Orthodeoxia and postural orthostatic tachycardia in patients with pulmonary arteriovenous malformations: a prospective 8-year series

ABSTRACT

Postural changes in 258 patients with pulmonary arteriovenous malformations (PAVMs) reviewed between 2005 and 2013 were evaluated prospectively using validated pulse oximetry methods. Of the 257 completing the test, 75 (29%) demonstrated orthodeoxia with an oxygen saturation fall of at least 2% on standing. None described platypnoea (dyspnoea on standing). The heart rate was consistently higher in the erect posture: 74 (29%) had a postural orthostatic tachycardia of $\geq 20$ min$^{-1}$, and in 25 (10%) this exceeded 30 min$^{-1}$. Orthostatic tachycardia was more pronounced in PAVM patients than controls without orthodeoxia (age-adjusted coefficient 5.5 (95% Cls 2.6, 8.4) min$^{-1}$, p<0.001). For PAVM patients, the age-adjusted pulse rise was 0.79 min$^{-1}$ greater for every 1% greater drop in oxygen saturation on standing (p<0.001). In contrast to the postural orthostatic tachycardia syndrome, in this population, there was a trend for more pronounced orthostatic tachycardia to be associated with better exercise tolerance.

To the editor

Pulmonary arteriovenous malformations (PAVMs) result in hypoxaemia due to right-to-left shunting. Recent studies highlight that chronic hypoxaemia in iron-replete patients leads to secondary erythrocytosis which preserves arterial oxygen content (CaO$_2$). Both shunt fraction and hypoxaemia severity may increase acutely on standing, a phenomenon ascribed to basally situated PAVMs. Platypnoea-orthodeoxia (dyspnoea and arterial desaturation on standing) has been described, particularly in patients with patent foramen ovale. However, platypnoea was not our experience in the PAVM population, suggesting they may be able to compensate for acute falls in CaO$_2$. The goal of the study was to quantify orthodeoxia and examine potential compensatory mechanisms to facilitate provision of appropriate information to PAVM patients.

The study was ethically approved by the Hammersmith, Queen Charlotte’s, Chelsea, and Acton Hospital Research Ethics Committee (LREC 2000/5764). Full methods are presented in the online supplementary data supplement. In all, 258 consecutive patients with CT-proven PAVMs were prospectively and newly recruited (2005–2013) and evaluated as described. Pulse and oxygen saturation (SaO$_2$) were measured by pulse oximetry in supine and erect postures for 10 min. Exercise capacity was stratified to a modified Medical Research Council (MRC) dyspnoea scale, with individuals classified as grade 1a if they participated in intense sporting activity at least three times per week.

Full patient demographics are presented in online supplementary Table 1. Ages ranged from 16 to 90 (median 48) years. A total of 89 (34.5%) were male. For 239
(92.6%), PAVMs were attributable to hereditary haemorrhagic telangiectasia (HHT). Overall, 50/221 (22.6%) were obese with a body mass index > 30. Comorbidities were more common in patients with higher grade dyspnoea (see online supplementary Figure 1). Replicate SaO2 and pulse values demonstrated high within-patient reproducibility (see online supplementary Table 2).

Overall, erect SaO2 was significantly lower than supine SaO2 (figure 1A). In 75/257 (29%) patients, the SaO2 fell by at least 2% on standing compared with the equivalent supine reading. A smaller fall of 1%–2% was present in a further 54/257 (21%) patients. None of these patients reported dyspnoea on standing (platypnoea), although one was unable to complete the 10 min standing due to dizziness. However, a further 24/257 (9%) patients reported dyspnoea on standing (platypnoea), although one was unable to complete the 10 min standing due to dizziness. As expected, obese patients had lower supine SaO2 for their erect SaO2, and correspondingly less evidence of orthodeoxia (see online supplementary Figure 4).

