

Pulmonary Embolism

/Pulmonary hypertension

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MD

PE

Introduction

Source of emboli

Pathogenesis & Risk factors

S&S

Management approach:

- Assess clinical probability

- Assess risk of mortality

- Investigation

- * Diagnostic

- *Non diagnostic (helpful test)

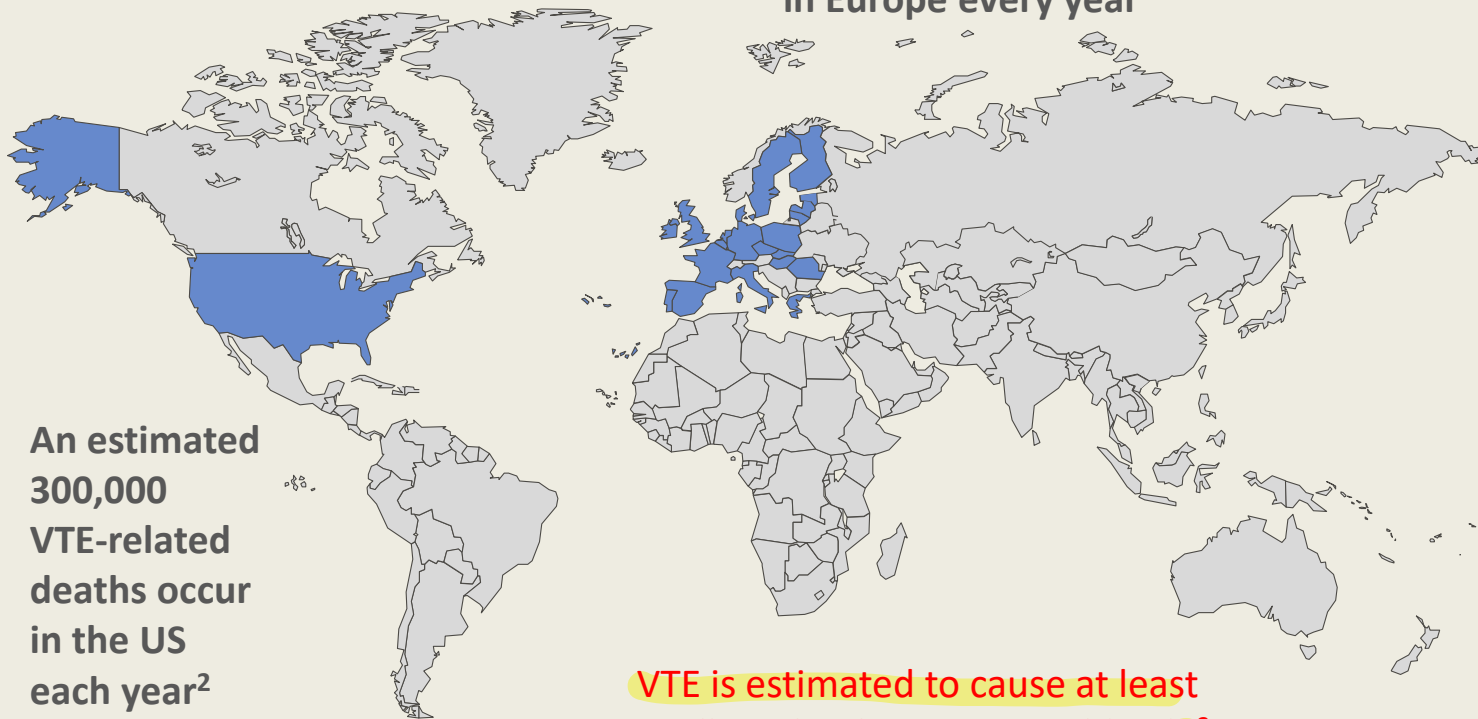
- Treatment (medications and duration of treatment)

Introduction

- Partial or complete **occlusion** of a pulmonary arterial branch by blood clot(thrombus or multiple thrombi).
- Deep vein thrombosis and PE are different presentations of the same underlying pathophysiological event, venous thromboembolism (VTE). *There should be shunting.*

VTE Is a Leading Cause of Death Worldwide

VTE is estimated to cause >500,000 deaths
in Europe every year¹



An estimated
300,000
VTE-related
deaths occur
in the US
each year²

VTE is estimated to cause at least
3 million deaths a year worldwide³

1. Cohen AT *et al*, *Thromb Haemost* 2007;98:756–764; 2. Heit JA *et al*, *Blood* 2005;106:Abstract 910;
3. ISTH Steering Committee for World Thrombosis Day *J Thromb Haemost* 2014;12:1580–1590

