Connective tissue disease-associated pulmonary arterial hypertension

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Abstract

Although rare in its idiopathic form, pulmonary arterial hypertension (PAH) is not uncommon in association with various associated medical conditions, most notably connective tissue disease (CTD). In particular, it develops in approximately 10% of patients with systemic sclerosis and so these patients are increasingly screened to enable early detection. The response of patients with systemic sclerosis to PAH-specific therapy appears to be worse than in other forms of PAH. Survival in systemic sclerosis-associated PAH is inferior to that observed in idiopathic PAH. Potential reasons for this include differences in age, the nature of the underlying pulmonary vasculopathy and the ability of the right ventricle to cope with increased afterload between patients with systemic sclerosis-associated PAH and idiopathic PAH, while coexisting cardiac and pulmonary disease is common in systemic sclerosis-associated PAH. Other forms of connective tissue-associated PAH have been less well studied, however PAH associated with systemic lupus erythematosus (SLE) has a better prognosis than systemic sclerosis-associated PAH and likely responds to immunosuppression.

Introduction

Pulmonary hypertension is defined as a mean pulmonary arterial pressure (mPAP) of ≥25 mmHg when measured at right heart catheterization [1]. Current classification describes five main groups with shared clinical and pathophysiological characteristics (Figure 1): group 1, PAH; group 2, pulmonary hypertension associated with left heart disease; group 3, pulmonary hypertension associated with lung disease; group 4, chronic thromboembolic pulmonary hypertension (CTEPH); and group 5, miscellaneous with unclear mechanisms [2]. Although PAH is rare in the general population (idiopathic PAH having an incidence of 1–2/million/year) [3] it is more common in several associated conditions, most noticeably CTD. As a consequence of the high prevalence of both left heart disease and interstitial lung disease in CTD, the accurate diagnosis of PAH is particularly challenging. Although PAH in other forms of CTD (CTD-PAH) will be briefly discussed, PAH is most commonly seen in association with systemic sclerosis and is the main focus of this review. In particular the differences between systemic sclerosis with PAH and idiopathic PAH will be explored.

Systemic sclerosis

Epidemiology

The prevalence of systemic sclerosis within the general population ranges from 80/million in the UK [4] to 240/million in the USA [5]. Prospective right heart catheter-based studies have observed a prevalence of PAH in patients with systemic sclerosis of 7.8–12% while a recent meta-analysis calculated the prevalence to be 9% [6,7]. The French Itiner-Air group observed an annual incidence of PAH in patients with systemic sclerosis of 0.61% [8].

Pathogenesis

Systemic sclerosis-PAH is characterized by intimal hyperplasia, medial hypertrophy and adventitial fibrosis as in other forms of PAH although, compared to idiopathic
PAH, a lower number of plexiform lesions are observed [9]. Recent histological data suggest a surprisingly high involvement of pulmonary venules, although the proportion of patients with clinically overt pulmonary veno-occlusive disease (PVOD) is lower [7,9,10]. Mutations in bone morphogenetic receptor type 2 (BMPR-2), which are well described in idiopathic and heritable PAH, have not been demonstrated in systemic sclerosis-PAH. An increased frequency of a polymorphism in the gene coding for endoglin, a component of the transforming growth factor beta (TGF-B) receptor complex, which is associated with hereditary haemorrhagic telangectasia and has been identified in patients with idiopathic PAH [11], has been recently identified in patients with systemic sclerosis-PAH [12].

Systemic sclerosis is an autoimmune condition and, as such, it seems likely that autoimmunity and inflammation will play an important role in the development of PAH. In support of this hypothesis, lymphocytes, macrophages and leucocytes have been identified in pulmonary arterial vascular lesions in systemic sclerosis-PAH [10]. The association between several auto-antibodies and the presence of isolated systemic sclerosis are well recognized, including anticentromere, antitopoisoasemase-1 (SCI-70), anti-RNA-polymerase-III and anti-Th/To antibodies. However, their exact role in the pathogenesis of PAH is unclear [13]. Studies in systemic sclerosis patients without PAH have identified high levels of molecules associated with endothelial cell activation and apoptosis (VCAM) and angiogenesis (vascular endothelial growth factor [VEGF]), which would be consistent with the risk of subsequent PAH development [14,15]. Anti-fibroblast IgG from sera of systemic sclerosis patients has been shown to activate normal fibroblasts, and it is hypothesized that fibroblast activation may lead to the induction of collagen synthesis, which may promote subsequent vascular remodeling [16]. Becker et al. have recently reported elevated levels of stimulating anti-endothelin receptor type A antibodies.
and anti-angiotensin receptor type-1 antibodies in systemic sclerosis-PAH and CTD-PAH when compared with idiopathic PAH [17]. Serum levels of both antibodies were prognostic, despite a lack of correlation with pulmonary haemodynamics, and also predicted the development of PAH in a separate cohort of systemic sclerosis patients without PAH at baseline.