Sudden falls in SaO2 reduce CaO2 per unit blood volume. However, there was a consistent increase in heart rate on standing (figure 1B). Orthostatic tachycardia was more pronounced in PAVM patients than 40 controls (figure 1C, and online supplementary Figures 2 and 3), and in patients exhibiting greater falls in SaO2/CaO2 (see online supplementary Figure 5). Postural orthostatic tachycardia is normally viewed in a detrimental manner, but in this study, more marked orthostatic tachycardia was observed in patients with better exercise tolerance (figure 1D), whether analysed in five groups as shown, or in three groups of athletes (grade 1a), normal (grade 1b) and all dyspnoeic patients (grades 2–4).

To conclude, we provide an extensive consecutive series demonstrating that orthodeoxia is common in PAVM patients, though may be masked by obesity. Exuberant postural orthostatic tachycardia may be part of acute compensatory mechanisms that maintain tissue oxygen delivery when CaO2 falls suddenly on standing, and is associated with better exercise tolerance in PAVM patients.

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Contributors VS generated the 2005–2010 database and analysed data. BC generated the 2005–2010 database. HM, HCT, LCW and JS performed pulmonary function measurements. HRT and AJW generated the control and HHT databases. CLS reviewed all patients, contributed to database generation, performed all statistical analyses, generated the figures/tables and wrote the article. All authors reviewed and approved the final article.

Competing interests None.

Ethics approval Hammersmith, Queen Charlotte’s, Chelsea, and Acton Hospital Research Ethics Committee (LREC 2000/5764).

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Additional material is published online only. To view please visit the journal online (http://dx.doi.org/10.1136/thoraxjnl-2014-205289).


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METHODS:

Study populations
258 consecutive patients with radiologically-proven PAVMs were prospectively and newly recruited June 2005-January 2013. Demographics are provided in Table1. The series excludes any patient seen between May 1999-May 2005, findings for which have been previously reported, and precipitated the current series. Full details of the structured clinic assessments are provided elsewhere. All data reported are from the presentation assessment. Oxygen saturations (SaO2) and heart rate were measured by pulse oximetry (Ohmeda Biox 3900, Boulder, Colorado) while breathing room air. Additionally, 221 PAVM patients had same day spirometry evaluations, and 230 PAVM patients had same day blood tests: full blood counts were measured on XE Series Analysers (Sysmex, UK) and biochemical indices including serum iron, transferrin saturation index (Tf/SI) and ferritin Ci1600 Architect Analysers (Abbott Diagnostics, Ireland).

Control groups were generated retrospectively from other individuals who had been evaluated in the clinical service between April 2000 and September 2013. Controls were selected if they had no evidence of PAVMs on a dedicated thoracic CT scan, and no current/recent pregnancy or concurrent cardiorespiratory pathology or treatments anticipated to influence SaO2 or pulse. Inclusion within these groups was determined by final diagnostic status, blinded to postural measurements. The main control group comprised 40 individuals (24 women, and 16 men aged 19-66 (median 45.5)ys in whom PAVMs and/or HHT had initially been considered diagnostic possibilities. They had undergone standard SaO2/CT evaluations as part of clinical work up. Subsequent investigations had revealed alternate causes for their symptoms (haemoptysis, stroke, transient ischaemic attack), with HHT either not suspected clinically, or not present in their branch of the family by pedigree or mutational studies. A second control group comprised 87 individuals who had definite hereditary haemorrhagic telangiectasia (HHT) but no evidence of PAVMs on dedicated thoracic CT scans. These 53 women and 34 men ranged in age from 19-79 (median 44)ys.

Study methodologies
The potential imprecision of pulse oximetry measurements is well recognised. To improve precision, measurements were made continuously for 10 minutes in both supine and erect postures, recorded at one minute intervals. All except one individual with PAVMs completed the full evaluation. Individual SaO2 and heart rate measurements at minutes 7, 8, 9 and 10 were compared to evaluate intrapatient variability, and demonstrated high reproducibility (Table 2). For the evaluations presented in this study, the mean values from minutes 7, 8, 9 and 10 were used for final comparisons, and to generate the differences in SaO2 or pulse between supine and erect postures. Data from one control outlier (change in pulse of 25min^-1 recorded as a fall on standing) was excluded.

