Borderline pulmonary arterial pressure in systemic sclerosis patients: a post-hoc analysis of the DETECT study

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Abstract

Introduction: Patients with mean pulmonary artery pressures (mPAP) of 21 to 24 mm Hg have a so-called borderline elevation of mPAP (BoPAP)—a condition thought to represent early-stage pulmonary arterial vasculopathy. Based on the DETECT study, this post-hoc analysis examined patient characteristics of systemic sclerosis (SSc) patients with normal mPAP, BoPAP, and elevated mPAP, fulfilling pulmonary arterial hypertension (PAH) criteria.

Methods: Adult patients with a duration of SSc more than 3 years, a diffusing capacity of the lung for carbon monoxide less than 60% predicted, and no previous diagnosis of any form of pulmonary hypertension (PH) underwent screening tests followed by right heart catheterization. Subjects were divided into three groups: normal mPAP, BoPAP, and PAH. Exploratory comparative and binary logistic regression analyses were performed for the BoPAP versus normal mPAP and PAH versus BoPAP groups.

Results: Of 244 patients evaluated, 148 (60%) had normal mPAP, 36 (15%) had BoPAP, and 60 (25%) had definite PAH. Univariable logistic regression (ULR) showed the mean tricuspid regurgitation velocity in patients with BoPAP to be intermediate between normal mPAP and PAH. In the ULR analyses BoPAP versus normal mPAP and PAH versus BoPAP, the statistically significant predictors were, amongst others: demographic, clinical, pulmonary function, echocardiographic and hemodynamic variables.

Conclusions: In this exploratory post-hoc analysis of the DETECT study population patients with BoPAP could be distinguished from patients with normal mPAP and PAH, and it appears that BoPAP may be an intermediate stage on the continuum between normal PA pressures and PAH.

Introduction

Systemic sclerosis (SSc, scleroderma) is an autoimmune connective tissue disorder characterized by inflammation, fibrosis and vasculopathy. Pulmonary arterial hypertension (PAH) is a leading cause of death in SSc [1,2]. Based on registry data, modern treatment has improved two-year survival estimates of SSc-associated PAH (SSc-PAH) from 40% [3] to 58% [4], a clear improvement which, however, lags gains seen in other forms of PAH [5-7]. Screening for SSc-PAH is capable of identifying earlier stages of the disease, resulting in earlier intervention that may improve survival [8]. Thus, early diagnosis and treatment of SSc-PAH is of paramount importance [9,10].

The DETECT study was a multi-center study that systematically evaluated 466 SSc patients at increased risk for development of SSc-PAH [11]. DETECT was the first SSc-PAH detection study to evaluate all subjects with right heart catheterization (RHC), the gold standard test for the diagnosis of PAH [9]. Detailed demographic, clinical,
Echocardiographic, serologic and functional testing data were also collected.

Measurements in healthy individuals show a normal mean pulmonary arterial pressure (mPAP) to be 14 ± 3.3 mmHg [12], with 2 SD extending the normal range up to 20.6 mm Hg. The consensus definition of PAH requires mPAP ≥25 mmHg and pulmonary artery wedge pressure (PAWP) ≤15 mmHg [9,13]. Thus, a borderline range of mPAPs exists between 21 and 24 mm Hg that may represent an early, milder stage of pulmonary vasculopathy in those at high risk of developing PAH [13-15]. This concept is especially relevant in connective tissue diseases (CTD), such as SSc, where progressive vasculopathy is highly prevalent and may be an important distinction from idiopathic PAH [13]. The objective of this post-hoc analysis was to compare the demographics and clinical features of patients with normal PAP, borderline mPAP (BoPAP), and elevated PAP (PAH) in a large international cohort of patients with SSc who participated in the DETECT study. The identification of characteristics specific to SSc patients with BoPAP would facilitate future investigations into the natural history of this condition, and provide insights into the proportion of patients who develop PAH.

Methods

Study design

Our study was conducted in accordance with the Declaration of Helsinki and its amendments, followed the International Conference on Harmonization Guideline for Good Clinical Practice, and was approved by local institutional review boards/ethics committees (a complete list is included as Additional file 1). RHC and echocardiography protocols were standardized and conducted systematically, and serum laboratory testing as well as data management were performed centrally. Data quality was monitored rigorously. All patients provided written informed consent. Patients were eligible for inclusion in DETECT if they were aged ≥18 years and had: 1) a definite diagnosis of SSc [16] of ≥3 years’ duration from first non-Raynaud’s symptom; 2) a diffusing capacity of the lung for carbon monoxide (DLCO) <60% of predicted; 3) a forced vital capacity (FVC) ≥40% of predicted; and 4) no left atrial hypertension confirmed by RHC prior to enrolment.