Risk factors
The classical systemic sclerosis patient considered to be at high-risk of developing PAH has limited cutaneous disease of >5 years duration, a low lung gas transfer (DLCO) and is anticentromere antibody positive [18–21]. More recent data have challenged some of these suppositions. A recent study of 78 consecutively diagnosed systemic sclerosis-PAH patients reported the onset of PAH <5 years from the onset of limited cutaneous systemic sclerosis in 55% of patients [22]. Similarly, more recent studies have suggested that PAH is almost as common in diffuse disease [17,23]. A reduced and falling gas transfer (DLCO) is common in patients with systemic sclerosis-PAH, with a mean DLCO of 52% predicted 4.5 years prior to the diagnosis of PAH, compared with a mean DLCO of 80% predicted in non-PAH systemic sclerosis patients [19,24]. Interestingly, patients with systemic sclerosis-PAH have lower DLCO than patients with idiopathic PAH, despite having less severe pulmonary haemodynamics at presentation (discussed below) [25]. As well as the recognized risk of PAH in patients with anticentromere antibody, associations between the presence of anti-RNA-polymerase-III and anti-Th/To antibodies and PAH have also been reported [13].

Prognosis
Several large registries have reported survival in systemic sclerosis-PAH over recent years (Table 1): the UK national registry enrolled 259 incident patients and observed 1- and 3-year survival of 78% and 47% [26]; the French Registry enrolled 156 incident patients and observed a 1-year survival of 82% [27]; and the PHAROS registry enrolled 131 incident patients (with a high proportion in World Health Organisation [WHO] Functional Class [FC] I and II) and observed 1- and 3-year survival of 93% and 75% [28]. The different outcomes reported are likely to be impacted by differences in patient characteristics (e.g., in the PHAROS study a high proportion were in WHO FC I and II, reflecting a higher proportion of patients identified by screening). Lefevre et al. recently performed a meta-analysis to adjust for these differences, and observed 1- and 3-year survival of 81% and 52% [29].

Prognostic factors identified at multivariate analysis have also differed between studies, but have included age [26,28], male gender [26,28], DLCO [26,28], functional class [26,28], pulmonary vascular resistance [21], pulmonary capacitance [21], stroke volume index [21] and estimated glomerular filtration rate [21].

It is unclear whether survival in systemic sclerosis-PAH has improved with the introduction of PAH-specific therapies. Historical case series prior to the widespread availability of PAH-specific therapies reported median survival rates of around one year, but are limited by their extremely small size [30]. Rubenfire et al. recently reported no difference in survival in patients diagnosed before 2002 (when only prostacyclin was available) than in patients diagnosed from 2002 onwards (when oral therapies were made available), whereas survival in idiopathic PAH did improve over the same period [31]. Similarly, in their meta-analysis Lefevre et al. also found no relationship between survival and years of enrollment of each study, which may also suggest no progressive improvement in survival over the last decade [29].

Detection
Given the relatively high prevalence of PAH in systemic sclerosis, active screening of patients is advocated [32]. Humbert et al. demonstrated that patients identified with systemic sclerosis-PAH via a screening programme had milder disease and superior survival than those patients who presented with symptomatic disease [33].