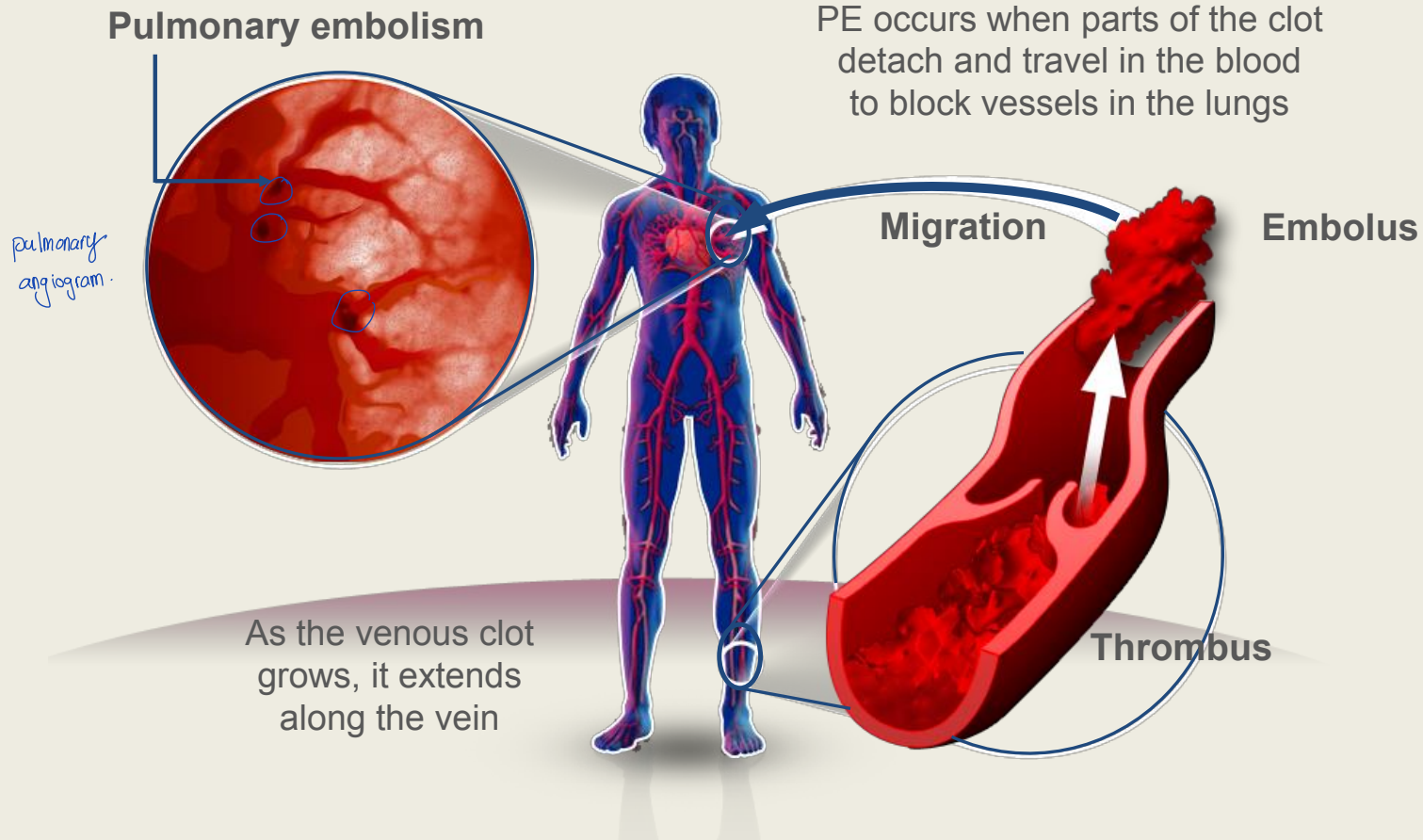
Source of emboli

Thrombotic non thrombotic

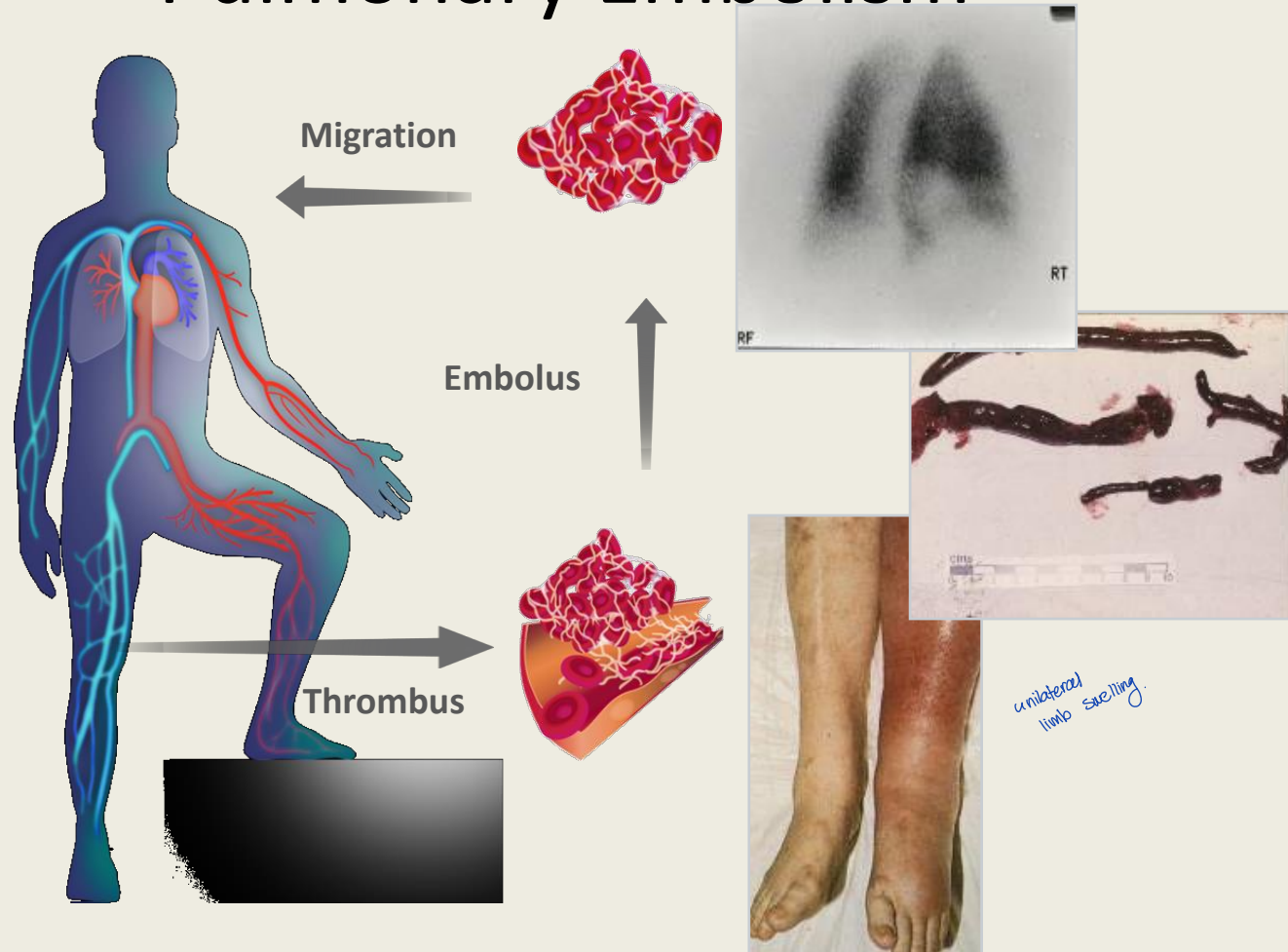
- **Thrombotic**


- Most cases (80–95 percent) as a result of thrombus **originating** in the lower extremity
- Most thrombi originate in the deep veins of the calf and propagate proximally to the popliteal and femoral veins. *(iliofemoral + deep veins of the pelvis)*
- Calf-limited(**below knee**) thrombi pose a minimal embolic risk
- Emboli may also originate from **atypical sites** such as upper-extremity thrombosis associated with central venous catheters or *IV abusers*, intravascular cardiac 2*devices, or may be associated with thoracic outlet obstruction or effort thrombosis *(Tumor)*

VTE: Deep Vein Thrombosis and Pulmonary Embolism



VTE: Deep Vein Thrombosis and Pulmonary Embolism





**Non
thrombotic**

BOX 61-2 Causes of Nonthrombotic Pulmonary Emboli

Fat Embolism

Amniotic fluid embolism

Air Embolism

- Venous

- Arterial

Tumor embolism

Septic pulmonary embolism

(5% only)

Table 3 Predisposing factors for VTE (1)

Strong risk factors (OR >10)
Fracture of lower limb
Previous VTE
Spinal cord injury
Hospitalization for heart failure or atrial fibrillation/flutter (within previous 3 months)
Hip or knee replacement
Major trauma
Myocardial infarction (within previous 3 months) imp

@ESC

VTE = venous thromboembolism.

Table 3 Predisposing factors for VTE (2)

Moderate risk factors (OR 2–9)

Arthroscopic knee surgery

Autoimmune diseases

Blood transfusion

Central venous lines

Intravenous catheters and leads

Chemotherapy

Congestive heart failure or respiratory failure

Erythropoiesis-stimulating agents

Hormone replacement therapy (depends on formulation)

VTE = venous thromboembolism.

Table 3 Predisposing factors for VTE (3)

Moderate risk factors (cont'd)

In vitro fertilization

Oral contraceptive therapy

Postpartum period

Infection (specifically pneumonia, urinary tract infection, and HIV)

Inflammatory bowel disease

Cancer (highest risk in metastatic disease)

Paralytic stroke

Superficial vein thrombosis

Thrombophilia

VTE = venous thromboembolism.

Table 3 Predisposing factors for VTE (4)

Weak risk factors (OR <2)

Bed rest >3 days

Diabetes mellitus

Arterial hypertension

Immobility due to sitting (e.g. prolonged car or air travel)

Increasing age

Laparoscopic surgery (e.g. cholecystectomy)

Obesity

Pregnancy

Varicose veins

VTE = venous thromboembolism.

→ could be congenital or acquired

TABLE 61-1 Inherited Thrombophilias

Disorder	Prevalence (%)		Inheritance	Relative Risk	Clinical Features
	General Population	Patients with VTE			
AT deficiency (most risky)	0.2	1–3	AD	20	VTE, heparin resistance
Protein C deficiency	0.2–0.4	3–5	AD	10	VTE
Protein S deficiency	0.03–0.1	1–5	AD	10	VTE and ATE
Factor V Leiden MC	5	10–50	AD	5	VTE and ATE
Prothrombin G20210A	2–5	6–18	AD	3	VTE
Hyperhomocysteinemia	5	10	Not known	3	VTE and premature ASCVD
Elevated factor VIII	11	25	Not known	5	VTE

(Data from Franchini M, Veneri D, Salvagno GL, et al: Inherited thrombophilia. Crit Rev Clin Lab Sci 2006;43:249–290.)

AT, Antithrombin; VTE, venous thromboembolism, ATE, arterial thromboembolism, ASCVD, atherosclerotic cardiovascular disease.

* presentation of PE is variable.

Once detached from their point of origin, emboli travel via the systemic venous system, through the right chambers of the heart, and **eventually reach the pulmonary arterial system.**

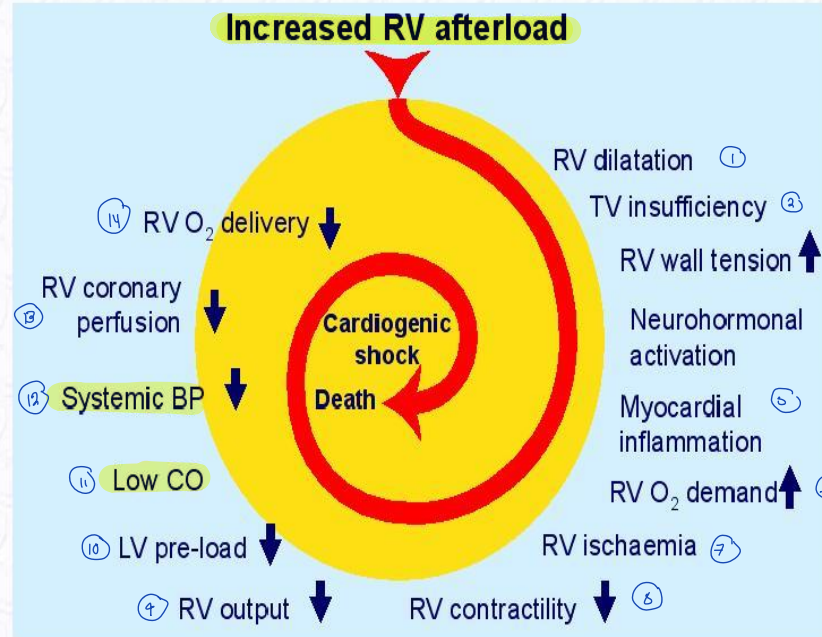
Physiologic effects and clinical consequences of pulmonary thromboembolism **vary widely**, ranging from asymptomatic disease to hemodynamic collapse and death

- Major factors that determine the **outcome** include:

- ☐ Size and location of emboli
- ☐ Coexisting cardiopulmonary **diseases**
- ☐ Secondary humoral mediator release and vascular hypoxic **responses**
- ☐ The rate of **resolution** of emboli.

Hemodynamic consequences

Key factors contributing to haemodynamic collapse in acute pulmonary embolism



*once you have PE with hemodynamic instability = massive PE.

BP = blood pressure; CO = cardiac output; LV = left ventricular; RV = right ventricular; TV = tricuspid valve.

