolysis or embolectomy)
- Oxygen saturations <90% on air
- Active bleeding or risk of major bleeding (eg recent GI bleed or surgery, previous intracranial bleeding, uncontrolled hypertension)
- On full-dose anticoagulation at the time of the PE
- Severe pain (for example requiring intravenous analgesia)
- Chronic kidney disease stage 4 or 5 (eGFR <30 mL/min) or severe liver disease
- Heparin-induced thrombocytopenia within the last year and where there is no alternative to repeating heparin treatment
- Other medical or social reasons necessitating inpatient management

eGFR, estimated glomerular filtration rate; GI = gastrointestinal; HR = heart rate; PE = pulmonary embolism; SBP, systolic blood pressure.
References


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