Table 1. Prognosis of SSc-PAH in selected major registries

| Registry | Year | n   | Age (yrs) | Incident cases (%) | WHO FC I/II/III/IV (%) | mPAP (mmHg) | PVR (dyn.s.cm5) | 1 yr survival (%) | 3 yr survival (%) |
|----------|------|-----|-----------|-------------------|-----------------------|-------------|----------------|------------------|------------------|
| UK [26]  | 2009 | 259 | 64        | 100               | 16/68/16              | 42          | 715            | 78               | 47               |
| REVEAL [20] | 2010 | 399 | 62        | 18                | 25/60/15             | 45          | 768            | 82               | n/a              |
| ASPIRE [25] | 2012 | 156 | 66        | 100               | 19/67/14             | 43          | 678            | 82               | 52               |
| French [27] | 2013 | 85  | 65        | 100               | 21/67/12             | 41          | 680            | 90               | 56               |
| PHAROS [28] | 2014 | 131 | 60        | 100               | 56/38/6              | 36          | 448            | 93               | 75               |

Age and haemodynamic data presented as mean.
Abbreviations: mPAP, mean pulmonary arterial pressure; PAH, pulmonary arterial hypertension; PVR, pulmonary vascular resistance; SSc, systemic sclerosis; WHO FC, World Health Organisation Functional Class
patients with mildly symptomatic disease have been shown to benefit from PAH therapy, the contribution lead-time bias plays in the superior survival of screened patients is still unclear [34]. Current European Respiratory Society (ERS)/European Society of Cardiology (ESC) guidelines recommend annual screening with echocardiography in patients with systemic sclerosis [32]. Mukerjee et al. demonstrated a DLco of <55% to have high sensitivity for pulmonary hypertension, but no threshold for either echocardiography-derived systolic pulmonary artery pressure or DLco could be identified which could confidently exclude pulmonary hypertension [35]. The recently published multicentre DETECT study enrolled 466 systemic sclerosis patients enriched for an increased risk of PAH (>3 years from diagnosis of systemic sclerosis with DLco <60%) and derived a score based on 6 non-echocardiographic variables (forced vital capacity [FVC] % predicted/DLco % predicted, present/past teleangiectasia, anticitrullinated antibody, N-terminal pro-brain natriuretic peptide (NT-proBNP), serum urate and right axis deviation on electrocardiogram (ECG) to guide whether or not an echocardiogram is subsequently required [36]. A second score based on right atrial area and tricuspid regurgitant velocity is then added to the first score to produce a total score that determines whether right heart catheterisation is required. The DETECT protocol resulted in more patients in this enriched cohort being referred for right heart catheterisation but resulted in less false negatives than an approach using echocardiography alone (4% versus 29%). There are important limitations to the DETECT approach: the protocol has not been validated in a separate cohort, has not been studied in patients <3 years from diagnosis of systemic sclerosis or with DLco >60% and importantly has not been compared with a more holistic approach to identifying patients with systemic sclerosis-PAH currently used in many centers, which incorporates echocardiography, DLco and symptoms to identify patients requiring right heart catheterisation. Furthermore, it is not clear how practical the DETECT algorithm will be in widespread clinical practice, especially in terms of accurate measurement of right atrial area and availability of NT-proBNP assays.

A proportion of patients undergoing will be found to have borderline mean pulmonary arterial pressures of 21–24 mmHg at right heart catheterization. A recent study of 228 patients in the Royal Free Hospital, London, observed progression to PAH in 31% of such patients at subsequent right heart catheterization, suggesting that patients with borderline pressures require careful monitoring [37]. Bae et al. repeated right heart catheterisation in 11/28 patients with the same borderline pulmonary pressures and found that 6/11 had developed resting pulmonary hypertension after a mean period of 14 months [38]. Of note, associated interstitial lung disease and restrictive spirometry was more common in those patients with borderline pressure than in those with normal pressure.

Exercise may have a role in the earlier identification of patients with pulmonary vasculopathy in connective tissue disease, with differential patterns of gas exchange on non-invasive exercise testing distinguishing pulmonary vascular disease from left heart disease in systemic sclerosis [39]. Patients with elevation of mPAP >30 mmHg on exercise appear to be at risk of developing PAH with 19% of patients developing PAH at rest in the UK national CTD-PAH registry [26]. Additional evidence demonstrating improved haemodynamics and exercise capacity in response to the endothelin-1 receptor antagonist, ambrisentan, in patients with exercise-induced systemic sclerosis-pulmonary hypertension further suggests this is a precursor to PAH [40]. The normal response in mPAP at exercise in patients without PAH at rest is not known [41] and so routine exercise at right heart catheterization is not currently recommended [1].