Current analysis population

Patients were included in the current analysis if they had: 1) a pulmonary artery wedge pressure (PAWP) ≤15 mmHg by RHC; 2) no significant interstitial lung disease (ILD; defined as FVC <60% or FVC between 60 and 70% with moderate-to-severe fibrosis on high resolution computed tomography); 3) no systemic hypertension (stage-I hypertension defined as systolic blood pressure ≥140 mmHg or diastolic blood pressure ≥90) [17]; and 4) no left atrial enlargement (defined as 3.9 cm for women and 4.1 cm for men) [18]. DETECT screened 646 SSc patients, and enrolled 488 SSc patients with 466 who underwent RHC. A total of 222 of 466 were excluded from the present analysis due to: 1) a pulmonary artery wedge pressure (PAWP) >15 mmHg or significant ILD (138 patients), and 2) an enlarged left atrium or stage-I or greater hypertension (84 patients). Eligible patients comprise the PAP analysis set and were categorized into three PAP groups for analysis, based on the mPAP by RHC: 1) normal mPAP (<21 mmHg); 2) BoPAP (mPAP 21 to 24 mmHg); and 3) PAH (mPAP ≥25 mmHg). Analysis of variables was performed using these three PAP groups.

Data collection and analysis

The DETECT study collected 112 variables, collated into four groups: 1) demographic and clinical parameters (68 variables); 2) serum tests (13 variables); 3) electrocardiography (ECG) (3 variables); and 4) echocardiography (28 variables). In the present study, variables were selected for further analysis based on a review of numerical descriptive differences between the three groups, as well as input from the authors on feasibility of the variables and their clinical relevance to PAH. Two additional variables were computed: 1) the transpulmonary gradient (TPG) which is the difference between mPAP and left atrial pressure (estimated by the PAWP); and 2) the diastolic wedge gradient (DWG) which is the difference between diastolic PAP and left atrial pressure (estimated by the PAWP). The selected variables were described using summary statistics: sample size, mean, SD, median, upper and lower quartiles, minimum and maximum for quantitative data and frequencies (counts and percentages) for qualitative and categorical data. The distributions of the variables were compared using non-parametric tests: Wilcoxon rank-sum and chi-square/Fisher’s exact test for continuous and categorical data, respectively. Univariable logistic regression (ULR) analysis was performed using BoPAP versus normal mPAP and PAH versus BoPAP as binary outcomes in separate models. The odds ratio (OR) and 95% CI for each variable was calculated and statistical significance was examined by the Wald chi-square test. A Forest plot was constructed to display the odds ratios and the 95% CI for 13 variables in the BoPAP versus normal mPAP and PAH versus BoPAP groups. Variables were selected for the plot if they met the statistical significance criteria or had potential clinical utility as screening tests for BoPAP and PAH. RHC hemodynamics were used for exploratory purposes only, as they were part of the group definitions.

Results

Among the 244 SSc patients included in the PAP analysis set, 60% (n = 148) had a normal mPAP, 15% (n = 36) had BoPAP and 25% (n = 60) had an elevated mPAP (PAH) (Figure 1).
Baseline characteristics of patients with normal mPAP, BoPAP and PAH

Statistically significant differences between the BoPAP and other groups were found in clinical, serologic, echocardiographic, and invasive hemodynamic parameters, including World Health Organization functional class (WHO FC) (PAH versus BoPAP), presence of peripheral edema (BoPAP versus normal mPAP), presence of telangiectasias (PAH versus BoPAP), ratio of FVC percent predicted/DLCO percent predicted (PAH versus BoPAP), TLC percent predicted (BoPAP versus normal mPAP), presence of anti-centromere antibodies (PAH versus BoPAP), log_{10} NT-proBNP (BoPAP versus normal mPAP), serum urate (PAH versus BoPAP), left atrium diameter (BoPAP versus normal mPAP, tricuspid annular plane systolic excursion (TAPSE, PAH versus BoPAP) and TR velocity (BoPAP versus both normal PAP and PAH; Table 1). Differences in the exploratory comparison of hemodynamic variables (PAWP, TPG, DWG and pulmonary vascular resistance (PVR)) were also noted.

