P1.4: HEMODYNAMICS OF PULMONARY HYPERTENSION: APPLICATION OF THE RESERVOIR-WAVE APPROACH

Anukul Ghimire*, Mads Andersen, Lindsay Burrowes, J. Christopher Bouwmeester, Andrew Grant, Israel Belenkie, Nowell Fine, Barry Borlaug, John Tyberg

To cite this article: Anukul Ghimire*, Mads Andersen, Lindsay Burrowes, J. Christopher Bouwmeester, Andrew Grant, Israel Belenkie, Nowell Fine, Barry Borlaug, John Tyberg (2015) P1.4: HEMODYNAMICS OF PULMONARY HYPERTENSION: APPLICATION OF THE RESERVOIR-WAVE APPROACH, Artery Research 12:C, 4–4, DOI: https://doi.org/10.1016/j.artres.2015.10.198

To link to this article: https://doi.org/10.1016/j.artres.2015.10.198

Published online: 7 December 2019
isometric conditions. Pacing experiments done in rodents and humans show that arterial compliance is highly cyclic stretch frequency-dependent. The Rodent Oscillatory Tension Set-up to study Arterial Compliance (ROTSAC) is a mouse developed organ bath that clamps aortic segments (width 2mm, diameter 0.5-3mm) to imposed preloads at physiological rates up to 600bpm. The technique enables us to acquire pressure-diameter loops (derived from simultaneous force-displacement measurements) and calculate biomechanical parameters such as Peterson’s modulus (Ep) and compliance. To our knowledge, this is the first set-up that facilitates the study of active vessel wall components, physiological stretch frequency and pressure variations and its effect on the biomechanical properties of the aorta.

Arterial stiffness is generally considered to be determined mainly by structural components. However, using this device, we were able to show by isobaric determination of compliance and Ep while changing pressure and vascular smooth muscle cells (VSMCs) tone—that active vessel wall components are highly important in determining biomechanical properties of the aorta. Ep values for WT mouse aorta (350.3 ± 8.2 mmHg) were in accordance with literature data and increased 29% upon a rise in diastolic pressure of 40 mmHg, while isobaric Ep increased 47% upon maximal contraction of the VSMCs. We believe that this set-up can significantly contribute to a better understanding how active vessel wall components influence arterial stiffening, hypertension and its associated cardiovascular complications.

**P1.4**
**HEMODYNAMICS OF PULMONARY HYPERTENSION: APPLICATION OF THE RESERVOIR-WAVE APPROACH**

Anukul Ghimire 1,2, Mads Andersen 3,3, Lindsay Burrowes 1, J. Christopher Bouwmeester 3, Andrew Grant 3, Israel Bellenkie 1, Nowell Fine 3, Barry Borlaug 3, John Runciman 2, Luis Arroyo 1

Using the reservoir-wave approach, previously we characterized pulmonary vascular mechanics with multiple interventions in a canine model. In the present study, we measured high-fidelity pulmonary arterial (PA) pressure, Doppler flow velocity, and pulmonary capillary wedge pressure in 11 patients referred for evaluation of exertional dyspnea. The analysis was performed using the reservoir-wave approach; wave intensity analysis was subsequently utilized to characterize the PA wave pattern. Our objective was to identify specific abnormalities associated with pulmonary hypertension. Seven patients with varying PA pressures had reduced pulmonary vascular conductance (i.e., the amount of flow that the lungs can accept per pressure gradient), suggesting that these patients might benefit from pulmonary vasodilator therapy, some even in the absence of markedly elevated PA pressure.

Right ventricular (RV) performance was assessed by examining the work done by the wave component of systolic PA pressure. Wave work, the non-recoverable energy expended by the RV to eject blood, varied directly with mean PA pressure. Wave pressure was partitioned into two components: forward-travelling and reflected backward-travelling waves. Among patients with lower PA pressures, we found pressure-decreasing backward waves that aided the RV during ejection, as previously reported in normal experimental animals. Among patients with higher PA pressures, we detected pressure-increasing backward waves that impede RV ejection.

We conclude that it is important to measure pulmonary vascular conductance to properly assess the pulmonary vasculature. The reservoir-wave approach and wave intensity analysis may prove to be valuable tools to evaluate RV performance and may facilitate development of therapeutic strategies.

**P1.5**
**AGE AND HYPERTENSION STRONGLY REDUCE AORTIC VISCO-ELASTIC PROPERTIES IN RATS AT BASAL AND MATCHED BLOOD PRESSURE LEVELS**

George Lindsey 1*, Christophe Ragionet, Stefano Chimenti, Nicole Villeneuve, Christine Vayssettes-Courchay

IDRS, 91430, France

Age and hypertension are major causes of large artery stiffening, a cardiovascular risk factor for heart and kidney damage. Long-term hypertension induces vascular remodeling, accelerating vascular aging. The aged Spontaneously Hypertensive Rat (SHR) model is recognized for human cardiovascular pathology but discrepancies are apparent in studies of arterial stiffness.