**Treatment**

Current PAH-specific therapies target 3 pathways (prostacyclin, endothelin-1 and nitric oxide) [42]. Several subgroup analyses of CTD-PAH patients enrolled in the pivotal randomized controlled trials (RCTs) have been published which, apart from one exception [43], suggest a less marked response to PAH therapy in CTD-PAH. It is possible that this may be partly due to the fact that the six minute walk distance (6MWD), which has been used as the primary endpoint in the majority of RCTs of PAH-specific therapies, may particularly be affected by age and musculoskeletal pathology in patients with CTD.

**Prostanoids**

An unblinded RCT of continuous intravenous epoprostenol (which is administered by a portable pump via an indwelling central line), involving 111 patients with systemic sclerosis-PAH, observed a significant improvement in mean 6MWD of 46 metres following 12 weeks of treatment, while patients receiving placebo experienced a mean decrease of 48 metres (P<0.05) [43]. Due to its much longer half-life, the prostacyclin analogue treprostinil can be administered via continuous subcutaneous infusion, thus removing the risk of line infection. Ninety patients with CTD-PAH (including 45 patients with systemic sclerosis-PAH) from the original RCT (which also included 270 patients with idiopathic PAH and 109 patients with congenital heart disease) were analysed separately [44]. Unlike in the original study, the between
treatment group’s difference in median 6MWD did not reach statistical significance (+25 metres, \( P=0.06 \)) in the CTD-PAH group. Furthermore, improvements in the secondary endpoints of Borg dyspnoea score, mean right atrial pressure, mean pulmonary artery pressure, and mixed venous oxygen saturation seen in the original study, were not observed in the subgroup analysis.

**Endothelin-1 receptor antagonist**

Bosentan, an endothelin-1 receptor antagonist, was studied in an RCT involving 213 PAH patients \[45\]. In the 150 idiopathic PAH patients, the 6MWD increased by 46 metres after 16 weeks of therapy, while it decreased by 5 metres in the placebo group (\( P<0.05 \)). A different picture was seen in patients with systemic sclerosis-PAH; among the 14 patients in the placebo group there was a decline of 40 metres, while in the 33 patients receiving active therapy deterioration was prevented with an improvement of 3 metres (\( P=\) not significant). No RCT data specifically on CTD-PAH have been published to date for the other currently available endothelin-1 receptor antagonists, ambrisentan and macitentan.

**Phosphodiesterase-5 inhibitors**

A subgroup analysis of the 84 patients with CTD-PAH (including 50 patients with systemic sclerosis) who were included in the SUPER-1 trial of sildenafil was subsequently published \[46,47\]. In the original study of 278 patients, in which patients received 20 mg, 40 mg or 80 mg three times a day, the mean placebo-controlled treatment effects on 6MWD were +45 metres, +46 metres and +50 metres after 12 weeks. Direct comparison is difficult especially since patients with SLE may have a greater response to treatment. However, the treatment effect in the subgroup analysis tended to be smaller, with the 6MWD increasing by +42 metres, +36 metres and +15 metres in patients receiving the three different doses.

**Comparison of systemic sclerosis-PAH with idiopathic PAH**

Survival in systemic sclerosis-PAH is significantly worse than in idiopathic PAH, despite less severe pulmonary haemodynamics (Figure 2) \[20,25,48–50\]. There are several potential reasons for this disconnect between pulmonary hypertension severity and outcome (Table 2).

**Age**

Patients with systemic sclerosis-PAH are up to a decade older than patients with idiopathic PAH at diagnosis \[25\]. Age is an independent prognostic factor in systemic sclerosis-PAH \[26\] and was also found to be an independent predictor of mortality in a study involving 50 systemic sclerosis-PAH patients and 41 idiopathic PAH patients \[50\].

![Figure 2. Survival of IPAH and SSc-PAH in the ASPIRE registry](image)

Adapted with permission \[25\]. Abbreviations: IPAH, idiopathic pulmonary arterial hypertension; PAH, pulmonary arterial hypertension; SSc, systemic sclerosis.