Univariable logistic regression analysis

**BoPAP versus normal PAP**

As shown in Table 2 and Figure 2, ULR models of SSc patients with BoPAP versus normal mPAP identified the following variables as being statistically significant predictors of BoPAP (P <0.05): older age, presence of peripheral edema, a higher log_{10} NT-proBNP, greater left atrium diameter, and greater tricuspid regurgitation (TR) velocity. Of note, 6-minute walk distance was not statistically different.

The RHC variables PAWP, TPG and PVR (which defined the groups) were all statistically significant, though DWG was not.

**PAH versus BoPAP**

ULR models of SSc patients with PAH versus BoPAP identified the following variables as being statistically significant (P <0.05) predictors of PAH: higher WHO FC (III or IV), diffuse versus limited disease subtype, the presence of telangiectasias, higher FVC percent predicted/DLCO percent predicted ratio, presence of anti-centromere antibodies, higher serum urate, a lower TAPSE and a higher TR velocity. As was the case with the BoPAP versus normal PAP analysis, 6-minute walk distance was not statistically different. As expected, the PAH group had a higher TPG, a higher DWG, and a higher PVR (Table 2 and Figure 2). Of note, the PAWP was not significantly different between these groups.

**Discussion**

Expert consensus opinion from the recent 5th World Symposium on Pulmonary Hypertension acknowledges that a borderline mPAP between 21 and 24 mmHg in SSc patients is associated with ‘a high risk of future development of manifest PAH’, and recommends careful follow up of these patients [13]. In our current study of a large, well-defined, high-risk SSc cohort we identified clinical and laboratory features that differentiate the BoPAP group from groups with normal mPAP and PAH. Our study provides a crucial first step towards the development of future longitudinal studies of BoPAP, as we
Table 1. Baseline characteristics of patients with normal mPAP, BoPAP and PAH