Gas exchange abnormality

- Gas exchange abnormalities...
 - Hypoxemia
 - Increased A–a gradient.
 - V/Q mismatch.
 - Increased dead space
 - Respiratory alkalosis from hyperventilation
- Often a sign of increased dead space and impaired minute ventilation
- *may suggest massive PE*

S&S

TABLE 61-3 Frequency of Signs and Symptoms in Acute Pulmonary Embolism

Symptoms	Frequency (%)
Dyspnea	73
Pleuritic chest pain	66
Cough	37
Leg swelling	33
Hemoptysis	13
Wheezing	9
Chest pain	4
Signs	
Respiratory rate $\geq 20/\text{min}$	70
Crackles	51
Heart rate $\geq 100/\text{min}$	30
Third or fourth heart sound	26
Loud pulmonary component of second heart sound	23
Temperature $> 38.5^\circ \text{C}$	7
Pleural rub	3

loud P2

, you expect low grade fever but this doesn't rule out PE.

dullness on percussion

TABLE 61-2 Differential Diagnosis of Acute Pulmonary Embolism

Pneumonia or bronchitis	Rib fracture
Asthma or exacerbation of chronic obstructive lung disease	Pulmonary edema/ Congestive heart failure
Pleuritis	Thoracic malignancy
Pericarditis/Cardiac tamponade	Pulmonary hypertension
Pneumothorax	Myocardial infarction
Musculoskeletal pain	Aortic dissection
Costochondritis	Anxiety

Management approach

- Assess clinical probability
- Assess risk of mortality
- Investigation
 - Diagnostic
 - Non diagnostic (helpful test)
- Treatment (medications and duration of treatment)

Assessment of pre-test probability

Clinical prediction rules for pulmonary embolism

	Clinical decision rule points	
Wells rule	Original version	Simplified version
Previous PE or DVT	1.5	1
Heart rate ≥ 100 b.p.m.	1.5	1
Surgery or immobilization within the past 4 weeks	1.5	1
Haemoptysis	1	1
Active cancer	1	1
Clinical signs of DVT	3	1
Alternative diagnosis less likely than PE	3	1
Clinical probability		
<i>Three-level score</i>		
Low	0-1	N/A
Intermediate	2-6	N/A
High	≥ 7	N/A
<i>Two-level score</i>		
PE unlikely	0-4	0-1
PE likely	≥ 5	≥ 2

Regardless
of the chr.

Assessment of pre-test probability (cont'd)

Clinical prediction rules for pulmonary embolism (cont.)		
	Clinical decision rule points	
Revised Geneva score	Original version	Simplified version
Previous DVT or PE	3	1
Heart rate		
75-94 b.p.m.	3	1
≥95 b.p.m.	5	2
Surgery or fracture within the past month	2	1
Haemoptysis	2	1
Active cancer	2	1
Unilateral lower limb pain	3	1
Pain on lower limb deep venous palpation and unilateral oedema	4	1
Age >65 years	1	1
Clinical probability		
<i>Three-level score</i>		
Low	0-3	0-1
Intermediate	4-10	2-4
High	≥11	≥5
<i>Two-level score</i>		
PE unlikely	0-5	0-2
PE likely	≥6	≥3

Assess risk of mortality

- **High Risk:**
 - Hemodynamically Unstable.
 - Early mortality is 15%.
- **Non-High Risk**
(According to RVD and Myocardial injury)
 - Intermediate Risk
 - Low Risk

Table 4 Definition of haemodynamic instability

(1) Cardiac arrest	(2) Obstructive shock	(3) Persistent <u>hypotension</u>
Need for cardiopulmonary resuscitation	Systolic BP <90 mmHg, or vasopressors required to achieve a BP ≥90 mmHg despite adequate filling status <i>if you stop them he would collapse.</i>	Systolic BP <90 mmHg, or systolic BP drop <u>≥40</u> mmHg, either lasting longer than 15 minutes and not caused by new-onset arrhythmia, hypovolaemia, or sepsis <i>إذا لم يرفع عنده Hypovolemia وتسمى على ج fluids This is not هذها hypovolemia.</i>
	And	
	End-organ hypoperfusion (altered mental status; cold, clammy skin; oliguria/anuria; increased serum lactate)	

Table 8 Original and simplified PESI (1)

→ Severity index
بعض اعراض العيوب بدرجة الأرقام

Parameter	Original version	Simplified version
Age	Age in years	1point (if age >80 years)
Male sex	+10 points	—
Cancer	+30 points	1point
Chronic heart failure	+10 points	1point
Chronic pulmonary disease	+10 points	
Pulse rate ≥ 110 b.p.m.	+20 points	1point
Systolic BP <100 mmHg	+30 points	1point

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BP = blood pressure; PESI = Pulmonary Embolism Severity Index.

Table 8 Original and simplified PESI (2)

Parameter	Original version	Simplified version
Respiratory rate >30 breaths per min	+20 points	—
Temperature <36 °C	+20 points	—
Altered mental status	+60 points	—
Arterial oxyhaemoglobin saturation <90%	+20 points	1point

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PESI = Pulmonary Embolism Severity Index.

Table 8 Original and simplified PESI (3)

	Risk strata	
كل ما زاد الـ ٥ نقاط كل ما كان اسوأ	Class I: ≤65 points very low 30-day mortality risk (0–1.6%) Class II: 66–85 points low mortality risk (1.7–3.5%)	0 points = 30-day mortality risk 1.0% (95% CI 0.0–2.1%)
	Class III: 86–105 points moderate mortality risk (3.2–7.1%) Class IV: 106–125 points high mortality risk (4.0–11.4%) Class V: >125 points very high mortality risk (10.0–24.5%)	≥1 point(s) = 30-day mortality risk 10.9% (95% CI 8.5–13.2%)

PESI = Pulmonary Embolism Severity Index.

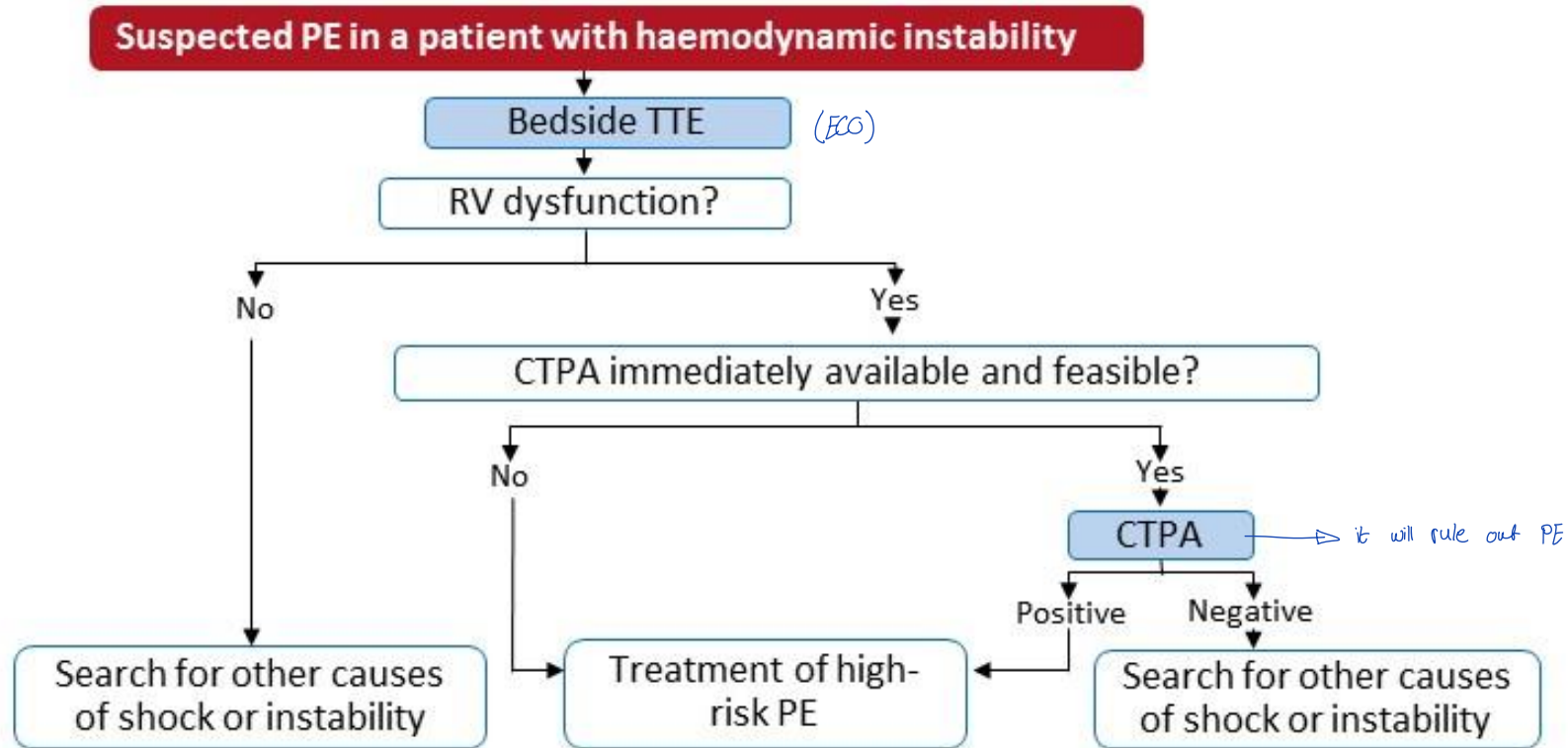
Table 9 Classification of PE based on early mortality risk

Early mortality risk		Indicators of risk			
		Haemo- dynamic instability	Clinical parameters of PE severity/ comorbidity: PESI III–V or sPESI ≥ 1	RV dysfunction on TTE or CTPA <i>(enlarged RV)</i>	Elevated cardiac troponin levels
<i>admit to ICU.</i> Intermediate	High	<i>Massive PE 30% death</i> +	(+)	+	(+)
	Intermediate-high	-	+	+	+
	Intermediate-low	-	+	One (or none) positive	
	Low <i>I can discharge the patient safely on medications.</i>	-	-	-	Assessment optional; if assessed, negative

CTPA = computed tomography pulmonary angiography; PESI = Pulmonary Embolism Severity Index; TTE = transthoracic echocardiography.

