

## An Unwelcome Embrace Adverse Pulmonary–Aortic Interactions in Pulmonary Hypertension

Michael A. Quail, MBChB, PhD; Vivek Muthurangu, MD

Vascular remodeling in pulmonary hypertension (PH) causes increased ventricular afterload, resulting in right ventricular failure and high mortality.<sup>1</sup> Afterload is the mechanical impedance imposed by the arterial tree, against which the ventricle must work to eject blood. Despite the tendency to conflate afterload with blood pressure, it cannot be reduced to a single number or variable. Instead it is more appropriately defined in terms of pressure-flow relations.<sup>2,3</sup>

### See Article by Schäfer et al

Afterload is divided into several steady and pulsatile components, including vascular resistance, total arterial compliance, characteristic impedance, and wave reflections. Each of these has been shown to be abnormal in the pulmonary circulation in PH resulting in increased right ventricular afterload.<sup>4–8</sup> However, abnormalities of the systemic vasculature that result in increased left ventricular (LV) afterload have been relatively understudied in PH.

In this issue of *Circulation: Cardiovascular Imaging*, Schäfer et al<sup>9</sup> demonstrate elevated proximal aortic pulse wave velocity and other metrics of local aortic stiffness in children with PH. This elevation was strongly associated with the degree of dilatation of the main pulmonary artery (PA) and proximal branches, suggesting that the aortic root is mechanically constrained by the embrace of the PAs.

In addition, the authors noted a modest correlation between aortic relative area change:  $([Area_{max} - Area_{min}] / Area_{max})$  and the main PA:aorta ratio with decreased LV ejection fraction. They went on to suggest that increased aortic stiffness because of pulmonary arterial constraint resulted in LV dysfunction in this patient group. This is an intriguing hypothesis, and we think that it is worth exploring the mechanisms by which this could occur.

When the LV ejects, it must accelerate blood into the aorta, overcoming a combination of inertance and vessel

compliance. This is the characteristic impedance ( $Z_c$ ) of the vessel, and  $Z_c$  governs ventricular load in early systole. The  $Z_c$  is directly related to pulse wave velocity ( $c$ ) as follows:

$$Z_c = \rho c / A_d, \quad (1)$$

where  $A_d$  is vessel diastolic area, and  $\rho$  is blood density. As can be seen from Equation 1, ventricular load in early systole is increased in vessels with smaller areas and elevated pulse wave velocity.<sup>10</sup> Therefore, Schäfer et al<sup>9</sup> results imply increased aortic  $Z_c$  in pediatric PH patients and increased LV afterload. However, elevated  $Z_c$  occurs in a range of conditions, including normal aging, and, at this degree of elevation, would not be expected to be associated with impaired LV function. Indeed, diastolic aortic area was not different between the groups, and pulse wave velocity was not associated with LV ejection fraction, suggesting that it may not be a prime candidate for the association.

Given that there was no evidence of increased systemic vascular resistance or total aortic compliance between PH patients and controls, alternative causes of LV dysfunction must be sought. The other element of afterload that is often neglected is wave reflections. Several types of wave reflections occur in the systemic vasculature and can be identified by wave intensity analysis in the time domain.<sup>11</sup> Backwards compression waves, which decelerate flow and increase pressure increase LV afterload and have been associated with heart failure events in adults.<sup>12</sup> In addition, we have recently shown that backwards compression waves arising in the repaired aortic arch are an important determinant of LV hypertrophy after coarctation repair.<sup>13</sup> Reflections arise in the vasculature at areas of impedance mismatch (eg, branches, changes in wall stiffness, or caliber). In this study, the locally increased stiffness and reduced aortic expansion at the level of PA branches is a likely area of impedance mismatch. Therefore, it is possible that pediatric PH patients have increased backwards compression waves, which increase LV afterload and result in myocardial dysfunction.<sup>14</sup> Evaluation of this hypothesis would be challenging, requiring simultaneous measurement of pressure and flow in the proximal aortic root. Alternatively, a non-invasive approach using magnetic resonance area and flow data may be possible, but the close proximity of the reflecting site would require high temporal resolution approaches that may not be generally available.<sup>7</sup>

Of course, PA size does increase with worsening PH, and the causes of LV dysfunction—collinear with relative area change and main PA:aorta ratio—but not operating via afterload-related mechanisms, are possible. Alternative explanations might include adverse ventricular–ventricular interactions or reduced myocardial perfusion because of increased

The opinions expressed in this article are not necessarily those of the editors or of the American Heart Association.