| Variable                                      | Patient groups (N=244) | P-value          |
|-----------------------------------------------|------------------------|-----------------|
|                                              | Normal mPAP (N=148)    | BoPAP (N=36)    | PAH (N=60) |
| Demographic and clinical parameters          |                        | BoPAP versus normal mPAP | PAH versus BoPAP |
| Age, years, N                               | 148                    | 36              | 60          |
| Mean ± SD                                    | 54.3 ± 12.0            | 58.4 ± 9.5      | 61.6 ± 9.7  |
| Median, Q1 to Q3                             | 55.0, 46.0 to 63.5     | 58.5, 52.5 to 63.5 | 62.0, 56.0 to 68.0 |
| Minimum, maximum                             | 260, 78.0              | 43.0, 80.0      | 33.0, 80.0  |
| Gender, n (%)                                | 148                    | 36              | 60          |
| Male, n (%)                                  | 18 (12.2)              | 6 (16.7)        | 12 (20.0)   |
| Female, n (%)                                | 130 (87.8)             | 30 (83.3)       | 48 (80.0)   |
| WHO FC, N                                     | 139                    | 34              | 60          |
| I/II, n (%)                                   | 127 (91.4)             | 28 (82.4)       | 37 (61.7)   |
| III/IV, n (%)                                 | 12 (8.6)               | 6 (17.6)        | 23 (38.3)   |
| 6-minute walk distance, N                    | 108                    | 27              | 48          |
| Mean ± SD                                    | 422.4 ± 110.4          | 397.4 ± 102.3   | 391.4 ± 109.8 |
| Median, Q1 to Q3                              | 432.5, 364.5 to 502.0  | 414.5, 341.0 to 472.0 | 395.0, 318.0 to 460.5 |
| SSc disease duration (years), N              | 147                    | 36              | 60          |
| Mean ± SD                                    | 110.0 ± 8.0            | 100.0 ± 5.0     | 140.0 ± 12.0|
| Median, Q1 to Q3                              | 80.0, 5.0 to 13.0      | 9.0, 6.0 to 15.0 | 11.0, 6.0 to 16.0 |
| SSc subtype, N                                | 146                    | 36              | 59          |
| Limited, n (%)                                | 90 (61.6)              | 20 (55.6)       | 44 (74.6)   |
| Diffuse, n (%)                                | 45 (30.8)              | 13 (36.1)       | 10 (16.9)   |
| Overlap/mixed CTD, n (%)                      | 11 (7.5)               | 3 (8.3)         | 5 (8.5)     |
| Presence of peripheral edema, n/N (%)         | 8/147 (5.4)            | 7/36 (19.4)     | 11/60 (18.3) |
| Presence of telangiectasias, n/N (%)          | 108/148 (73.0)         | 22/36 (61.1)    | 52/60 (86.7) |
| FVC % predicted, N                            | 148                    | 36              | 60          |
| Mean ± SD                                    | 92.5 ± 17.6            | 88.7 ± 14.5     | 92.0 ± 19.7 |
| Median, Q1-Q3                                 | 91.9, 79.9 to 102.7    | 87.1, 80.0 to 96.5 | 89.5, 77.2 to 102.5 |
| FVC % predicted /DLCO % predicted, N          | 148                    | 36              | 60          |
| Mean ± SD                                    | 1.9 ± 0.4              | 1.9 ± 0.4       | 2.3 ± 0.7   |
| Median, Q1 to Q3                              | 1.8, 1.5 to 2.1        | 2.0, 1.5 to 2.2 | 2.2, 1.8 to 2.5 |
| TLC % predicted, N                            | 127                    | 28              | 54          |
| Mean ± SD                                    | 92.8 ± 21.9            | 84.8 ± 14.4     | 88.1 ± 21.6 |
| Median, Q1 to Q3                              | 92.5, 80.0 to 106.0    | 83.0, 74.0 to 95.0 | 87.3, 78.0 to 100.3 |
| Serum laboratory tests                        |                        |                 |             |
| Presence of anti-centromere antibodies         |                        |                 |             |
| n/N (%)                                       | 49/141 (34.8)          | 10/34 (29.4)    | 30/56 (53.6) |
| Log_{10} NT-proBNP (pg/ml), N                 | 141                    | 34              | 56          |
| Mean ± SD                                    | 2.0 ± 0.5              | 2.2 ± 0.5       | 2.4 ± 0.5   |
| Median, Q1 to Q3                              | 2.1, 1.7 to 2.3        | 2.2, 1.9 to 2.5 | 2.3, 2.0 to 2.8 |
| Serum urate, mg/100 ml, N                     | 141                    | 34              | 56          |
| Mean ± SD                                    | 4.7 ± 1.4              | 4.7 ± 1.3       | 5.6 ± 1.4   |
| Median, Q1 to Q3                              | 4.4, 3.7 to 5.3        | 4.5, 3.7 to 5.4 | 5.6, 4.6 to 6.5 |
Formalized early detection of SSc-PAH has been shown to identify milder forms of the disease, resulting in opportunities for earlier management [8,19]. As algorithms and practice guidelines are developed to improve early detection of SSc-PAH, it is likely that referrals for RHC will increase (in DETECT the rate increased from 40% to 62%). The increase in RHCs will likely result in more patients identified with both PAH and BoPAP. The finding of BoPAP is significant, as a number of recent studies have shown an association between BoPAP and worse clinical outcomes [14,15,20]. Thus, further characterization of BoPAP is important. In the current study we identify significant differences in clinical, serologic, echocardiographic and hemodynamic parameters useful in differentiating BoPAP from normal PAP and PAH from BoPAP groups.

### Table 1 Baseline characteristics of patients with normal mPAP, BoPAP and PAH (Continued)

| Log_{10} estimated GFR | 141 | 34 | 55 |
|------------------------|-----|----|----|
| 1.9 ± 0.1              | 1.9 ± 0.1 | 1.9 ± 0.1 |
| 1.9, 1.9 to 2.0        | 1.9, 1.8 to 2.0 | 1.9, 1.8 to 2.0 |

#### Echocardiography

| Left atrium diameter, mm, N | 148 | 36 | 60 | <0.001 |
|-----------------------------|-----|----|----|--------|
| Mean ± SD                  | 30.2 ± 5.1 | 33.9 ± 3.6 | 31.8 ± 5.4 |
| Median, Q1 to Q3            | 31.0, 27.8 to 34.0 | 33.5, 31.0 to 37.0 | 32.0, 30.0 to 36.0 |

| Right atrium area, cm², N | 142 | 33 | 58 | <0.05 |
|---------------------------|-----|----|----|-------|
| Mean ± SD                 | 128 ± 4.7 | 14.7 ± 5.9 | 16.8 ± 5.8 |
| Median, Q1 to Q3           | 120, 10.0 to 14.3 | 124, 10.5 to 18.0 | 15.8, 13.4 to 20.0 |

| TAPSE (mm), N | 138 | 33 | 53 |
|---------------|-----|----|----|
| Mean ± SD     | 232 ± 4.3 | 23.8 ± 5.1 | 21.1 ± 4.3 |
| Median, Q1 to Q3 | 23.7, 20.4 to 26.0 | 23.0, 20.0 to 27.0 | 20.8, 19.0 to 23.0 |

| TR velocity (m/s), N | 140 | 35 | 58 | <0.001 | <0.05 |
|----------------------|-----|----|----|--------|-------|
| Mean ± SD            | 2.3 ± 0.4 | 2.7 ± 0.3 | 3.0 ± 0.7 |
| Median, Q1 to Q3     | 2.3, 2.2 to 2.6 | 2.7, 2.5 to 2.9 | 2.9, 2.5 to 3.5 |