The reported exercise tolerance for patients reviewed between 2005-2010 was assigned to the Medical Research Council (MRC) Dyspnoea scale, blinded to all other patient parameters. Normal individuals with no dyspnoea except on extreme exertion were classified as Grade 1a if they were highly athletic individuals participating in intense sporting activity at least three times per week. Grade 1b represented normal (dyspnoeic only on strenuous exertion); Grades 2-5 progressively lower exercise tolerance. The primary clinician’s awareness of interim analyses in late 2010 implied these data could not be assigned to the 2010-2013 cohort in an unbiased manner. Arterial oxygen content (CaO2) on air was calculated by 1.34*haemoglobin *SaO2, where 1.34mls is the empirically determined amount of oxygen carried per gram of haemoglobin.

STATA IC version 12 (Statacorp, Texas) and GraphPad Prism 5 (Graph Pad Software Inc, San Diego) were used to calculate distributions of participant-specific variables, to perform comparisons between groups, and to generate graphs. Two group comparisons were by Spearman rank or Mann Whitney; three group repeated measures comparisons by Friedman with post-test Dunns corrections. Univariate and multivariate regression analyses were performed in STATA IC version 12 (Statacorp, Texas).
RESULTS:

Table 1: Demographics of the 258 PAVM Patients

<table>
<thead>
<tr>
<th>Binary variables</th>
<th>N §</th>
<th>Present</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hereditary haemorrhagic telangiectasia</td>
<td>258</td>
<td>239</td>
<td>92.6</td>
</tr>
<tr>
<td>Gender (female)</td>
<td>258</td>
<td>169</td>
<td>61.5</td>
</tr>
<tr>
<td>Ever smoked</td>
<td>240</td>
<td>87</td>
<td>36.3</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Continuous variables</th>
<th>N §</th>
<th>Range</th>
<th>Median</th>
<th>IQR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (yr)</td>
<td>258</td>
<td>16-90</td>
<td>48</td>
<td>36, 61</td>
</tr>
<tr>
<td>Body mass index (kg/m²)</td>
<td>220</td>
<td>16.6-46.3</td>
<td>26.3</td>
<td>23.0, 29.7</td>
</tr>
<tr>
<td>FEV1 (L)</td>
<td>221</td>
<td>1.0-5.2</td>
<td>2.9</td>
<td>2.4, 3.6</td>
</tr>
<tr>
<td>FEV1 (% predicted*)</td>
<td>221</td>
<td>36-127</td>
<td>94</td>
<td>83, 104</td>
</tr>
<tr>
<td>FEV1/VC (%)</td>
<td>221</td>
<td>38-99</td>
<td>78</td>
<td>73, 83</td>
</tr>
<tr>
<td>SaO₂ supine at presentation (%)</td>
<td>257</td>
<td>81.5-100</td>
<td>95.3</td>
<td>92.8, 96.5</td>
</tr>
<tr>
<td>Pulse supine at presentation (min⁻¹)</td>
<td>257</td>
<td>45-110</td>
<td>74</td>
<td>66, 81</td>
</tr>
<tr>
<td>SaO₂ erect at presentation (%)</td>
<td>257</td>
<td>78-99</td>
<td>95</td>
<td>91.8, 96.3</td>
</tr>
<tr>
<td>Pulse erect at presentation (min⁻¹)</td>
<td>257</td>
<td>52.3-142</td>
<td>89</td>
<td>80, 99</td>
</tr>
<tr>
<td>Change in SaO₂ on standing (%)</td>
<td>257</td>
<td>-11 to 7.5</td>
<td>-1</td>
<td>-2.3, 0.5</td>
</tr>
<tr>
<td>Change in pulse on standing (min⁻¹)</td>
<td>257</td>
<td>-2.5 to 49</td>
<td>14.25</td>
<td>10.5, 21.3</td>
</tr>
<tr>
<td>Haemoglobin (g/dL)</td>
<td>230</td>
<td>5.9-20.9</td>
<td>14.1</td>
<td>12.7, 15.5</td>
</tr>
<tr>
<td>Serum iron (µmol/L)</td>
<td>214</td>
<td>0-64</td>
<td>14</td>
<td>7, 19</td>
</tr>
<tr>
<td>Transferrin saturation index (TfSI, %)</td>
<td>214</td>
<td>0-79</td>
<td>23</td>
<td>12.32</td>
</tr>
<tr>
<td>Ferritin (ug/L)</td>
<td>176</td>
<td>0-409</td>
<td>30</td>
<td>15, 63</td>
</tr>
<tr>
<td>Supine CaO₂ (mls/dl)</td>
<td>229</td>
<td>7.7-24.8</td>
<td>18.2</td>
<td>16.1, 19.6</td>
</tr>
<tr>
<td>Erect CaO₂ (mls/dl)</td>
<td>229</td>
<td>7.6-22.9</td>
<td>17.9</td>
<td>16.1-19.3</td>
</tr>
<tr>
<td>Change in CaO₂ on standing</td>
<td>229</td>
<td>-2.8 to 1.11</td>
<td>-0.16</td>
<td>-0.47, 0.07</td>
</tr>
</tbody>
</table>