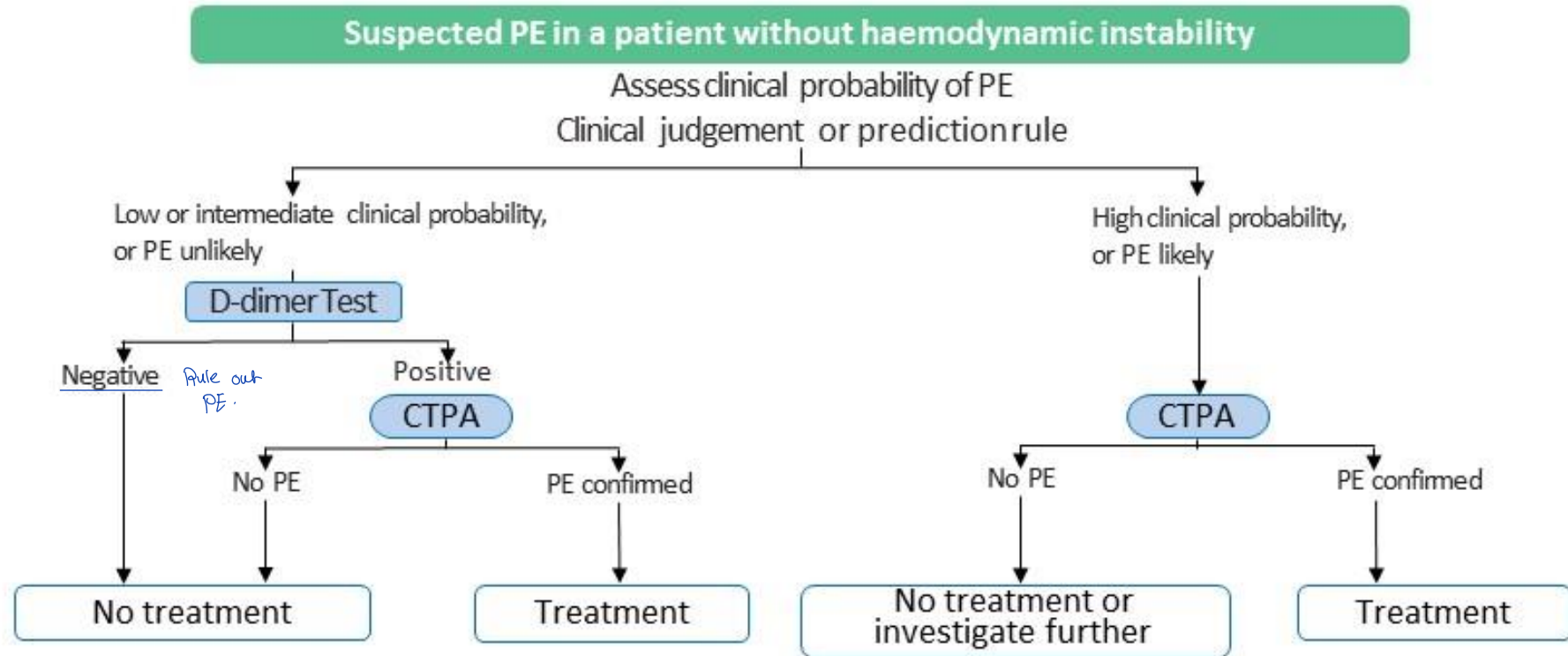
Figure 3 Diagnostic algorithm for suspected high-risk PE

hemodynamic instability.



CTPA = computed tomography pulmonary angiography; RV = right ventricular; TTE = transthoracic echocardiography

Figure 4 Diagnostic algorithm for suspected PE without haemodynamic instability



CTPA = computed tomography pulmonary angiography

Diagnostic investigation

- **D-dimer** for unlikely PE.
 - Non specific measure of fibrinolysis
 - High sensitivity (positive in presence of dx)
 - High negative predictive value (dx is absent when test is negative) in the outpatient setting
 - Useful in outpatient setting/emergency room, not an inpatient test for ruling out PE
 - Need to be adjusted with age the optimal age-adjusted cut-off was defined as the patient's age multiplied by 10 in patients aged ≥ 50 years.

- Currently reserved for

→ you have to have normal CXR.
due to less sensitivity with abnormal CXR.

V/Q scan

can be equal to CTPA.

- Renal impairment
- IV contrast allergies
- Pregnancy
- Hospital resources

Washout
Stage.

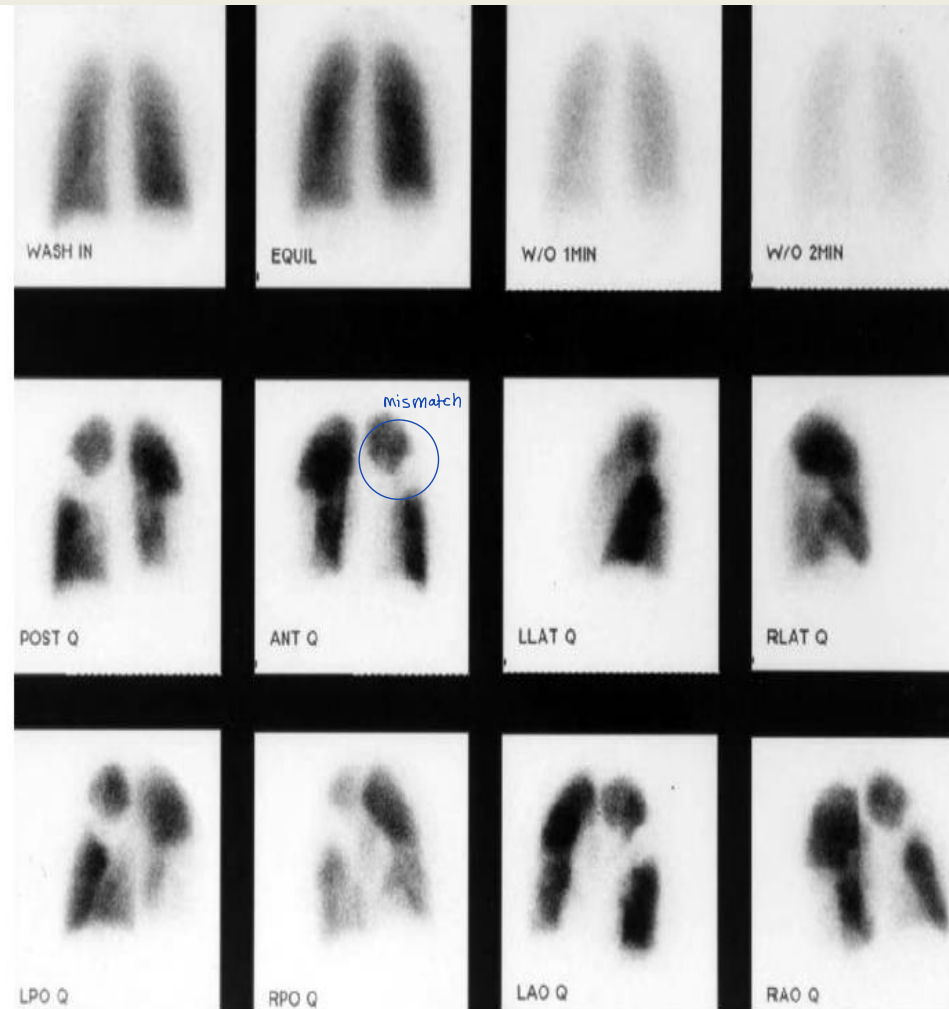


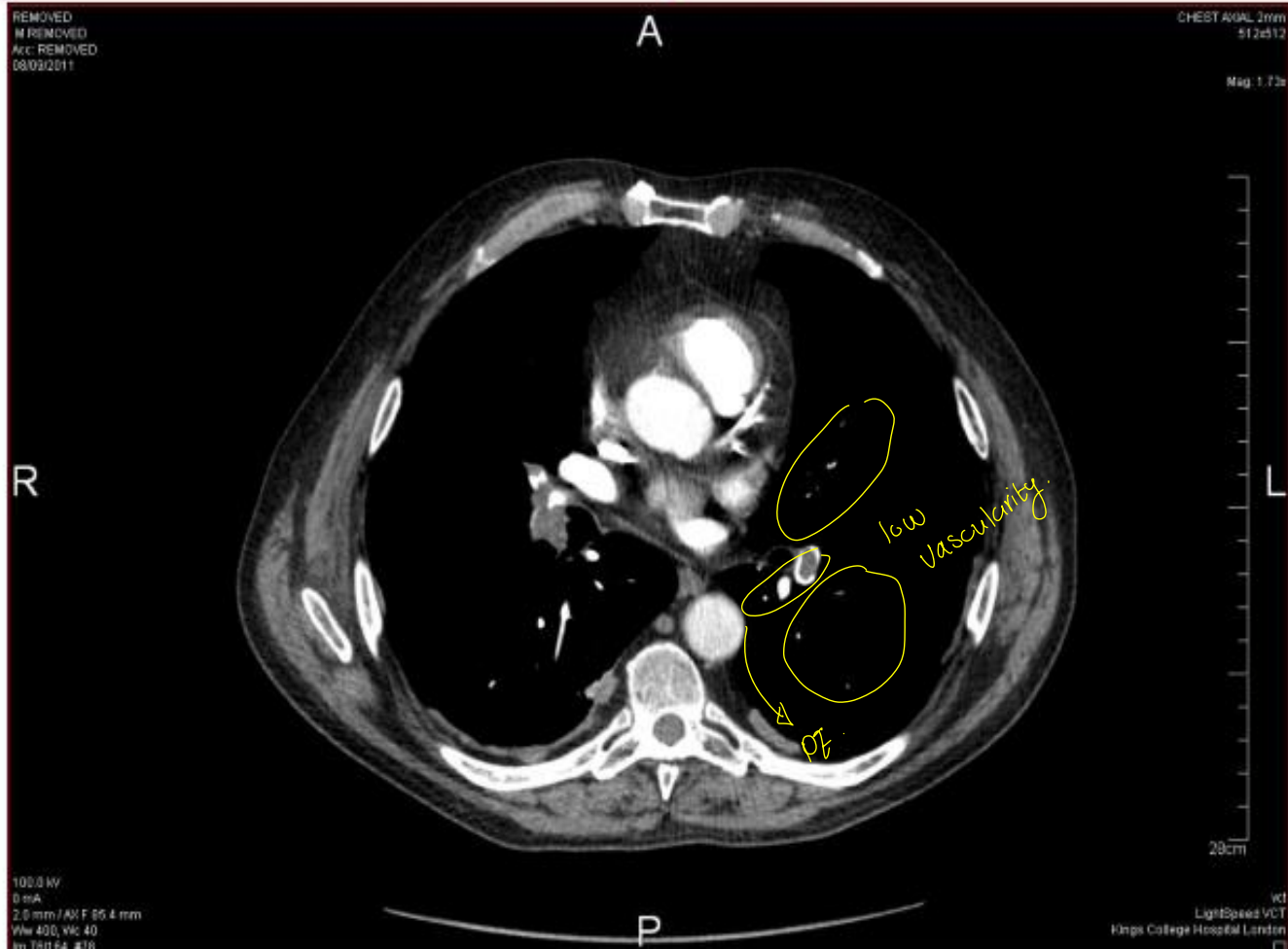
Figure 82-9 "High probability" ventilation/perfusion scan demonstrating normal ventilation and multiple mismatched segmental and larger defects.