#### Electrocardiography

| Presence of right axis deviation, n/N (%) | 2/138 (1.4) | 1/30 (3.3) | 8/58 (13.8) |
|------------------------------------------|-------------|------------|-------------|

| PAWP, mmHg, N                       | 148 | 36 | 60 | <0.001 |
|-------------------------------------|-----|----|----|--------|
| Mean ± SD                           | 7.7 ± 3.3 | 10.8 ± 3.0 | 9.9 ± 3.3 |
| Median, Q1 to Q3                    | 8.0, 5.0 to 10.0 | 11.0, 9.0 to 13.0 | 10.5, 7.5 to 12.5 |

| TPG, mmHg, N                        | 148 | 36 | 60 | <0.001 | <0.001 |
|-------------------------------------|-----|----|----|--------|-------|
| Mean ± SD                           | 7.9 ± 2.7 | 11.6 ± 2.8 | 22.3 ± 9.7 |
| Median, Q1 to Q3                    | 7.0, 6.0 to 10.0 | 11.0, 10.0 to 13.0 | 20.0, 15.0 to 28.0 |

| DWG, mmHg, N                        | 148 | 36 | 60 | <0.001 |
|-------------------------------------|-----|----|----|--------|
| Mean ± SD                           | 1.8 ± 2.7 | 3.0 ± 5.4 | 10.7 ± 8.1 |
| Median, Q1 to Q3                    | 1.0, 0.0 to 3.0 | 2.0, 0.5 to 5.0 | 8.0, 5.0 to 17.0 |

| PVR, dyn.sec/cm², N                 | 148 | 36 | 60 | <0.001 | <0.001 |
|-------------------------------------|-----|----|----|--------|-------|
| Mean ± SD                           | 128.0 ± 50.8 | 179.8 ± 58.7 | 375.8 ± 217.0 |
| Median, Q1 to Q3                    | 121.5, 91.3 to 153.9 | 170.3, 147.8 to 187.8 | 295.5, 236.0 to 419.1 |

*N for each variable represents the number of patients with available data. Statistical tests were Wilcoxon rank-sum for continuous data and chi-square/Fisher’s exact for categorical data. These variables were analysed for completeness, but it should be borne in mind that they are part of the definition of the analysis groups. BoPAP, borderline pulmonary arterial pressure; CTD, connective tissue disease; DLCO, diffusing capacity of carbon monoxide; DWG, diastolic wedge gradient; FVC, forced vital capacity; GFR, glomerular filtration rate; NT-proBNP, N-terminal pro-brain natriuretic peptide; PAH, pulmonary arterial hypertension; PAWP, pulmonary artery wedge pressure; PVR, pulmonary vascular resistance; Q, quartile; SSc, systemic sclerosis; TAPSE, tricuspid annular plane systolic excursion; TLC, total lung capacity; TPG, transpulmonary gradient; TR, tricuspid regurgitation; WHO FC, World Health Organization functional class.*

provide a panel of variables that can be tested to determine their prognostic utility.

Formalized early detection of SSc-PAH has been shown to identify milder forms of the disease, resulting in opportunities for earlier management [8,19]. As algorithms and practice guidelines are developed to improve early detection of SSc-PAH, it is likely that referrals for RHC will increase (in DETECT the rate increased from 40% to 62%). The increase in RHCs will likely result in more patients identified with both PAH and BoPAP. The finding of BoPAP is significant, as a number of recent studies have shown an association between BoPAP and worse clinical outcomes [14,15,20]. Thus, further characterization of BoPAP is important. In the current study we identify significant differences in clinical, serologic, echocardiographic and hemodynamic parameters useful in differentiating BoPAP from normal PAP and PAH from BoPAP groups.