§N, number of datasets; values <258 imply that data was not available for a subgroup of patients. Note one patient could not complete the 10 minutes standing, and their incomplete pulse oximetry assessments were not included in the final calculations. IQR, interquartile range. FEV1, forced expiratory volume in one second. VC, vital capacity. CaO₂: arterial oxygen content on air, calculated by SaO₂*Hb*1.34/100.⁶

Table 2: Variability of pulse oximetry measurements in 257 PAVM patients

<table>
<thead>
<tr>
<th>Overall (mean (SD))</th>
<th>4 replicate measurements</th>
</tr>
</thead>
<tbody>
<tr>
<td>SaO₂ erect (%)</td>
<td>93.3 (4.6)</td>
</tr>
<tr>
<td>SaO₂ supine (%)</td>
<td>94.5 (3.0)</td>
</tr>
<tr>
<td>Pulse erect (min⁻¹)</td>
<td>89.9 (14.2)</td>
</tr>
<tr>
<td>Pulse supine (min⁻¹)</td>
<td>73.4 (11.1)</td>
</tr>
</tbody>
</table>

The first column presents the mean and standard deviation for SaO₂ and pulse in the two postures overall. The second column presents the standard deviations (SD) of the four replicate values for SaO₂ and pulse in the two
postures, calculated for each individual dataset within the respective group. Note that one patient was unable to complete measurements due to dizziness.

**Dyspnoea in 165 PAVM patients by modified MRC dyspnoea grade**

The dypsnoea grading system for the 165 patients first assessed between 2005-2010 was derived from the MRC dyspnoea scale. The distributions of oxygen saturation (SaO₂) and arterial oxygen content (CaO₂) across these dyspnoea grades are presented elsewhere: There was no clear relationship with SaO₂, but patients with higher dyspnoea grades had lower CaO₂ (defined as SaO₂* haemoglobin*1.34/100).

There was also a striking correlation with comorbidities unrelated to PAVMs (online supplementary Figure 1). For these 165 patients, 15 (9.1%) had significant co-existing disease, including eight (4.8%) with obstructive spirometry due to either asthma or COPD; one with emphysema and previous bullectomy; two with severe pulmonary arterial hypertension; one with obstructive sleep apnoea, and three with congestive cardiac failure or severe cardiac valvular disease. No PAVM patient had severe (Grade 4/5) dyspnoea without significant co-existing cardio-pulmonary disease (online supplementary Figure 1).