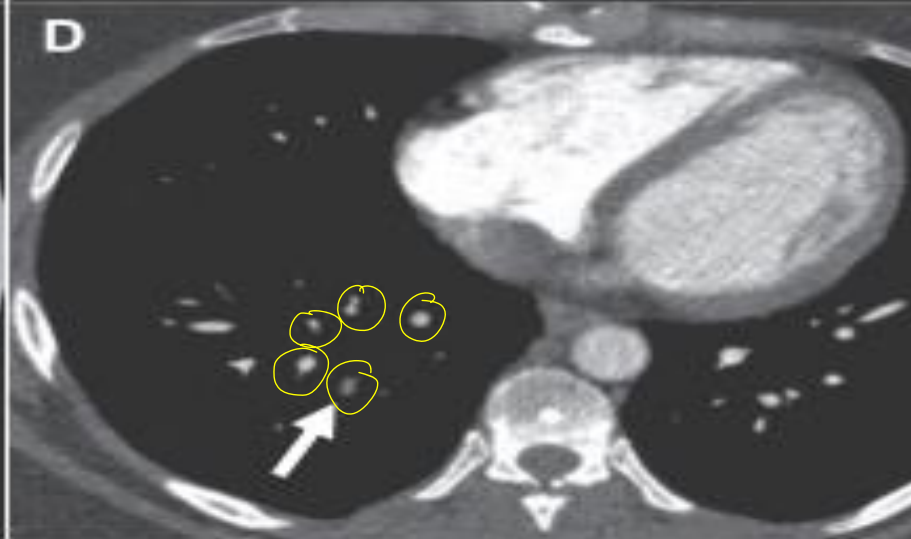
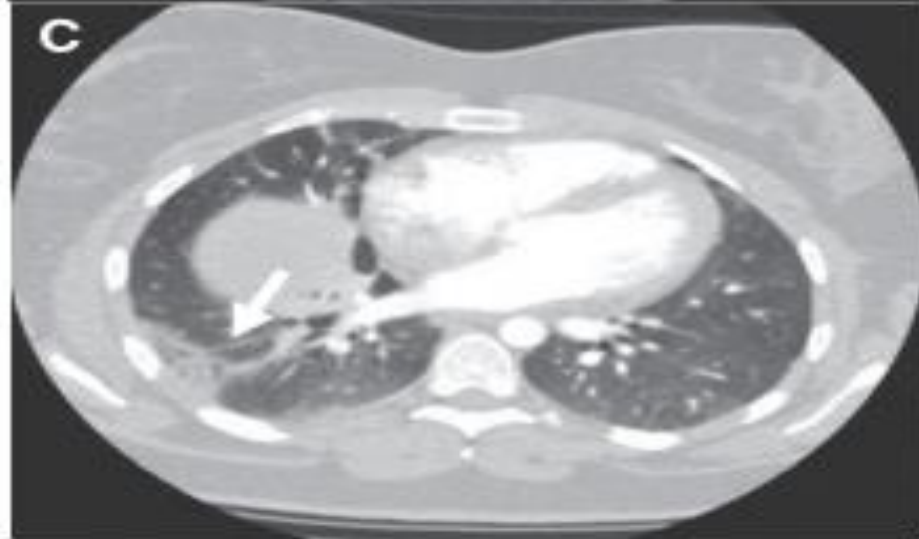
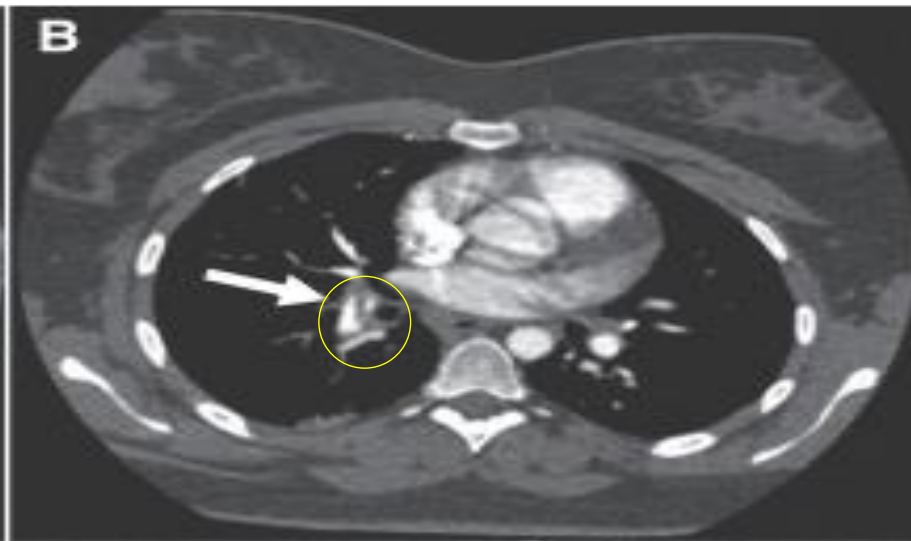
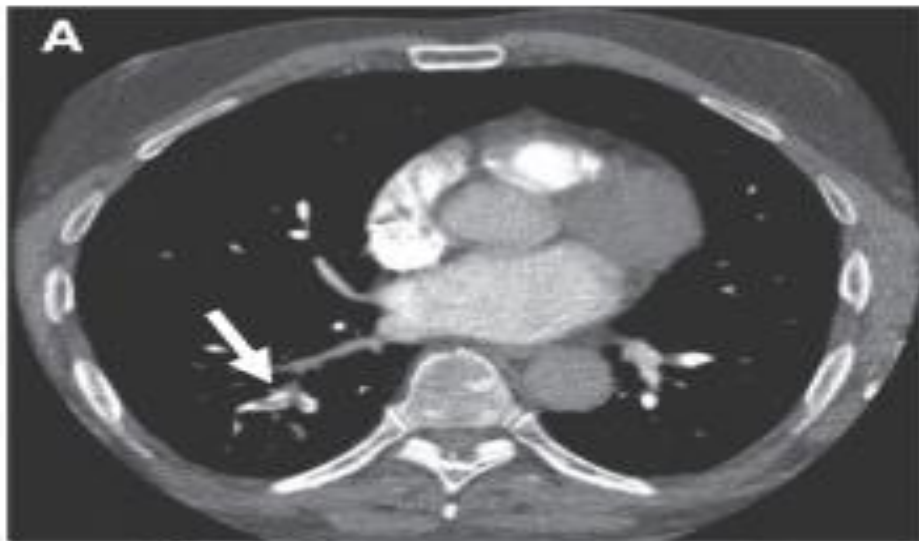
CT with PE protocol

Spiral CT (CTPA) .

- Larger dose of Contrast
 - Rapid rate(timed) of contrast
 - Effective dose at pulmonary CT angiography, without significant loss of objective or subjective image quality.
- } characteristics .