A number of recent studies have shown an association between BoPAP and worse clinical outcomes. One small study compared SSc patients with mPAPs >17
but <25 mmHg (group mean of 20 ± 2 mmHg) to those with mPAPs ≤17 mmHg (group mean 14 ± 2 mmHg), and found the former group to be associated with both shorter mean 6-minute walk distances (396 ± 71 meters versus 488 ± 77 meters, \( P < 0.005 \)) and lower mean percent predicted peak oxygen consumption (\( \text{VO}_2 \)) values (76 ± 11% versus 90 ± 24%, \( P = 0.05 \)) [20]. The PHAROS registry included 206 SSc patients at increased risk for PAH who underwent RHC [21]. After excluding patients with significant interstitial lung disease, a comparison of SSc patients with normal mPAP (group median 16 mmHg) and borderline mPAP (group median 23 mmHg) showed the latter group to have significantly higher right ventricular systolic pressures on echocardiogram, higher pulmonary vascular resistance and a higher transpulmonary gradient [14].

### Table 2 Univariable logistic regression analysis of patients with BoPAP versus normal PAP and those with PAH versus BoPAP

| Variable                                      | Odds ratios (95% confidence interval) | \( P \)-value |
|-----------------------------------------------|--------------------------------------|---------------|
| **Demographic and clinical parameters**       |                                      |               |
| Age, years                                    | 1.03 (1.00, 1.07)                     | 1.04 (0.99, 1.08) |
| Gender, female versus male                    | 0.69 (0.25, 1.89)                     | 0.80 (0.27, 2.36) |
| WHO FC, III/IV versus I/II                    | 2.27 (0.78, 6.56)                     | 2.90 (1.04, 8.08) |
| 6-minute walk distance                        | 1.00 (0.99, 1.00)                     | 1.00 (1.00, 1.00) |
| SSc disease duration, years                   | 0.99 (0.94, 1.04)                     | 1.05 (1.00, 1.11) |
| SSc disease subtype                           |                                      | <0.05         |
| dcSSc versus lcSSc                            | 1.30 (0.59, 2.85)                     | 0.35 (0.13, 0.93) |
| Overlap/mixed CTD versus lcSSc                | 1.23 (0.31, 4.81)                     | 0.76 (0.17, 3.48) |
| Presence of telangiectasias                   | 0.58 (0.27, 1.25)                     | 4.13 (1.52, 11.26) |
| Presence of peripheral edema                  | 4.19 (1.41, 12.46)                    | 0.93 (0.32, 2.67) |
| FVC predicted                                 | 0.99 (0.97, 1.01)                     | 1.01 (0.99, 1.04) |
| FVC % predicted /DLCO % predicted             | 1.24 (0.54, 2.82)                     | 3.41 (1.36, 8.54) |
| TLC % predicted                               | 0.98 (0.96, 1.00)                     | 1.01 (0.99, 1.04) |
| **Serum laboratory**                          |                                      |               |
| Presence of anti-centromere antibodies        | 0.78 (0.35, 1.77)                     | 2.77 (1.12, 6.85) |
| \( \log_{10} \) NT-proBNP, pg/ml             | 2.65 (1.17, 5.98)                     | 1.69 (0.71, 4.05) |
| Serum urate, mg/100 ml                        | 1.01 (0.76, 1.33)                     | 1.73 (1.20, 2.48) |
| \( \log_{10} \) estimated GFR               | 0.54 (0.03, 9.44)                     | 0.04 (0.00, 1.77) |
| **Echocardiography**                          |                                      |               |
| Left atrium diameter, mm                      | 1.22 (1.10, 1.35)                     | 0.89 (0.80, 1.00) |
| Right atrium area, cm²                        | 1.07 (1.00, 1.14)                     | 1.07 (0.99, 1.17) |
| TAPSE, mm                                     | 1.03 (0.94, 1.12)                     | 0.88 (0.80, 0.98) |
| TR velocity, m/s                              | 25.10 (6.07, 103.74)                  | 3.15 (1.30, 7.67) |
| **Electrocardiography**                       |                                      |               |
| Right axis deviation                          | 2.35 (0.21, 26.74)                    | 4.64 (0.55, 38.99) |
| **Right heart catheterization**               |                                      |               |
| PAWP, mmHg                                    | 1.39 (1.21, 1.61)                     | 0.91 (0.79, 1.04) |
| TPG, mmHg                                     | 1.57 (1.33, 1.85)                     | 1.72 (1.35, 2.19) |
| DWG, mmHg                                     | 1.10 (0.99, 1.21)                     | 1.27 (1.12, 1.44) |
| PVR, dyn.sec/cm²                              | 1.01 (1.01, 1.02)                     | 1.03 (1.01, 1.04) |

