Analysis of Aortic Energetics from Pulse Wave Examination in a Porcine Study of Septic Shock

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Abstract: Aortic pressure (Pao) can be separated into two components representing the wave phenomena of flow, called excess pressure (Pex), and the storage capacity of the elastic arteries, called reservoir pressure (Pwk). Subject-specific aortic models, identified from measurements from a porcine study on septic shock, were used to calculate the hydraulic work associated with the excess and reservoir pressures (Wex, Wwk). Changes in these energies were compared to a metric derived from left ventricular pressure-volume analysis. Total aortic work (Wao=Wex+Wwk) compared well to clinically assessed left ventricular work (R²=0.88). However, only a weak relationship of R²=0.24 was found between ventricular arterial coupling (Ees/Ea) and Wex/Wwk. Although a strong relationship (R²=0.76) was noticed between the inverse of afterload (1/Ea) and Wex. As septic shock progressed, a drop in Wwk was seen, indicating the arterial system loses its ability to store the stroke volume (SV) from the ventricle for release during diastole, resulting in a flattening of the diastolic pressure. These results indicate that one of the main reasons left ventricular afterload decreases during septic shock is because arterial system loses its ability to act as a storage reservoir.

Keywords: Physiological models, septic shock, hemodynamic monitoring, parameter identification.

1. INTRODUCTION

Arterial pressure and cardiac output (CO) are two of the main measures of which intensive care cardiovascular management is based on. However, recently, the efficiency of hemodynamic monitoring to affect outcome has been questioned (Pinsky, 2003). The positive impact of CO and/or stroke volume (SV) monitoring has not been proven (Mutoh et al., 2007, Pinsky, 2007). Furthermore, although continuous blood pressure monitoring of the arterial waveform is commonplace, the most commonly used blood pressure indices are calculated from discrete measurements of systolic and diastolic pressure, neglecting the morphological information available from the shape of the arterial pulse. All these results suggest these measurements are underutilised in the intensive care unit (ICU).

This study looks at utilising SV and the shape of the aortic pressure waveform to better understand the dynamics of arterial flow and ventricular arterial coupling. It is based on the work of (Wang et al., 2003, Davies et al., 2010, Tyberg et al., 2009) where the aortic pressure is separated into two components representing the reservoir pressure and the excess pressure (Fig. 1). These components were identified using measurements from a porcine study of septic shock (Lambermont et al., 2006) to create subject-specific models of aortic blood flow.

It is hypothesised that the work of the excess pressure represents the contractile state of the ventricle, and reservoir pressure component is the energy that needs to be overcome for forward flow. Hence, the ratio of the excess pressure and reservoir work may represent a metric of ventricular arterial coupling. To test this hypothesis the modelled hydraulic energies of these components were compared to metrics obtained from left ventricular pressure-volume (P-V) loops, to understand their relationship to ventricular arterial (de)coupling during septic shock.

2. METHODS

2.1 Porcine trials and measurements

All procedures and protocols used were approved by the Ethics Committee of the Medical Faculty at the University of Liege, Belgium. Experiments were performed on 4 healthy pure pietrain pigs weighing between 20-30kg. The animals were premedicated, anesthetised, and ventilated as explained in (Lambermont et al., 2006). The animals received a 0.5-mg/kg endotoxin infusion over 30 minutes (from T0 to T30) to model septic shock. Micro-tipped catheters were used to record continuous hemodynamic measurements during the study. Every 30 minutes (T0 to T120) 6-12 central aortic pressure waveforms (Pao) were recorded. At the same time, descending left ventricular pressure-volume (P-V) loops (Pbl, Vbl) were stored during transient occlusion of the inferior
Describing represents the compliance of the whole arterial tree. It is proposed that during diastole, the reflected or backwards travelling waves in the aorta are assumed to be negligible. Hence, in Equation (5), to and $P_0$ represent the time and pressure at the start of ejection and $\tau$ the time decay constant. Note that during diastole $Q_{in} = 0$, as it is assumed there is no inflow during diastole.