**Figure 1: Dyspnoea in 165 PAVM patients by modified MRC dyspnoea grade**

![Dyspnoea Distribution Graph]

**Legend:** Grey bars indicate individuals with significant co-existing cardiopulmonary disease. Note that Grade 1 (normal) is separated into 1a (highly athletic individuals participating in intense sporting activity at least three times per week), and 1b (normal). Grades 2-5 represent dyspnoea up a hill (Grade 2); after a mile on the flat (Grade 3); after a few minutes (~100m, Grade 4), and at rest or on minimal effort (Grade 5).
Postural changes in $\text{SaO}_2$ in individuals with and without CT evident PAVMs.

Patients with CT-proven PAVMs were more hypoxaemic than controls in both postures (Figure 2A). There was no trend for the control or HHT groups to exhibit orthodeoxia (Figure 2B).

**Figure 2: Postural $\text{SaO}_2$ assessments in PAVM patients and controls**

Legend: Comparison of $\text{SaO}_2$ values in patients with and without PAVMs: PAVM patients (n=257), HHT patients with no CT scan evidence of PAVMs (n=87) and controls (n=40). **A** Absolute $\text{SaO}_2$ values in supine and erect postures. **B** Postural changes. Error bars represent median and interquartile range. P values were calculated by Kruskal Wallis and Dunn’s post test. ***: p<0.001, ns: not significant.

Postural tachycardia in individuals with and without CT evident PAVMs.

The supine heart rate was no different in patients with and without CT-proven PAVMs (Figure 3A). However, the pulse was consistently higher after assuming the erect posture in all three groups (Figure 3A, Figure 3B), particularly in younger individuals: On average for each year of life added, the postural tachycardia diminished by 0.61 min$^{-1}$ in controls (p=0.011), 0.57 min$^{-1}$ in HHT patients with no evidence of PAVMs on CT (p=0.001), and 0.75 min$^{-1}$ in the PAVM group (p<0.0001). Age alone explained only 18.7% of the variance in postural tachycardia in PAVM patients. Overall postural tachycardia was more pronounced in PAVM patients than in controls (Figure 3B).

**Figure 3: Postural pulse assessments in PAVM patients and controls**

Legend: Three way comparison of heart rates (pulse) in PAVM patients and control groups. Comparison of **A** absolute pulse and **B** postural change in pulse in PAVM patients (n=257), HHT patients with no CT scan
evidence of PAVMs (n=87) and controls (n=40). Error bars represent median and interquartile range. P values were calculated by Kruskal Wallis and Dunn’s post test. **: p<0.01, ns: not significant.

**Obesity and postural changes in PAVM patients**

In the general population, obese individuals have smaller total lung capacity attributed to greater compression of the chest wall and reduced diaphragmatic expansion. Variable reports on SaO$_2$ have been described. In the current study, PAVM patients with higher body mass index (BMI) tended to have lower supine SaO$_2$ for their erect SaO$_2$, and correspondingly less evidence of orthodeoxia (Figure 4A). Patients with higher BMI also demonstrated less pronounced postural tachycardia (Figure 4B). The associations remained evident once adjusted for age and gender (data not shown).

**Figure 4: Postural data in 220 PAVM patients stratified by body mass index (BMI).**

![Graph A: Change in SaO$_2$ on standing. Linear regression r$^2$ = 4.4%, p = 0.0017.](image1)

**Legend:** A) Change in SaO$_2$ on standing. Linear regression r$^2$ = 4.4%, p = 0.0017. B) Change in pulse on standing. Linear regression r$^2$ = 4.9%, p = 0.0010.

**Acute hypoxaemia and postural tachycardia in PAVM patients**

Patients who demonstrated greater orthodeoxia tended to have more pronounced postural orthostatic tachycardia (Figure 5). The associations remained evident once adjusted for age, body mass index, haemoglobin, and all other parameters in online supplementary Table 1 (data not shown).