We performed experiments using a robust aortic visco-elasticity analysis via echotomodelling in 20 (n=6) 26-week-old SHR (n=8), and respective control. Wistar Kyoto rats (WKY, n=6-6) at basal and matched levels of blood pressure (BP). After anesthesia (pentobarbital), abdominal aortic diameter and pressure were recorded and BP was decreased by clonidine i.v. At basal BP, aortic pulse distension, compliance, distensibility (AD) and wall viscosity (AWJ) were reduced and stiffness index increased with age and hypertension and further altered with age + hypertension. BP being adjusted to 130 and 100 mmHg between groups, there was no difference between 20w old SHR and WKY but importantly the age effect was maintained in both WKY and SHR and accentuated by hypertension in old rats. At 130 mmHg, AD = 24.2±1.1 vs 20w WKY, 20.3±1.8 in 20w SHR, 12.4±1.3 in 80w WKY and 6.1±0.7 in 80w SHR; AWJ = 58.5±5, 58.9±9, 29±1 and 10±2 in the same groups. In conclusion reduced distensibility i.e. stiffening due to age is clearly shown here in both WKY and SHR as well as the effect of hypertension in aged rats. It will allow new investigations of the mechanisms and possible effect of drugs on aortic stiffness.

**P1.8**
**DEVELOPMENT OF A TECHNIQUE FOR DETERMINATION OF PULMONARY ARTERY PULSE WAVE VELOCITY IN HORSES**

Gonçalo Silva 1,2, Bruce Guest 2, Diego Gomez 1, Martine McGregor 3, John Bunciman 1, Luis Arroyo 1

1Department of Clinical Studies, Ontario Veterinary College, University of Guelph, Guelph, Ontario, Canada
2School of Engineering, University of Guelph, Guelph, Ontario, Canada

Calcification of the tunica media of the main pulmonary arteries has been observed in a large proportion of young racehorses. In humans, medial calcification is the most important cause of increased arterial stiffness, and has been implicated in the pathogenesis of microvascular diseases. Pulse wave velocity (PWV) is a marker of arterial stiffness. This study aimed to develop a technique for determination of pulse wave velocity of the main pulmonary arteries of horses.

Six healthy adult horses were sedated, and continuously monitored with echocardiography during the procedure. The pulmonary artery (PA) trunk was cannulated via right heart catheterization, with a catheter introducer sheath (9Fr x 100cm). Introducer placement was guided with echocardiography. A custom-made dual pressure sensor catheter (PSC) (7Fr x 170cm) was inserted through the introducer sheath, and into one of the main branches of the PA. The position of the PSC in one of the main branches of the PA was confirmed with thoracic radiography and pressure measurements were recorded. The time delay of the pulse waves between the two sensors was used to calculate PWV.

The PSC placement was successfully achieved in all horses (6/6), without significant complications, aside from transient arrhythmias. The catheter was more commonly located on the left PA (5/6). The mean ±SD PWV was 3.0±1.3m/s.

This study demonstrated the feasibility of a technique to determine PA-PWV in standing horses. The technique developed may allow further investigation of the effect of calcification of large pulmonary arteries in the development of microvascular disorders in horses.

**P1.9**
**PLATELET-LOCALIZED FXI PROMOTES A GLYCOPROTEIN Ibα DEPENDENT FEEDBACK LOOP IN ARTERIAL HYPERTENSION AND VASCULAR INFLAMMATION**

Jeremy Lagrange 1,2, Sabine Kossmann 1,2, Moritz Ehliken 1,2, Brett Monia 3, Wolfram Ruf 1, Philip Wenzel 1,2

1Center for Thrombosis and Hemostasis, Mainz, Germany
2Department of Medicine 2 University Medical Center, Mainz, Germany
3ISIS Pharmaceuticals Inc, Gazelle Court Carlsbad, USA

Background: Interactions of platelets, leukocytes and the vessel wall play pivotal roles in activating coagulation and precipitating thrombosis. High levels of angiotensin II (ATII) cause arterial hypertension by a complex inflammatory pathway requiring leukocyte recruitment and reactive oxygen species production within the vessel wall.

Objective: The aim of this work was to explore the role of platelet glycoprotein Ibα dependent thrombin-FXI feedback loop in arterial hypertension.

Methods: FXII−/−, FXI−/−, and hIL-4R/Ibα and 5/6 nephrectomized rats were used for this study. Mice where treated with ATII (1mg/kg) −/− for 7