In this paper, the reflected or backwards travelling waves in the aorta are assumed to be negligible. Hence, $Q_{in}$ is assumed to be linearly proportional to $P_{ex}$, as indicated by the work of (Wang et al., 2003);

$$P_{ex}(x,t) = Q_{in}(t)R_{prox}$$  \hspace{1cm} (6)

where $R_{prox}$ represents the proximal resistance of the aorta. The substitution of (5) and (6) into (1) then gives:

$$P_{ao}(x,t) = P_{wk}(t) + P_{ex}(x,t)$$ \hspace{1cm} (1)

An ODE describing the windkessel pressure is derived considering a mass balance across the aortic:

$$\frac{dP_{wk}(t)}{dt} = \frac{Q_{in}(t) - Q_{out}(t)}{C}$$ \hspace{1cm} (2)

where $Q_{in}$ and $Q_{out}$ are the flows in and out of the aorta and $C$ represents the compliance of the whole arterial tree. Describing $Q_{out}$ using Ohm’s law gives:

$$Q_{out} = \frac{P_{wk}(t) - P_{ao}}{R}$$ \hspace{1cm} (3)

In Equation 3, $R$ is the resistance of the peripheral circulation and $P_{ao}$ is the asymptotic pressure approached during decay in diastole. Substituting this definition of $Q_{out}$ into (1) gives:

$$\frac{dP_{wk}(t)}{dt} + \frac{P_{wk}(t) - P_{ao}}{RC} = \frac{Q_{in}(t)}{C}$$ \hspace{1cm} (4)

with the general solution defined:

$$P_{wk}(t) - P_{ao} = (P_0 - P_{ao})e^{-\frac{t}{R\tau}} + e^{-\frac{t}{R\tau}} \int_{t_0}^{t} \frac{Q_{in}(t')}{C} e^{\frac{t'}{R\tau}} dt'$$ \hspace{1cm} (5)

In Equation (5), $t_0$ and $P_0$ represent the time and pressure at the start of ejection and $\tau = R\tau$, the time decay constant. Note that during diastole $Q_{in} = 0$, as it is assumed there is no inflow during diastole.

In this paper, the reflected or backwards travelling waves in the aorta are assumed to be negligible. Hence, $Q_{in}$ is assumed to be linearly proportional to $P_{ex}$, as indicated by the work of (Wang et al., 2003):

$$P_{ex}(x,t) = Q_{in}(t)R_{prox}$$ \hspace{1cm} (6)

where $R_{prox}$ represents the proximal resistance of the aorta. The substitution of (5) and (6) into (1) then gives:

$$P_{ao}(t) = P_{ao} + (P_0 - P_{ao})e^{-\frac{t}{\tau}} + e^{-\frac{t}{\tau}} \int_{t_0}^{t} \frac{Q_{in}(t')}{C} e^{\frac{t'}{\tau}} dt' + Q_{in}(t)R_{prox}$$ \hspace{1cm} (7)

Equation (7) is now analogous to a 3-element windkessel model, where the added resistance, $R_{prox}$, represents the flow resistance of the large arteries.

Fig. 1. Example of aortic pressure separated into reservoir pressure ($P_{wk}$) and excess pressure ($P_{ex}$) in a healthy subject.
2.4 Data analysis

Hemodynamic measurements are presented as mean +/- standard error of the mean (SEM). A paired sample t-test was used to check temporal variance over T0 - T60 to analyse the effect of the endotoxin intervention. P<0.05 was considered statistically significant result. Windkessel models were fitted to 36 sets of measurements from the 4 pigs, over all time points.

Hydraulic work ($W_{ao}$) was calculated for each measurement by summing the work of the reservoir and excess pressure components of the aortic model ($W_{ao} = W_{wk} + W_{ex}$). Where, $W_{ex}$ and $W_{wk}$ are evaluated from the start of ejection ($t_0$) to the end of systole ($t_{es}$). Left ventricular work ($W_{lv}$) was calculated from the area enclosed by the measured P-V loop. $W_{ao}$ and $W_{lv}$ are calculated:

$$ W_{ao} = W_{wk} + W_{ex} = \int_{t_0}^{t_{es}} P_{wk}(t)[Q_{in}(t) - Q_{on}(t)]dt' $$

$$ + \int_{t_0}^{t_{es}} P_{ex}(t)Q_{in}(t)dt' $$

$$ W_{lv} = \int_{V_0}^{V} P_{lv}(t)dv' $$

The ratio of the $W_{ex}$ to the $W_{wk}$ was analysed and compared to clinical gold standard ventricular arterial coupling metric, $E_{es}/E_{sa}$, calculated from ventricular P-V loop analysis, as shown in Fig. 2. End systolic elastance ($E_{es}$) is defined as the gradient of the linear least squares fit to the end systolic points in a series of descending P-V loops, representing the left ventricular contractility. Left ventricular afterload ($E_{so}$) is the gradient of the line from the origin in Fig. 2 to the end systolic point of the first P-V loop, recorded before vena cava occlusion, a highly invasive manoeuvre.

