Management of pulmonary arteriovenous malformations in pulmonary hypertensive patients: a pressure to embolise?

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Pulmonary arteriovenous malformations (PAVMs) carry significant risks [1, 2]. Pulmonary arterial blood passing through these right-to-left shunts cannot be oxygenated, leading to hypoxaemia, with arterial oxygen tension and arterial oxygen saturation (\(S_aO_2\)) inversely related to the size of the shunt fraction. The fragile wall of the PAVM sac may rupture, leading to haemoptysis or haemothorax [3]. In addition, the absence of a filtering capillary bed allows particulate matter to reach the systemic circulation where it impacts in other capillary beds, including those of the cerebral circulation, resulting in embolic cerebrovascular accidents and cerebral abscesses. In a recent series of 219 patients that corrected for ascertainment bias, the rates for ischaemic stroke and cerebral abscess were 11.3% and 9%, respectively, with risks unrelated to PAVM severity or symptoms [4]. Treatment of PAVMs by embolisation is an effective means of reducing lifetime risks of paradoxical embolic stroke and cerebral abscess [4], improving oxygenation [1, 5, 6], and treating PAVM-induced haemoptysis [3, 7]. Embolisation usually carries minimal risk in expert hands and, as a result, is generally offered to patients with PAVMs of a size amenable to embolisation, irrespective of the presence of respiratory symptoms.

PAVMs themselves have a lower vascular resistance than that of the surrounding normal lung, due to the absence of a microvascular network of capillary vessels. However, most individuals with PAVMs have underlying hereditary haemorrhagic telangiectasia (HHT), now recognised to carry an independent risk of pulmonary hypertension (PH) [8]. HHT is inherited as an autosomal dominant trait, and is most commonly caused by mutations in the genes for endoglin (HHT type 1) or activin receptor-like kinase-1 (HHT type 2) [9]. The condition, which affects one in 5–8,000 Europeans [10, 11], is classically diagnosed by the consensus criteria of nosebleeds, mucocutaneous telangiectasia, visceral arteriovenous malformations (AVMs) and family history [12]. HHT-related PH occurs in two main settings: 1) a post-capillary PH in the context of high-output cardiac failure secondary to hepatic AVMs with elevated pulmonary artery wedge pressure and normal or near-normal pulmonary vascular resistance (PVR); and 2) a true pulmonary arterial hypertension (PAH) phenotype [13] characterised by normal pulmonary artery wedge pressure and markedly raised PVR [6]. While severe PH in unselected HHT cohorts is rare [6, 14], and PAVMs and PH are more common in different HHT genotypes (type 1 for PAVMs, type 2 for both forms of PH [15]), a small group of HHT patients have both PAVMs and severe PH.

The presence of PAVMs does not modify the treatment of PH. The question is, should the presence of PH modify recommended treatments for PAVMs and, if so, for which patients?

**PH MODIFICATION OF THE NATURAL HISTORY OF PAVMS**

As noted in table 1, PAVM risks differ in the presence of PH. Recent data demonstrate that the risk of ischaemic stroke is substantially lower in PAVM patients with higher pulmonary artery pressure (\(P_{pa}\)); the hazard ratio fell with each 1 mmHg rise in mean \(P_{pa}\) [4]. Conversely, the presence of elevated PAVM sac perfusion pressures, either due to PH or in the setting of a systemic arterial supply to PAVM sacs [19], appears to place patients at higher risk of life-threatening haemorrhage from PAVMs, as has been described in the present issue of the European Respiratory Review [16]. While it is unclear whether PH precipitates the development of PAVMs in susceptible individuals, there are rare case reports suggesting PH may be associated with rapid growth of PAVMs [18].

**PH MODIFICATION OF PAVM TREATMENT-RELATED RISKS**

PAVM embolisation abolishes low resistance pathways for pulmonary blood flow and, therefore, might be expected to elevate \(P_{pa}\). Isolated case reports, including fatalities [20], highlight the potential risks of such an increase. However,
a recent study of 43 patients demonstrated that for most individuals, successful PAVM embolisation resulting in improved SaO₂ did not result in a sustained or acute change in Ppa [6]. In half of the cases, including one individual with post-capillary PH due to left ventricular disease, embolisation led to a fall in Ppa, which was attributed to the described reduction in cardiac output [5].

PAVM patients clearly differ in their haemodynamic responses to embolisation. Several authors have pointed out that deleterious rises may relate to underlying hepatic AVMs [6, 16, 20]. Temporary balloon occlusion of the PAVM before definitive embolisation has been suggested in order to identify which patients are at risk of such an increase [16, 20]; however, in the one reported individual, test balloon occlusion did not predict a substantial 22 mmHg rise in mean Ppa following definitive embolisation [6].

INTERPRETATION

So, should embolisation of PAVMs be performed in patients with severe pre-existing PH? The most common indications for PAVM embolisation are to reduce the risk of paradoxical embolic stroke and, for individuals with hypoxaemia, to improve dyspnoea and exercise tolerance. Since the risk of paradoxical embolic stroke is substantially lower in individuals with higher Ppa [4], and symptomatic relief from dyspnoea should not be expected for patients with PH and SaO₂ >90%, as was the case in all four patients with severe PH in a recent study [6], Shovlin et al. [6] suggested that for the majority of patients with pre-existing severe PAH the risks of PAVM embolisation outweigh potential benefits. In the present issue of the Review, the case report by Montani et al. [16] highlights the risk of later development of haemothorax but, as has been reported in the pregnancy setting (where 1.02% (95% confidence interval 0.13–1.92%) of 484 PAVM/HHT pregnancies resulted in a major PAVM bleed), there is no reported evidence that embolisation of PAVMs for individuals who have not experienced a complication reduces the risk of later haemorrhage [7]. Thus, in general, for patients with severe PH, we would not interpret the risk–benefit considerations in favour of PAVM embolisation.

The most difficult judgements relate to individuals with severe pulmonary hypertension and active major haemoptysis or haemothorax in whom pulmonary arteriovenous malformation-related haemorrhage may be a terminal event. In a previous study of pregnant females (without pulmonary hypertension) [7], and in pulmonary hypertension cases known to the current authors, there was time for emergency intervention to be performed after the onset of herald symptoms (haemoptysis or chest pain). In such an emergency setting, patients and their physicians may consider the risks of precipitating a further increase in pulmonary artery pressure justifiable. In the present authors’ opinion, even if test balloon occlusion is performed and appears satisfactory, all patients should be warned of a potential fatal increase in pulmonary artery pressure. Further data are required to assess whether particular categories of pulmonary hypertension carry lower risks.

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