**Pulmonary vasculopathy**

Dorfmüller et al. studied lung tissue samples from 8 patients with various forms of CTD-PAH and compared them with 29 idiopathic PAH patients \[10\]. Obstructing lesions in pulmonary veins and venules were seen in 75% of the CTD-PAH patients and 17% of the idiopathic PAH patients (\( P<0.05 \)). Overbeek et al. subsequently studied lung tissue samples from 8 systemic sclerosis-PAH and 11 idiopathic PAH patients \[51\]. They observed plexogenic lesions in all the idiopathic PAH patients and none of the systemic sclerosis-PAH patients. Furthermore, they observed pulmonary vein and venule fibrosis in all of the systemic sclerosis-PAH patients (with a pattern consistent with PVOD in 50%) and only 3 of the idiopathic PAH patients. Gunther et al. observed at least 2 radiographic signs consistent with PVOD on high resolution computerized tomography (CT) imaging (centrilobular ground glass abnormalities, septal lines and lymphadenopathy) in 16/26 patients with systemic
Despite pulmonary haemodynamics being more severe poorer outcomes in systemic sclerosis-PAH. A higher proportion of pulmonary venule involvement in SSc-PAH [10,51]. This difference persisted even more abnormal than in systemic sclerosis-PAH patients. The underlying reason for these observations is not clear. It is known that abnormal collagen deposition is high in both ventricles in patients with systemic sclerosis and no evidence of left heart disease or PAH [56]. However, when Overbeek et al. examined right ventricle samples for five systemic sclerosis-PAH and nine idiopathic PAH patients they found no difference in extent of interstitial fibrosis although there was a greater inflammatory infiltrate in the systemic sclerosis-PAH group [9].

Right ventricular function
Despite pulmonary haemodynamics being more severe in the idiopathic PAH group, Mathai et al. observed significantly higher NT-proBNP levels in 55 systemic sclerosis-PAH patients when compared with 43 idiopathic PAH patients [53]. This difference persisted even when adjusting for renal function and age, and suggests a poorer ability of the right ventricle to cope with increased afterload. Overbeek et al. studied 13 systemic sclerosis-PAH and 17 idiopathic PAH patients using the pump function graph technique and demonstrated poorer right ventricle contractility in the systemic sclerosis-PAH group [54]. Tedford et al. performed pressure-volume loops in 5 idiopathic PAH, 7 systemic sclerosis-PAH and 7 systemic sclerosis non-pulmonary hypertension patients [55]. No significant difference in right ventricle afterload as assessed by pulmonary vascular resistance (PVR), compliance and arterial elastance (Ea) was identified between the idiopathic PAH and systemic sclerosis-PAH patients. Right ventricle contractility as measured using end-systolic elastance (Ees) and the coupling of right ventricle contractility with afterload (Ees:Ea) were, however, poorer in systemic sclerosis-PAH than in idiopathic PAH. Interestingly, Ees:Ea in systemic sclerosis patients without pulmonary hypertension was even more abnormal than in systemic sclerosis-PAH patients. The underlying reason for these observations is not clear. It is known that abnormal collagen deposition is high in both ventricles in patients with systemic sclerosis and no evidence of left heart disease or PAH [56]. However, when Overbeek et al. examined right ventricle samples for five systemic sclerosis-PAH and nine idiopathic PAH patients they found no difference in extent of interstitial fibrosis although there was a greater inflammatory infiltrate in the systemic sclerosis-PAH group [9].

Left heart disease
Hachulla et al. performed cardiac magnetic resonance imaging in 52 patients with systemic sclerosis and observed evidence of left ventricle systolic or diastolic dysfunction in approximately ⅓ of cases, raising the possibility that occult left heart disease may be responsible for a proportion of diagnosed systemic sclerosis-PAH [57]. Fisher et al. observed evidence of left heart diastolic dysfunction in approximately ⅓ of 50 patients fulfilling diagnostic criteria for systemic sclerosis-PAH (with a mean age of 59) but only in approximately 10% of patients diagnosed with idiopathic PAH (with a mean age of 48), although its presence was not predictive of higher mortality [49]. Fox et al. identified pulmonary hypertension in 53/107 systemic sclerosis patients who underwent left and right heart catheterization [58]. Originally, 29/53 patients were given a diagnosis of PAH, but after fluid challenge 11/29 (38%) were reclassified as pulmonary hypertension secondary to left heart disease, which was in keeping with radiological appearances of the left heart and risk factors for left heart disease. A subsequent study identified 207 patients fulfilling haemodynamic criteria for PAH, 49% of whom had underlying CTD [59]. Following fluid challenge, 46/207 (22%) were reclassified.