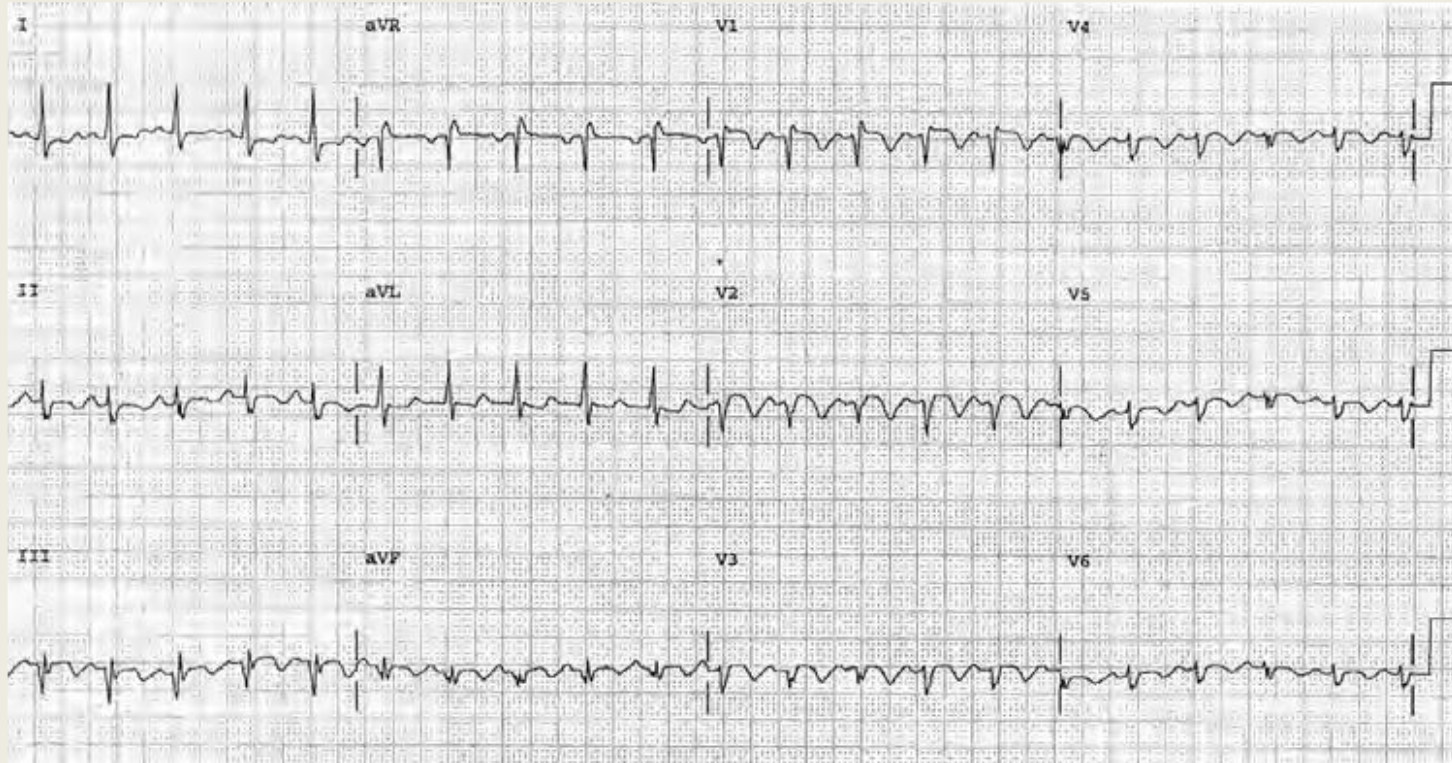
Pulmonary Emboli





Electrocardiogram demonstrating findings consistent with embolism including sinus tachycardia, incomplete right bundle branch block, **S1Q3T3 pattern**, and **inverted precordial T waves**.(minority of patients)

MC finding on ECG:
Sinus tachy



Others

- **CXR:**

Most patients with pulmonary embolism have abnormal

but nonspecific chest radiographic findings

- **Echocardiogram:**

Suspected massive pulmonary embolism who are too ill for transportation or

have an absolute contraindication to the administration of a contrast agent.

- **Troponin :**

Increase in right heart strain .

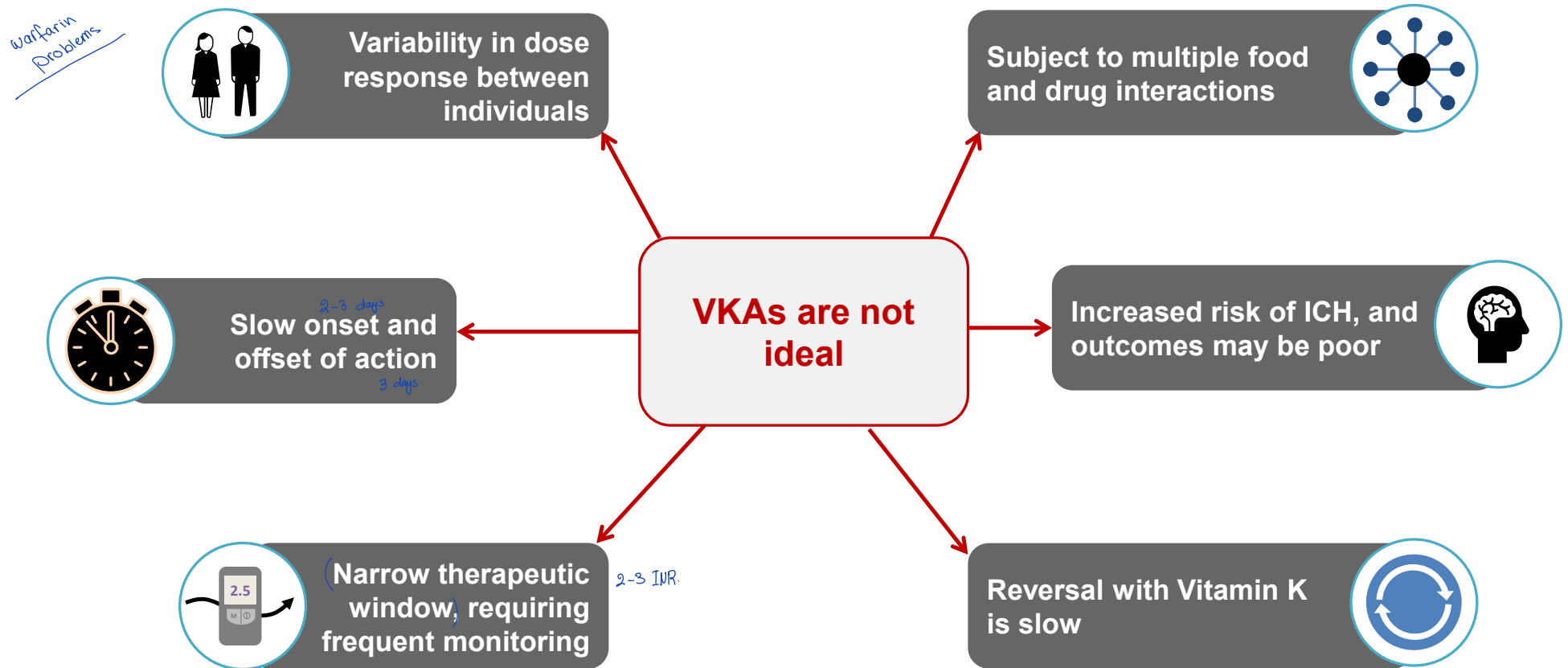
Treatment

Three phases: **Initiation phase** (5-21 days): The initial provision of anticoagulants following VTE diagnosis, **Treatment phase** (3 months): The period after initiation that completes treatment for the acute VTE event, **Extended phase** (3 months-no planned stop date): The period of anticoagulant use at full or reduced dose for the goal of secondary prevention

- Most recent 2016 ATS guidelines, ECS 2019 , ASH 2020 , 2021 ATS update **NOACs** become the recommended oral treatment and alternative is VKAs(warfarin).
- ECS2019 recommends decrease dose of NOACs after 3-6months of treatment while ASH you can use full or reduced dose of NOACs. ATS 2021 update recommends decrease the dose of NOACs after 3 months of treatment(treatment phase)

● In Summary : Main: NOAC , 2nd: warfarin in initial treatment low molecular weight heparin. on long treatment depends on the cause.

VKAs were the cornerstone of anticoagulation but they have multiple limitations



ICH, intracranial hemorrhage

Turpie. Eur Heart J 2008;29:155; Khoo et al. Int J Clin Pract 2009;63:630; Fang et al. Stroke 2012;43:1795; Hanley et al. J Clin Pathol 2004;57:1132

TABLE 1 Non-Vitamin K-Dependent Oral Anticoagulant Agents in the Treatment and Secondary Prevention of VTE

	Dosage and Interval			Not Recommended or Contraindicated*
	Initial Phase	Long-Term Phase	Extended Phase	
Rivaroxaban†	15 mg twice daily with food for 21 days	20 mg once daily with food		<ul style="list-style-type: none"> • CrCl <30 ml/min • Moderate or severe hepatic impairment (Child-Pugh B and C), or hepatic disease associated with coagulopathy • Concomitant use of combined P-gp and strong CYP3A4 inhibitors or inducers
Dabigatran etexilate‡	Initial therapy with parenteral anticoagulation for 5–10 days should precede administration of dabigatran etexilate	150 mg twice daily		<ul style="list-style-type: none"> • CrCl <30 ml/min • Concomitant treatment with P-gp inhibitors in patients with CrCl <50 ml/min • Concomitant treatment with P-gp inducers (i.e., rifampin)
Apixaban	10 mg twice daily for 7 days	5 mg twice daily	2.5 mg twice daily after at least 6 months of treatment	<ul style="list-style-type: none"> • CrCl <15 ml/min • Severe hepatic impairment (Child-Pugh C), or hepatic disease associated with coagulopathy • Strong dual inhibitors or inducers of CYP3A4 and P-gp
Edoxaban§	Initial therapy with parenteral anticoagulation for 5–10 days should precede administration of edoxaban	60 mg once daily 30 mg once daily can be considered in patients with ≥1 of the following factors: CrCl 15–50 ml/min; body weight ≤60 kg; concomitant use of P-gp inhibitors, cyclosporin, dronedarone, erythromycin, or ketoconazole		<ul style="list-style-type: none"> • CrCl <15 ml/min • Moderate or severe hepatic impairment (Child-Pugh B and C), or hepatic disease associated with coagulopathy • Concomitant treatment with rifampin

NOACs are not used

- **CKD** with Ccl less than 30 ml/min (apixaban & edoxaban can be used Ccl bet.15-30 ml/min with reduced dose)
- Moderate to severe **hepatic** impairment *bc they are secreted by the liver.*
- Pregnancy and lactation:
Still the use of LMWH is the standard of care in pregnant lady and VKAs can be used in lactating women
- PE with cancer :
LMWH
- Antiphospholipid syndrome :
not proved yet

Risk Factors Associated with VTE Recurrence

(needs anti-coagulations more than
3 months): "chronic PE"