BoPAP, borderline pulmonary arterial pressure; CTD, connective tissue disease; dcSSc, diffuse cutaneous SSc; DLCO, diffusing capacity of carbon monoxide; DWG, diastolic wedge gradient; FVC, forced vital capacity; GFR, glomerular filtration rate; lcSSc, limited cutaneous SSc; NT-proBNP, N-terminal pro-brain natriuretic peptide; PAH, pulmonary arterial hypertension; PAWP, pulmonary artery wedge pressure; PVR, pulmonary vascular resistance; SSc, systemic sclerosis; TAPSE, tricuspid annular plane systolic excursion; TLC, total lung capacity; TPG, transpulmonary gradient; TR, tricuspid regurgitation; WHO FC, World Health Organization functional class.
underwent repeat RHC at a later date showed that 32% of patients with normal mPAP and 55% of patients with borderline mPAP developed resting PH (P-value not significant) 14.92 ± 9.23 and 12.19 ± 6.82 months after the first RHC, respectively [14]. Another analysis compared the outcomes of SSc patients with mPAP ≤ 20 mmHg to those with mPAPs of 21 to 24 mmHg [15]. Within 228 patients without ILD, 142 had normal mPAP and 86 had mPAP of 21 to 24 mmHg. Clinically indicated repeat RHCs were performed in 38 patients from each group and the hazard ratio for PAH diagnosis on the subsequent RHC for the borderline mPAP group compared to the normal mPAP group was 3.7 (95% CI 1.7, 8.0, P <0.001). Within the borderline mPAP group, 18.5% (95% CI 8.3, 28.7) developed PAH within 3 years, and 27.1% (95% CI 13.9, 40.3) developed PAH within 5 years [15].

In the current study we identified significant differences in clinical, serologic, echocardiographic and hemodynamic parameters useful in differentiating BoPAP from normal PAP and PAH from BoPAP groups. Across both comparisons, TR velocity was the variable most strongly associated with BoPAP (versus normal mPAP) and PAH (versus BoPAP).

Our analysis also identified additional variables capable of differentiating BoPAP from the PAH group: WHO functional class, Ssc subtype, presence of telangiectasias, ratio of % predicted FVC/percent predicted DLCO, anti-centromere antibodies, and serum urate concentration. Some of these variables (telangiectasia [22], FVC/DLCO [22], anti-centromere antibodies [22-24], and serum urate [25]) have been useful in predicting SSc-PAH in other studies.

Invasive pressure measurements including TPG and DWG were explored, as these measurements have been addressed in other studies of BoPAP. Elevations in these gradients (≥12 mmHg and ≥5, respectively) [26] are believed consistent with pulmonary vascular disease, and thus, help separate individuals with PAH from those with pulmonary venous hypertension. As described previously [15], the TPG in our study was capable of separating the
BoPAP group from the normal PAP and PAH groups. We also show the DWG to be useful in differentiating the BoPAP group from the PAH group, and it has the added benefit of being able to differentiate PAH from post-capillary PH [26]. Though not in the abnormal range, the mean TPG and DWG measurements for the BoPAP group were intermediate between the normal PAP and PAH groups, perhaps suggesting that a subset of SSc patients with BoPAP may have early pulmonary vasculopathy.

In the present study we excluded patients with overt pulmonary venous hypertension based upon the finding of a PAWP >15 mmHg by RHC. It is important to acknowledge that occult pulmonary venous hypertension (OPVH) due to left heart disease may have been present in a proportion of patients who underwent a RHC while in a state of volume depletion, and were thus found to have underestimated left-heart filling pressures [27]. One study involving patients with scleroderma found that 6 out of 24 patients who originally met the criteria for PAH by RHC were subsequently diagnosed with OPVH after a fluid challenge [28]. As a fluid challenge was not included in our RHC study protocol, we minimized the likelihood of including patients with OPVH by excluding individuals with conditions commonly associated with OPVH: stage I or greater systemic hypertension and/or abnormally enlarged left atrium. Future BoPAP studies should consider the use of evocative maneuvers such as fluid challenge or exercise in order to identify patients with OPVH.

Our study has significant strengths. Ours is the first BoPAP study to mandate a diagnostic RHC in all subjects, allowing us to assign individuals into groups (normal mPAP, BoPAP and PAH) based on rigorous, objective criteria. Standardization of the study protocol and centralization of serum laboratory testing and data management assured homogeneity. In addition, it is the largest, most complete cohort of patients evaluated for SSc-PAH, providing us a sufficient sample size to conduct our analyses.