Table 2. Potential reasons for poorer outcome in SSc-PAH than in IPAH

| Factor                     | Evidence                                                                 |
|----------------------------|--------------------------------------------------------------------------|
| Age                        | Mean age of diagnosis of SSc-PAH <10 yrs > IPAH [25]. Age was independent prognostic factor in combined group of SSc-PAH and IPAH [30]. |
| Pulmonary vasculopathy     | Higher proportion of pulmonary venule involvement in SSc-PAH [10,51].    |
| Right ventricle (RV)       | Reduced RV contractility assessed by pump function graph and pressure-volume loops in SSc-PAH [54,55]. Higher NT-proBNP in SSc-PAH despite less severe pulmonary haemodynamics [53]. |
| Left ventricle (LV)        | High prevalence of LV systolic and diastolic dysfunction in SSc [50,57]. 1/3 of patients fulfilling haemodynamic criteria for SSc-PAH may have occult left heart disease based on response to fluid challenge and clinico-radiological characteristics [58]. |
| Interstitial lung disease (ILD) | ILD common in SSc. Different registries have included varying degrees of ILD in SSc-PAH cohorts [29]. Response to PH therapy in SSc in the presence of ILD appears poor [75]. |
| Multisystem disease        | Coexisting renovascular and gastrointestinal disease including iron deficiency and malnutrition more common in SSc than in general population. |
| Antibodies                 | Exact role of SSc-associated autoantibodies (e.g. anticentromere) in pathogenesis of PAH and poorer prognosis compared with idiopathic PAH not clear [76]. Elevated stimulating anti-endothelin receptor type A antibodies and anti-angiotensin receptor type-I antibodies in SSc-PAH compared with idiopathic PAH [17]. |

Abbreviations: IPAH, idiopathic pulmonary arterial hypertension; PAH, pulmonary arterial hypertension; PH, pulmonary hypertension; NT-proBNP, N-terminal pro-brain natriuretic peptide; SSc, systemic sclerosis.
as pulmonary hypertension secondary to left heart disease. It is, however, well recognized that elderly patients fulfilling haemodynamic criteria for idiopathic PAH also have a high prevalence of risk factors for left heart disease and so it is unclear whether the potential misdiagnosis of pulmonary hypertension with left heart disease as PAH is more common in idiopathic PAH than in CTD-PAH [3]. Furthermore, the role of fluid challenge in the assessment of pulmonary hypertension is not well defined, with the positive predictive value for the presence of pulmonary hypertension secondary to left heart disease being unknown; hence it is not part of the diagnostic algorithm in the most recent world pulmonary hypertension guidelines [1].

**Interstitial lung disease**

Interstitial lung disease (ILD) is common in patients with systemic sclerosis and occurs more frequently in diffuse than in limited cutaneous disease [60]. Defining pulmonary hypertension in systemic sclerosis patients as being isolated PAH rather than ILD-associated pulmonary hypertension is not always straightforward. In the absence of significant pulmonary hypertension, an approach combining radiological assessment (limited disease when the extent of ILD <10% and extensive disease when the extent of disease >30%) and a spirometric threshold of FVC 70% (when the extent of ILD is estimated to be 10–30%) has been shown to effectively identify patients with systemic sclerosis and ILD with good and poor prognoses [46]. Whether this approach also robustly differentiates systemic sclerosis-PAH from ILD-associated pulmonary hypertension is unclear. Several thresholds involving FVC or total lung capacity (TLC) of 60 or 70% and/or varyingly-defined minor/mild versus moderate/severe ILD have been used in different studies and it is very possible that cohorts of “systemic sclerosis-PAH” in the literature have been “contaminated” with a proportion of patients who actually have ILD-associated pulmonary hypertension [29]. Several groups have demonstrated that ILD-associated pulmonary hypertension in systemic sclerosis is associated with a significantly poorer survival than systemic sclerosis-PAH, despite similar pulmonary haemodynamics [26,61]. A retrospective study of 70 systemic sclerosis patients with ILD-associated pulmonary hypertension was unable to demonstrate any clear benefit from PAH-specific therapy. However, prospective data are required and many centres have a low threshold for PAH-specific therapy in this group of patients.