3. RESULTS

The windkessel model was fitted to the 36 sets of measurements from the 4 pigs. In this study, each animal acts as their own control with baseline (T0) measurements reflecting the undiseased state of the pig. Statistically significant temporal changes (P<0.005) were noticed in the measured mean arterial pressure (MAP), left ventricular end diastolic volume (LVEDV), heart rate (HR), and maximum left ventricular pressure ($P_{lv,max}$) over T0-T60, underlining the pathological changes occurring in the animals due to the endotoxin infusion. A summary of the main hemodynamic measurements is seen in Fig. 3.

![Fig. 3. Evolution of mean arterial pressure (MAP), left ventricular end diastolic volume (LVEDV), stroke volume (SV), heart rate (HR), and maximum left ventricular pressure ($P_{lv,max}$) measurements during the trials. * indicates P<0.05 for expected temporal changes over T0-T60 due to the induction of septic shock. Data presented as mean +/- SEM.](image)

3.1 Comparison of ventricular and arterial work

To verify the aortic model, the hydraulic energy of the aortic flow ($W_{ao}$), calculated from model, was compared to the measured left ventricular work ($W_{lv}$). The comparison between the hydraulic work and the measured P-V loops, is shown in Fig. 4. $W_{ao}$ matched $W_{lv}$ to a bias and 2 standard
deviations of -0.15 J +/- 0.13 J (as seen on Fig. 4.), and with a correlation coefficient of $R^2 = 0.88$. These results indicate that $W_{ao}$ underestimates ventricular work, but, more importantly, closely follows the changes in P-V area that occur during the trials.

$W_{ao} = 0.64X - 0.01$

$R^2 = 0.88$

$N = 36$

3.2 Effects of septic shock on aortic energetics and volume

The effects of septic shock on the aortic model were analysed by tracking the changes in $W_{ain}$, $W_{ex}$, and $W_{wk}$ throughout the duration of the experimental study. In this analysis, $W_{ex}$ can be interpreted as the work required to overcome the resistance of the large arteries, whereas $W_{wk}$ reflects the maximum potential energy stored in the arterial reservoir due to elastic distension. The change in magnitudes of $W_{ao}$, $W_{wk}$, and $W_{ex}$ (averaged across the 4 pigs) at 30 minute intervals during the experimental study are shown in Fig. 5. Initially, at T0, $W_{wk}$ makes up the majority of $W_{ao}$. However, after the endotoxin infusion, the magnitude of $W_{wk}$ decreases, and $W_{ex}$ increases to a point where they are approximately equal by the end of the experiment.

A decrease in $W_{wk}$ indicates that a smaller proportion of the SV is being stored in the arterial system, due to distension of elastic arteries, for release during diastole. Fig. 6 reinforces this point, showing the percentage of SV stored in the arterial reservoir, averaged across the 4 pigs. This ratio drops from a baseline level of 59.2% to 48.6% by the end of the study.

3.3 Estimation of ventricular arterial coupling

It was hypothesised that the ratio of $W_{ex}$ to $W_{wk}$ may be an indicator of ventricular arterial coupling, given that $W_{ex}$ is a function of the flow energy ejected from the ventricle, and $W_{wk}$ represents the work required to distend the arterial volume. Hence, $W_{ex}/W_{wk}$ was compared to $E_{es}/E_{a}$ to check this claim, as shown in Fig 7. Analysis of Fig. 7 showed a very weak relationship between these indices, with correlation of $R^2 = 0.24$, indicating they are not directly related, and proving our initial hypothesis wrong.

On further analysis, it was noticed that $W_{ex}$ is strongly related to the inverse of $E_{a}$ ($R^2 = 0.76$), as seen in Fig. 8. However, changes $W_{ex}$ or $W_{wk}$, or any combination of these two indices, were not reflective of changes in left ventricular contractility ($E_{es}$). Hence, demonstrating that the energetics of the aortic model are only capable of representing changes in the state of
left ventricular afterload and not changes in left ventricular contractility.

\[ Y = 1.47X + 0.28 \]
\[ R^2 = 0.24 \]
\[ N = 36 \]

Fig. 7. Comparison of pressure-volume loop derived ventricular vascular coupling \((E_v/E_a)\) with the ratio of excess work to windkessel work \((W_{ex}/W_{wk})\) derived from the aortic model.

\[ Y = 3.57X + 0.13 \]
\[ R^2 = 0.76 \]
\[ N = 36 \]

Fig. 8. Comparison of the inverse of afterload \((1/E_a)\) calculated from pressure-volume loop analysis with excess work \((W_{ex})\) derived from the aortic model.