**Figure 5: Postural tachycardia in 257 PAVM patients stratified by acute oxygen changes**

![Graph A: Change in SaO$_2$ on standing.](image2)

**Legend:** A) Change in SaO$_2$ on standing. r$^2$ = 9.4%, p < 0.0001. B) Change in arterial oxygen content (CaO$_2$) on standing (mL/dL).
**Legend:** Change in pulse on standing stratified by A) Change in SaO$_2$ on standing; B) Change in arterial oxygen content (CaO$_2$) on standing. The crude regression coefficients and p values are displayed.

**DISCUSSION**

The questions addressed by this study were whether orthodeoxia was commonly present in PAVM patients, whether patients were symptomatic as a result, and if not, why this might be.

Where PAVMs are present in dependent portions of the lung, gravitational forces will increase flow through the right-to-left shunts, a phenomenon first noted by auscultation, and quantified using nuclear medicine perfusion scans using $^{99m}$Tc-labelled albumin microspheres. In clinical practice, the oxygen saturation (SaO$_2$) provides a convenient, non invasive assessment of the right-to-left shunt. Orthodeoxia, a fall in arterial oxygenation on assuming the erect posture, is described in the PAVM literature, though its frequency has been questioned. Orthodeoxia is also described for patients with patent foramen ovale and other intracardiac shunts. We were intrigued by the apparent prevalence of the platypnoea-orthodeoxia syndrome in patients with patent foramen ovale (PFO), as despite clear evidence of orthodeoxia, the PAVM patients that we have reviewed do not report dyspnoea on standing, despite acute postural assessments being an integral part of their clinical assessment at our institution.

We have recently highlighted one of the primary mechanisms that help sustain tissue oxygen delivery in chronically hypoxaemic PAVM patients--a graded secondary erythrocytotic response which resolves following correction of hypoxaemia by embolisation of PAVMs. As a result, despite often substantial increments in SaO$_2$, the majority of patients report no difference in exercise tolerance when re-evaluated several months after embolisation.

Acute orthodeoxia could not be accompanied by a change in haemoglobin. We hypothesised that compared to the patients described in the literature with platypnoea-orthodeoxia syndrome, PAVM patients might be better able to utilise physiological compensatory methods. Considerations of the initial response to haemorrhage suggested an increase in heart rate might be one such potential compensatory mechanism, and the data in the current study would support such a hypothesis. Standing reduces venous return due to gravitational forces, with baroreceptors usually implicated in the activation of the sympathetic nervous system responsible for postural tachycardia. The PAVM patients remind of an additional potential increment from chemoreceptor activation. This appears to be relevant to the acute change: we have recently demonstrated no difference in heart rate at peak exercise in PAVM patients chronically adapted to their hypoxaemia, compared to the same patients retested several months after embolisation.

Why the PAVM patients might be better able to mount compensatory responses to acute orthodeoxia than the symptomatic patients described in the literature is not clear. Increasing age was associated with less pronounced orthostatic tachycardia in all groups. The continuous nature of the right-to-left shunt, in contrast to PFO where right-to-left shunting is intermittent, may facilitate adaptation. We cannot exclude HHT-specific enhancement of orthostatic tachycardia, and this might be suggested in view of the intermediate phenotype of the HHT patients compared to PAVM patients and controls (Figure3). However, it is important to note that from published data, and the endoglin (HHT1 bias) of the study population, it would be anticipated that at least half of this HHT ‘non PAVM’ group would have right-to-left shunting that would have been detectable by contrast echocardiography but not associated with PAVMs sufficiently large to be visualised on CT scans.

Finally, the delineation of changes in a disease population often leads to suggestions that the assessments might be helpful in diagnostic or screening strategies. As emphasised in 1999, and subsequently reiterated, the detection of hypoxaemia by age-defined PaO$_2$, or SaO$_2$ from postural assessments, is insufficiently sensitive to be used as the sole diagnostic screening test for PAVMs. SaO$_2$ measurements do have substantial clinical utility in the serial investigations required in follow up of individual patients. It remains to be seen whether the evaluation of postural tachycardia will
be similarly beneficial, or merely a tool to better understand the physiological mechanisms that permit tolerance of acute changes in arterial oxygen content.

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