From the Centre for Cardiovascular Imaging, Institute of Cardiovascular Science, University College London, United Kingdom, and National Paediatric Pulmonary Hypertension Centre, Great Ormond Street Hospital for Children, London, United Kingdom.

Correspondence to Vivek Muthurangu, MD, Level 6 Old Nurses Home, Cardiorespiratory Unit Great Ormond Street Hospital for Children, Great Ormond St, London WC1N 3JH, United Kingdom. E-mail v.muthurangu@ucl.ac.uk

*Circ Cardiovasc Imaging*. 2017;10:e006063.

DOI: 10.1161/CIRCIMAGING.117.006063.

© 2017 American Heart Association, Inc.

*Circ Cardiovasc Imaging* is available at  
<http://circimaging.ahajournals.org>

DOI: 10.1161/CIRCIMAGING.117.006063

right atrial pressure or coronary compression. Larger studies are required to ascertain if these factors are also important.

In conclusion, magnetic resonance flow imaging provides an excellent opportunity to assess both pulmonary and systemic vascular hemodynamics. The right ventricle in PH has been shown to negatively impact the function of the LV through ventricular interactions.<sup>15</sup> However, the observation of adverse PA and aortic interaction is intriguing and may explain some of the symptomatic consequences of PH in children.

### Disclosures

Dr Quail is funded by the 2016 British Heart Foundation-Fulbright award (FS/16/28/32327).