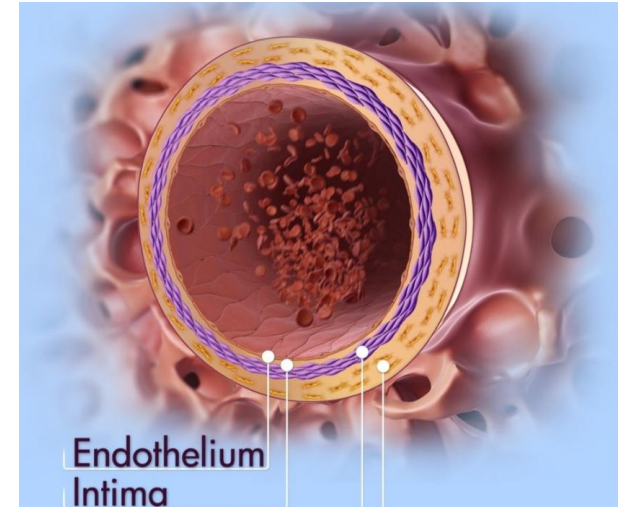
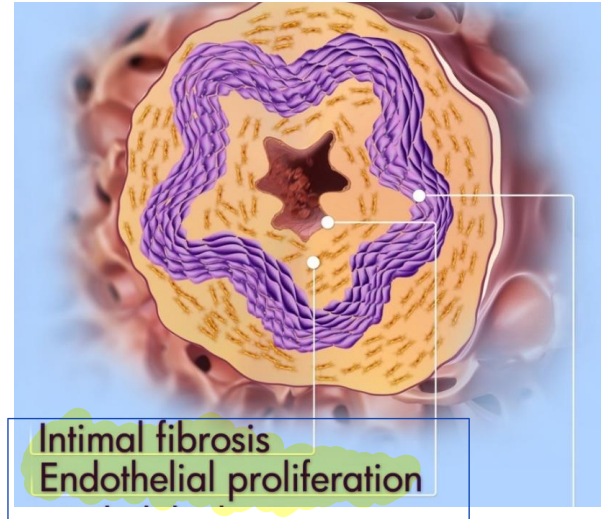
- Idiopathic presentation^{1,2}
- Presentation of primary DVT¹
- Increasing age¹
- Proximal DVT²
- Cancer²
- Residual thrombus mass³
- Male gender⁴
- High-risk thrombophilia¹
- Scoring systems such as the DASH score, the Vienna Prediction Model, and the 'Men continue and HERDOO2' score are used to predict PE.

Pulmonary hypertension

Haemodynamic definitions of pulmonary hypertension

Definition	Haemodynamic characteristics
PH	mPAP >20 mmHg <i>previously we used to say 25.</i>
(Pre-capillary PH) <i>To diagnose it you need right heart cath.</i>	mPAP >20 mmHg PAWP ^{widge pressure.} ≤15 mmHg <i>→ left atrial pressure.</i> PVR ^{Resistance.} >2 WU
Isolated post-capillary PH	mPAP >20 mmHg PAWP <u>>15</u> mmHg PVR ≤2 WU <i>(عكس فوف)</i>
Combined post- and pre-capillary PH	mPAP >20 mmHg PAWP >15 mmHg PVR >2 WU
Exercise PH	mPAP/CO slope between rest and exercise >3 mmHg/L/min

**Pulmonary
remodelling
in PAH leads
to narrowing
of the artery¹**



- PAH, pulmonary arterial hypertension; SMC, smooth muscle cell.
- 1. Figure adapted from Galiè N, *et al. Eur Heart J* 2010; 31:2080-6.

Clinical classification of pulmonary hypertension (1) *(2nd ML)*

GROUP 1 Pulmonary arterial hypertension (PAH)

1.1 Idiopathic

1.1.1 Non-responders at vasoreactivity testing

1.1.2 Acute responders at vasoreactivity testing

1.2 Heritable

1.3 Associated with drugs and toxins

1.4 Associated with:

1.4.1 Connective tissue disease *↳ Scleroderma, RA*

1.4.2 HIV infection

1.4.3 Portal hypertension

1.4.4 Congenital heart disease

1.4.5 Schistosomiasis *(endemic hypertension)*

1.5 PAH with features of venous/capillary (PVOD/PCH) involvement

1.6 Persistent PH of the newborn

Clinical classification of pulmonary hypertension (2) (Mc class)

GROUP 2 PH associated with left heart disease

2.1 Heart failure:

2.1.1 with preserved ejection fraction

2.1.2 with reduced or mildly reduced ejection fraction

2.2 Valvular heart disease

2.3 Congenital/acquired cardiovascular conditions leading to post-capillary PH

GROUP 3 PH associated with lung diseases and/or hypoxia

3.1 Obstructive lung disease or emphysema

3.2 Restrictive lung disease

3.3 Lung disease with mixed restrictive/obstructive pattern

3.4 Hypoventilation syndromes

3.5 Hypoxia without lung disease (e.g. high altitude)

3.6 Developmental lung disorders

Clinical classification of pulmonary hypertension (3)