4. DISCUSSION

A model of aortic blood flow, based on the works of (Davies et al., 2010, Tyberg et al., 2009, Wang et al., 2003), was identified from measurements from an experimental animal study on septic shock. Aortic models were fitted using only measurements of the aortic pressure waveform and SV, removing the reliance on knowing \(Q_{in}\) which can be highly invasive or difficult to measure clinically. In this study, the models were successfully fitted to 36 sets of these measurements and the energetics of these models were analysed.

The modelled derived \(W_{ao}\), representing the total hydraulic work of flow in the aorta, tracked changes in the measured \(W_{ao}\) well with \(R^2 = 0.88\), although \(W_{ao}\) underestimated \(W_{ao}\) on average by -0.13J. The bias in \(W_{ao}\) was probably a result of the extra work required by the ventricle to push the flow through the aortic valve, which is not accounted for in the aortic model. Fisher and Wheatley (1988) recorded an energy loss of around 0.16-0.33J for flow through bio-prosthetic porcine aortic valves using SV between 60-80mls. Indicating that a bias of -0.13J, given that measured SVs in this study are around half those used in (Fisher and Wheatley, 1988), is a reasonable approximation for aortic valve energy loss. This result verifies the accuracy of the fitted models.

The identified aortic models also appeared capable of capturing the effects septic shock on the pig’s hemodynamics. As the state of the pigs worsened, due to the inflammatory effects of the endotoxin, modelled systemic resistance decreased substantially and arterial compliance increased, as seen in Fig. 9. The drop in resistance was generally larger than the increase in compliance causing the decay constant, \(\tau\), to decrease as \(\tau = RC\). The combination of these factors caused the aortic reservoir to charge and discharge quicker than it would have done for a healthy subject. The volume in the arterial reservoir could not accumulate faster than it was discharging, resulting in a flattening in the \(P_{wk}\) component of the aortic pressure. Hence, in the subjects with a low decay constant when normalised by heartbeat \((\tau / T)\), the shape of the aortic pressure begins to look similar to the shape of \(Q_{in}\), as seen in Fig. 10, with a noticeable flattening of the diastolic pressure observed when compared to Fig. 1. This effect is responsible for the relative decrease in the proportion of \(W_{ao}\) compared to \(W_{ao}\) and \(W_{ex}\) that occurs during the study, as shown in Fig. 5. These factors could explain why left ventricular afterload \((E_a)\) decreases during septic shock as arterial pressure can never build up during the heartbeat because less energy and volume are stored in the arterial reservoir.

Fig. 9. Change in systemic resistance, R (top panel), aortic compliance, C (middle panel), and ratio of the decay constant to heartbeat period, \(\tau / T\) (bottom panel) during the trials. Data presented as mean +/- SEM.
Fig. 10. Example of aortic pressure, separated into reservoir pressure ($P_{wk}$) and excess pressure ($P_{ex}$) components, 180 minutes (T210) after endotoxin infusion.

The ratio of $W_{ex}/W_{wk}$ did not relate well to $E_{ex}/E_{a}$ as seen in Fig. 7. However, $W_{ex}$ by itself could be a useful metric for describing acute changes in afterload ($E_{a}$) due to septic shock as indicated by the strong correlation shown in Fig. 8.

The main, and most controversial, assumption made during the fitting process is that $P_{ex}$ is proportional to $Q_{in}$, and hence, the two can be represented by a purely resistive relationship ($P_{ex} = R_{prox}Q_{in}$). This assumption is based on the work of (Wang et al., 2003), which shows that the magnitudes of backwards travelling waves in the aorta are minimal. The authors feel justified in using this assumption in this study for two reasons: 1) the model was only fitted to central aortic pressure measurements, where backwards travelling waves are small during systole (Aguado-Sierra et al., 2008), due to the central aorta’s distance from major reflection sites; and 2) in sepsis vascular tone decreases, reducing the production of reflective waves (O'Rourke and Yaginuma, 1984). However, the authors realise the assumption may not be valid in subjects with high arterial stiffness, such as with hypertensive subjects.

5. CONCLUSIONS

This analysis of SV and morphological features of the aortic pressure waveform suggests that one of main reasons left ventricular afterload decreases during septic shock is because arterial reservoir loses its ability to store hydraulic energy in the form of stressed volume during systole. However, a larger more comprehensive study is required to validate the method and results of this paper.

REFERENCES