### References

- van de Veerdonk MC, Kind T, Marcus JT, Mauritz GJ, Heymans MW, Bogaard HJ, Boonstra A, Marques KM, Westerhof N, Vonk-Noordegraaf A. Progressive right ventricular dysfunction in patients with pulmonary arterial hypertension responding to therapy. *J Am Coll Cardiol*. 2011;58:2511–2519.
- Milnor WR. *Hemodynamics*. 2nd ed. Baltimore, MD: Williams & Wilkins; 1989.
- Nichols WW, O'Rourke MF, Vlachopoulos C. *McDonald's Blood Flow in Arteries: Theoretical, Experimental and Clinical Principles*. 6th ed. London, United Kingdom: Hodder Arnold; 2011.
- Pandya B, Quail MA, Steeden JA, McKee A, Odille F, Taylor AM, Schulze-Neick I, Derrick G, Moledina S, Muthurangu V. Real-time magnetic resonance assessment of septal curvature accurately tracks acute hemodynamic changes in pediatric pulmonary hypertension. *Circ Cardiovasc Imaging*. 2014;7:706–713. doi: 10.1161/CIRCIMAGING.113.001156.
- Muthurangu V, Atkinson D, Sermesant M, Miquel ME, Hegde S, Johnson R, Andriantsimiavona R, Taylor AM, Baker E, Tulloh R, Hill D, Razavi RS. Measurement of total pulmonary arterial compliance using invasive pressure monitoring and MR flow quantification during MR-guided cardiac catheterization. *Am J Physiol Heart Circ Physiol*. 2005;289:H1301–H1306. doi: 10.1152/ajpheart.00957.2004.
- Lankhaar JW, Westerhof N, Faes TJ, Marques KM, Marcus JT, Postmus PE, Vonk-Noordegraaf A. Quantification of right ventricular afterload in patients with and without pulmonary hypertension. *Am J Physiol Heart Circ Physiol*. 2006;291:H1731–H1737. doi: 10.1152/ajpheart.00336.2006.
- Quail MA, Knight DS, Steeden JA, Taelman L, Moledina S, Taylor AM, Segers P, Coghlan GJ, Muthurangu V. Noninvasive pulmonary artery wave intensity analysis in pulmonary hypertension. *Am J Physiol Heart Circ Physiol*. 2015;308:H1603–H1611. doi: 10.1152/ajpheart.00480.2014.
- Vonk Noordegraaf A, Westerhof BE, Westerhof N. The relationship between the right ventricle and its load in pulmonary hypertension. *J Am Coll Cardiol*. 2017;69:236–243. doi: 10.1016/j.jacc.2016.10.047.
- Schäfer M, Ivy DD, Abman SH, Barker AJ, Browne LP, Fonseca B, Kheyfets V, Hunter KS, Truong U. Apparent aortic stiffness in children with pulmonary arterial hypertension: existence of vascular interdependency? *Circ Cardiovasc Imaging*. 2017;10:e005817. doi: 10.1161/CIRCIMAGING.116.005817.
- Chirinos JA, Segers P. Noninvasive evaluation of left ventricular afterload: part 2: arterial pressure-flow and pressure-volume relations in humans. *Hypertension*. 2010;56:563–570. doi: 10.1161/HYPERTENSIONAHA.110.157339.
- Parker KH, Jones CJ. Forward and backward running waves in the arteries: analysis using the method of characteristics. *J Biomech Eng*. 1990;112:322–326.
- Chirinos JA, Kips JG, Jacobs DR Jr, Brumback L, Duprez DA, Kronmal R, Bluemke DA, Townsend RR, Vermeersch S, Segers P. Arterial wave reflections and incident cardiovascular events and heart failure: MESA (Multiethnic Study of Atherosclerosis). *J Am Coll Cardiol*. 2012;60:2170–2177. doi: 10.1016/j.jacc.2012.07.054.
- Quail MA, Short R, Pandya B, Steeden JA, Khushnood A, Taylor AM, Segers P, Muthurangu V. Abnormal wave reflections and left ventricular hypertrophy late after coarctation of the aorta repair [published online ahead of print January 23, 2017]. *Hypertension*. doi: 10.1161/hypertensionaha.116.08763. <http://hyper.ahajournals.org/content/early/2017/01/23/HYPERTENSIONAHA.116.08763> (accessed 2/5/2017).
- Kobayashi S, Yano M, Kohno M, Obayashi M, Hisamatsu Y, Ryoke T, Ohkusa T, Yamakawa K, Matsuzaki M. Influence of aortic impedance on the development of pressure-overload left ventricular hypertrophy in rats. *Circulation*. 1996;94:3362–3368.
- Knight DS, Steeden JA, Moledina S, Jones A, Coghlan JG, Muthurangu V. Left ventricular diastolic dysfunction in pulmonary hypertension predicts functional capacity and clinical worsening: a tissue phase mapping study. *J Cardiovasc Magn Reson*. 2015;17:116. doi: 10.1186/s12968-015-0220-3.

KEY WORDS: Editorials ■ aorta ■ blood pressure ■ hemodynamics ■ pulmonary artery ■ pulmonary hypertension

## An Unwelcome Embrace: Adverse Pulmonary–Aortic Interactions in Pulmonary Hypertension

Michael A. Quail and Vivek Muthurangu

*Circ Cardiovasc Imaging.* 2017;10:

doi: 10.1161/CIRCIMAGING.117.006063

*Circulation: Cardiovascular Imaging* is published by the American Heart Association, 7272 Greenville Avenue, Dallas, TX 75231

Copyright © 2017 American Heart Association, Inc. All rights reserved.

Print ISSN: 1941-9651. Online ISSN: 1942-0080

The online version of this article, along with updated information and services, is located on the World Wide Web at:

<http://circimaging.ahajournals.org/content/10/2/e006063>

**Permissions:** Requests for permissions to reproduce figures, tables, or portions of articles originally published in *Circulation: Cardiovascular Imaging* can be obtained via RightsLink, a service of the Copyright Clearance Center, not the Editorial Office. Once the online version of the published article for which permission is being requested is located, click Request Permissions in the middle column of the Web page under Services. Further information about this process is available in the [Permissions and Rights Question and Answer](#) document.

**Reprints:** Information about reprints can be found online at:  
<http://www.lww.com/reprints>

**Subscriptions:** Information about subscribing to *Circulation: Cardiovascular Imaging* is online at:  
<http://circimaging.ahajournals.org/subscriptions/>