Our study has important limitations. Results are cross-sectional, so longitudinal follow up to determine the incidence of PAH in the BoPAP group over time was not determined. Our inclusion criteria included a DLCO <60% to enrich for a higher likelihood of PAH. Thus, the groups analyzed in our study represent a high-risk SSc group, and may not be representative of the general SSc population. Future studies are needed in order to expand our findings into the general SSc population. Our study was not designed to compare and model relationships involving the BoPAP sub-group; thus, multivariable logistic regression for this post-hoc sub-analysis could be misleading, and was not performed. In addition, the three sub-groups analyzed (normal mPAP, BoPAP and PAH) were defined using mPAP (a continuous variable). Various pairwise sub-

groups using binary logistic regression may provide a different set of predictive covariables for each multiple regression model that may be inaccurate.

Conclusion

In conclusion, our study identifies clinical, echocardiographic, hemodynamic, pulmonary function, and serologic variables that allow SSc patients with BoPAP to be differentiated from SSc patients with normal mPAPs and PAH. Future longitudinal studies designed to reassess patients with BoPAP at pre-designated time points would be the ideal means of validating the prognostic value of our findings. Such studies would allow further characterization of the natural history of BoPAP, including the rate and frequency of conversion to pre-capillary or post-capillary PH. Our exploratory analyses of hemodynamic data suggest that the TPG and DWG should continue to be evaluated as potential predictors for the development of PAH in the SSc population.

Additional file

Additional file 1: Institutional ethics committees and review boards.

Abbreviations

BoPAP: borderline elevation of the mean pulmonary arterial pressure; CTD: connective tissue disease; DLCO: diffusing capacity of the lung for carbon monoxide; DWG: diastolic wedge gradient; FVC: forced vital capacity; GFR: glomerular filtration rate; HfPEF: heart failure with preserved ejection fraction; ILD: interstitial lung disease; mmHg: millimeters of mercury; mPAP: mean pulmonary arterial pressure; NT-proBNP: N-terminal pro-brain natriuretic hormone; OPVH: occult pulmonary venous hypertension; OR: odds ratio; PAH: pulmonary arterial hypertension; PAWP: pulmonary artery wedge pressure; PVR: pulmonary vascular resistance; RHC: right heart catheterization; ROC AUC: receiver operating characteristic area under the curve; SSc: systemic sclerosis (scleroderma); SSC-PAH: scleroderma-associated pulmonary arterial hypertension; TAPSE: tricuspid annular plane systolic excursion; TCC: total lung capacity; TPG: transpulmonary gradient; TR: tricuspid regurgitation; ULR: univariable logistic regression; VO2: oxygen consumption; WHO FC: World Health Organization functional class.

Competing interests

OD has/had consultancy relationship and/or has received research funding in the area of systemic sclerosis and related conditions from Actelion, Pfizer, Erganex, BMS, Sanofi-Aventis, United BioSource Corporation, Roche/Genentech, Medac, Bovivitrum, Boehringer Ingelheim Pharma, Novartis, A D Science, Active Biotec, Bayer-Scherering, Sinoxa, Serodapharm, EphiPharm, Biogen and GSK. JGC has consultancy relationships, received lecture honoraria and/or has received research funding from Actelion Pharmaceuticals Ltd, Pfizer, GlaxoSmithKline and United Therapeutics. CPD has received research funding or had consultancy relationship with Actelion, Pfizer, Sanofi-Aventis, Roche, Novartis and Merck-Serono. EG has received honoraria for consultations and/or speaking at conferences from Actelion Pharmaceuticals Ltd, Bayer, Gilead, GlaxoSmithKline, Lilly, Milteney, Novartis, Pfizer and Rojex Medica, and funding for clinical trials from Actelion Pharmaceuticals Ltd, Bayer, GlaxoSmithKline, Encysive, Lilly and Pfizer. DB has acted as a consultant for Actelion Pharmaceuticals Ltd. UM-L: UML is a speaker and advisor for Actelion and Roche, and a speaker for GSK and Pfizer. JEP has consulted for and/or had research grants from Actelion, Bayer, Biogen and Celgene. MCV has consultancy relationships, and has received lecture honoraria and/or research funding from Actelion Pharmaceuticals Ltd, Pfizer, GlaxoSmithKline, Therabel Pharma and United Therapeutics. JRS has received consulting fees from Actelion, Aires, Aprex, Bayer, Boehringer-Ingelheim, Celgene, DART, Eiccose, Egger, EMD Serono, FibroGen, Gilead, InterMune Novartis, Pfizer.
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