GROUP 4 PH associated with pulmonary artery obstructions

4.1 Chronic thrombo-embolic PH

4.2 Other pulmonary artery obstructions

GROUP 5 PH with unclear and/or multi-factorial mechanisms

5.1 Haematological disorders

5.2 Systemic disorders

5.3 Metabolic disorders

5.4 Chronic renal failure with or without haemodialysis

5.5 Pulmonary tumour thrombotic microangiopathy

5.6 Fibrosing mediastinitis

Figure 1
Central illustration

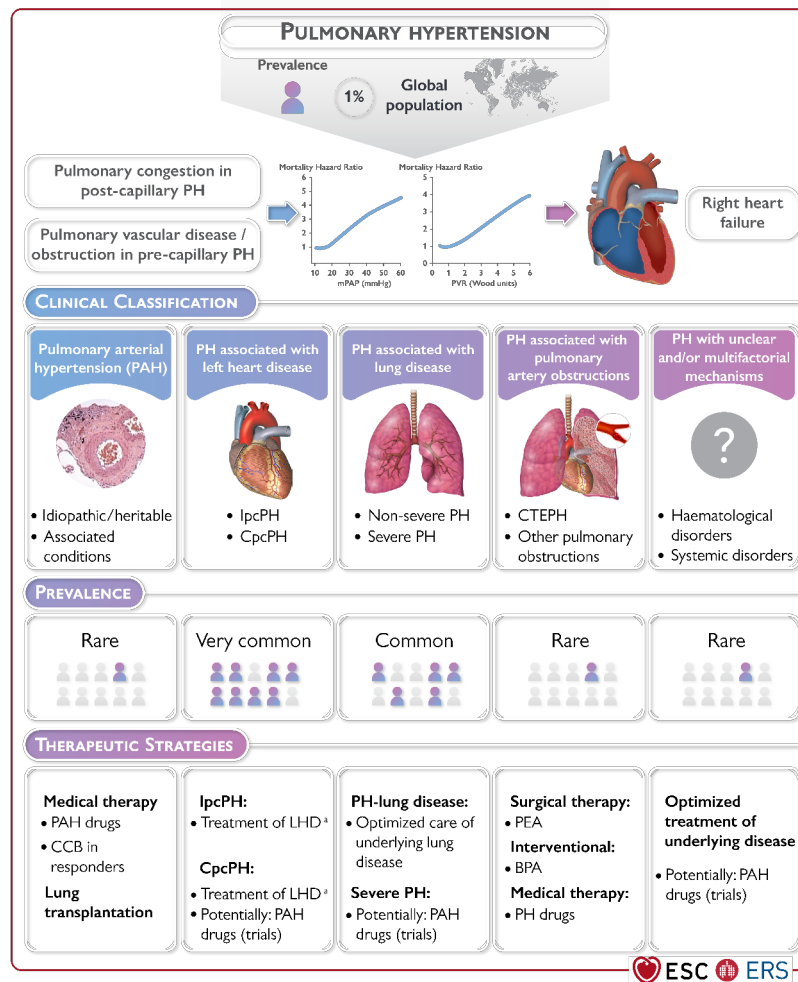


Figure 2

Symptoms in patients with pulmonary hypertension

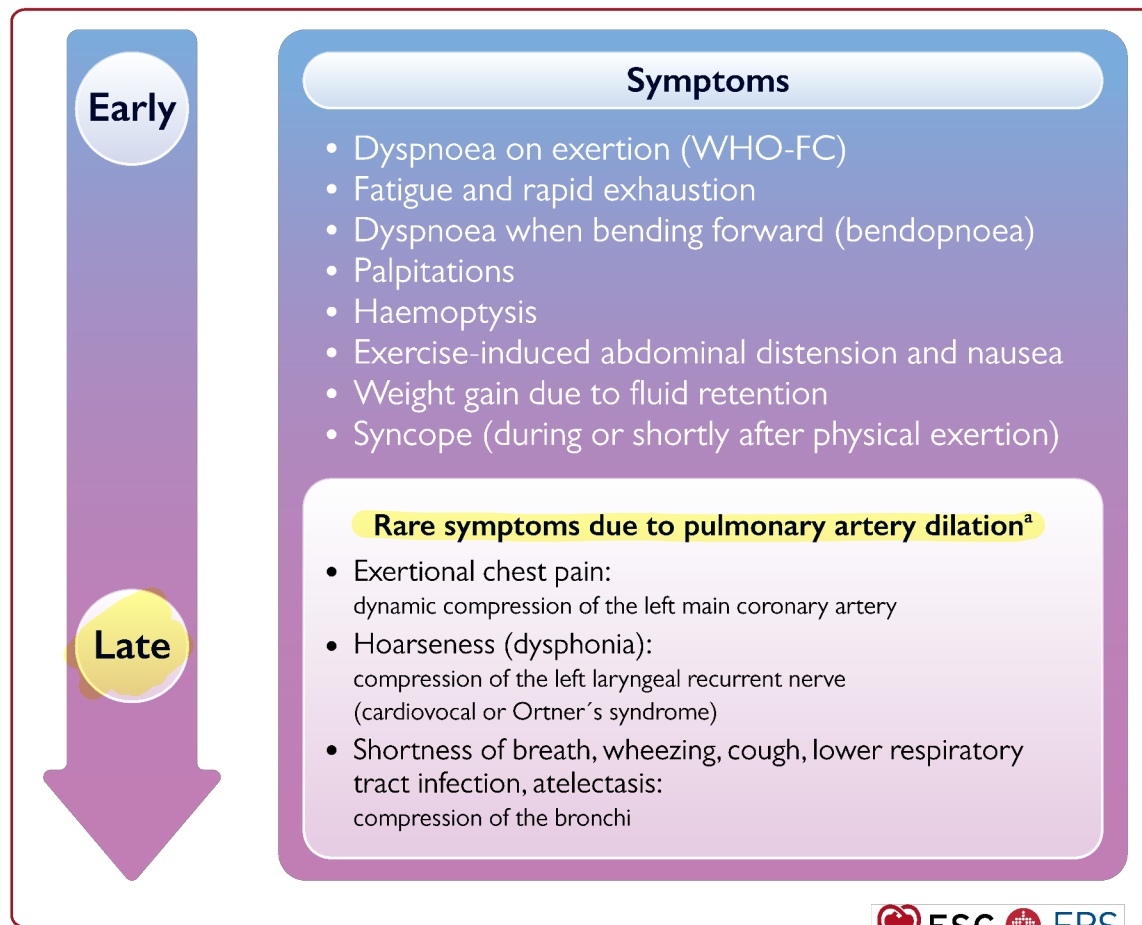


Figure 3

Clinical signs in patients with pulmonary hypertension

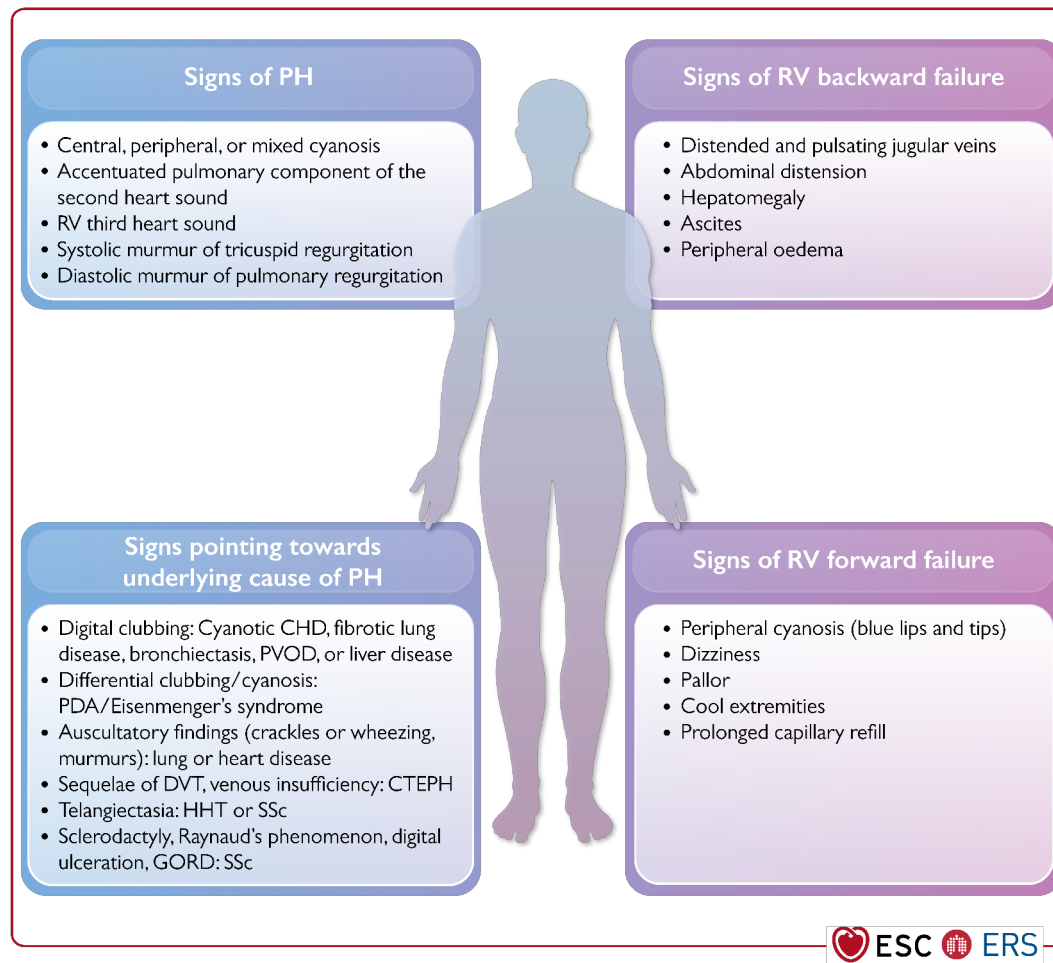
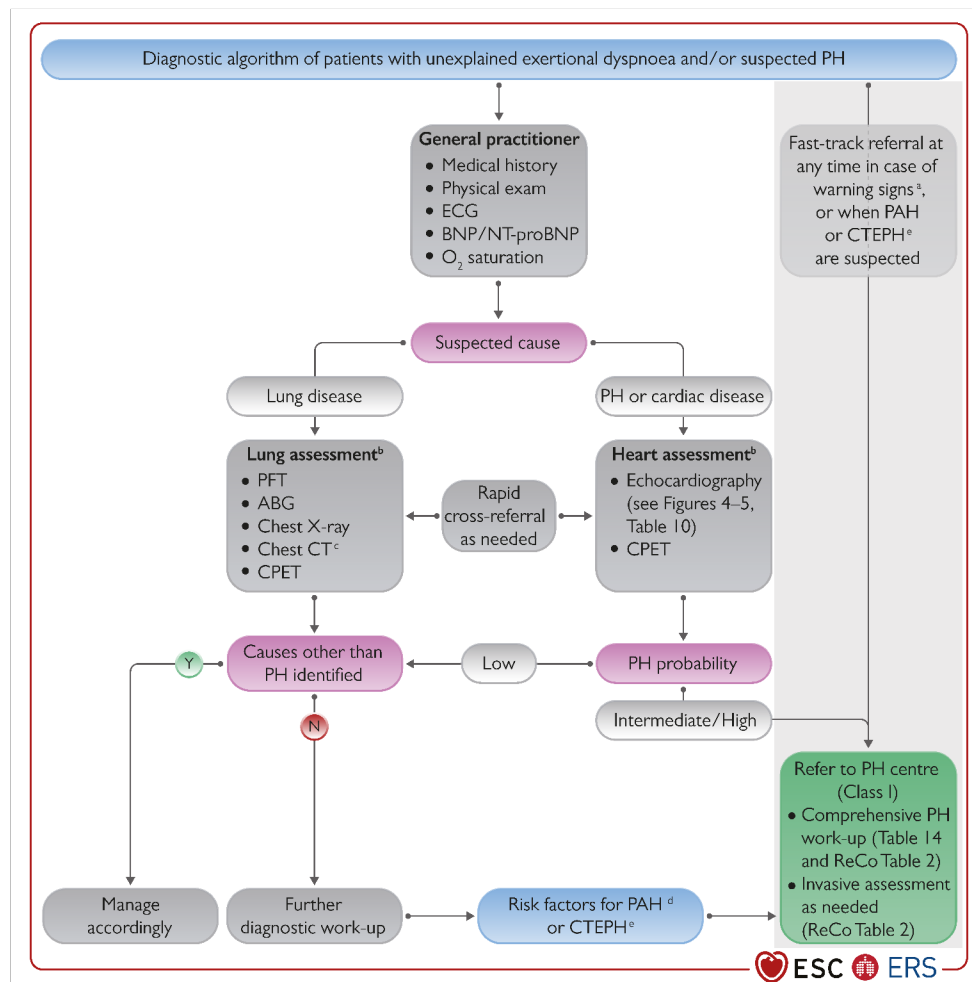


Figure 6

Diagnostic algorithm of patients with unexplained dyspnoea and/or suspected pulmonary hypertension

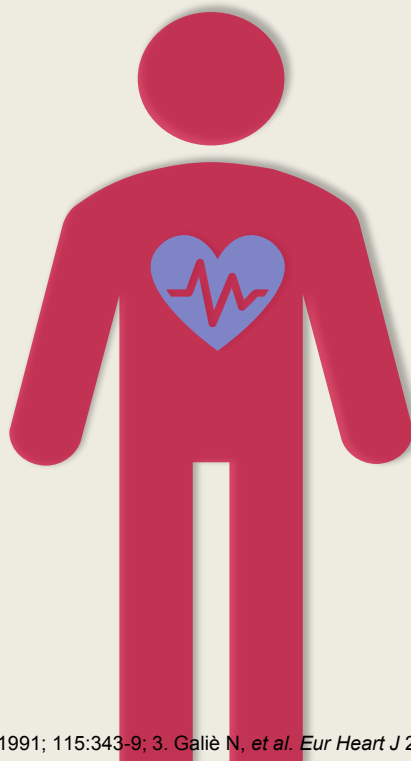


- **PAH is a rare, progressive disease with poor prognosis if left untreated**

Prevalence:
6.6 to 26 per million¹

Poor short- and
medium-term prognosis²

Severe and progressive²



Incidence:
1.1 to 7.6 per million¹

Complex management³

Estimated median survival of
2.8 years if left untreated²



Thank you