**Iron deficiency**

Iron deficiency has previously been shown to be common in idiopathic PAH, occurring in up to 61% of patients in the absence of overt anaemia in the majority of cases, and is associated with poorer outcome [62]. Inappropriate elevation in serum hepcidin levels with subsequent reduced iron gut absorption has been postulated as a possible mechanism. Iron deficiency has recently been reported as occurring in 46% of systemic sclerosis-PAH and 16% of systemic sclerosis non-pulmonary hypertension patients [63]. Both exercise capacity and survival were reduced in those systemic sclerosis-PAH patients with iron deficiency. Hepcidin levels were found to be elevated while underlying gastrointestinal angiodysplastic lesions may also predispose iron deficiency in systemic sclerosis patients. Although it is not clear whether iron deficiency is more severe in systemic sclerosis-PAH, it is interesting to note that haemoglobin levels were statistically significantly lower in a recent study comparing 228 systemic sclerosis-PAH patients and 279 idiopathic PAH patients [47].

**Multisystem disease**

Extra-cardiorespiratory manifestations of systemic sclerosis may impact on both morbidity and mortality. Scleroderma renal crisis was previously a major cause of mortality, although is now more treatable since the availability of ACE inhibitors. Gastrointestinal involvement can lead to severe nutritional deficiency.

**Selected other CTD-PAH**

The exact prevalence of pulmonary hypertension in patients with SLE is unknown and estimates have ranged from 2–43% depending on diagnostic methods used [64,65]. Echocardiography-based studies are associated with a significant risk of over-estimating pulmonary pressures. Despite the prevalence of SLE being at least 3 times that of systemic sclerosis in the UK [66], the number of SLE-associated PAH (SLE-PAH) patients in the UK national registry was only 11% of the number of systemic sclerosis-PAH patients, suggesting that the true prevalence of PAH in patients with SLE is likely to be <1% [26]. A recent prospective study of 245 patients with SLE demonstrated pulmonary hypertension associated with significant lung or left heart disease in 12 patients (5%), but no cases of PAH were identified [67]. Patients with SLE-PAH are younger than those with systemic sclerosis-PAH, with higher pulmonary pressures and lower Dlco [26,68]. Survival appears to be superior with 3-year survival in the UK national registry of 74%, compared to 47% in systemic sclerosis-PAH [26]. Significant improvements in PAH have been described in patients with SLE-PAH (as well as some patients with mixed connective tissue disease-associated PAH) treated with immunosuppression alone, although no good RCT data exist [69]. Our approach is to treat active lupus aggressively and ensure that all patients with SLE-PAH are adequately immunosuppressed. Antiphospholipid syndrome is a risk factor
for the development of chronic thromboembolic pulmonary hypertension and so, if SLE-PAH is suspected, then particular attention to the exclusion of thromboembolic disease is required [70].

Antisynthetase syndrome is characterised by myositis and interstitial lung disease in the presence of anti-tRNA-synthetase antibodies (most commonly Jo-1). A recent French study calculated a right heart catheter-based prevalence of pulmonary hypertension of 7.9% [71]. The majority of patients had severe pre-capillary pulmonary hypertension in the presence of significant coexisting ILD while 3-year survival was 58%. PAH appears to be less common in other forms of CTD, with an estimated prevalence of 2% in mixed connective tissue disease [72], while studies estimating the prevalence of PAH in other forms of CTD (including Sjogren’s syndrome and rheumatoid arthritis) have used echocardiography to calculate systolic pulmonary arterial pressures with significant risk of over-estimation [73,74].

**Conclusion**

Survival in systemic sclerosis-PAH remains poor. This may be as a result of the underlying pulmonary arterial vasculopathy and the poor ability of the right ventricle to cope with increased afterload, together with the multi-system nature of the condition including the involvement of myocardium and lungs. Although early detection of systemic sclerosis-PAH in patients is required, further work to demonstrate a definite impact on the natural history of the condition is needed. More extensive investigation of the underlying molecular and genetic pathogenesis is warranted, while specific trials of currently available and novel PAH-specific therapies in systemic sclerosis-PAH are urgently required.

**Abbreviations**

6MWD, six minute walk distance; CTD, connective tissue disease; CTEPH, chronic thromboembolic pulmonary hypertension; DLCO, lung gas transfer; Ea, arterial elastance; Ees, end-systolic elastance; FC, Functional Class; FVC, forced vital capacity; ILD, interstitial lung disease; mPAP, mean pulmonary arterial pressure; PAH, pulmonary arterial hypertension; NT-proBNP, N-terminal pro-brain natriuretic peptide; PVOD, pulmonary veno-occlusive disease; PVR, pulmonary vascular resistance; RCT, randomized controlled trial; SLE, systemic lupus erythematosus; TGF-β, transforming growth factor beta; WHO, World Health Organisation.

**Disclosures**

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