Measurement of cardiac output by use of noninvasively measured transient hemodilution curves with photoacoustic technology

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Abstract: We present the theoretical basis and experimental verification for cardiac output measurements using noninvasively measured hemodilution curves afforded with an indicator dilution technique and the emerging photoacoustic technology. A photoacoustic system noninvasively tracks a transient hemodilution effect induced by a bolus of isotonic saline as an indicator. As a result, a photoacoustic indicator dilution curve is obtained, which allows to estimate cardiac output from the developed algorithm. The experiments with a porcine blood circulatory phantom system demonstrated the feasibility of this technology towards the development of a noninvasive cardiac output measurement system for patient monitoring.

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OCIS codes: (170.0170) Medical optics and biotechnology; (170.3890) Medical optics instrumentation; (170.5120) Photoacoustic imaging; (170.4580) Optical diagnostics for medicine; (170.3880) Medical and biological imaging.

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#203412 - $15.00 USD  Received 24 Dec 2013; revised 13 Feb 2014; accepted 28 Feb 2014; published 7 Apr 2014 (C) 2014 OSA 1 May 2014 | Vol. 5, No. 5 | DOI:10.1364/BOE.5.001445 | BIOMEDICAL OPTICS EXPRESS 1445
1. Introduction

Cardiac output (CO) is an established important physiological parameter that enables physicians to optimize fluid status in hemodynamically instable and critically ill patients, such as to improve the patient outcomes [1, 2]. Conventional technologies for measuring CO involve using indicator dilution techniques [3–6]. In these techniques, a bolus of indicator, such as cold saline, is typically introduced into a patient’s central circulation through a central venous catheter or pulmonary arterial catheter (PAC). An indicator dilution curve is measured downstream with a highly invasive femoral arterial catheter [7] or the distal port of the PAC [8] as the indicator is washed out in the patient’s circulation system, which is used to calculate the patient’s CO [3, 4, 6]. However, since the established indicator dilution methods for the estimation of CO are highly invasive and potentially cause medical complications, there is an increasing need for developing noninvasive devices [9, 10] to assess CO in critically ill patients. In this paper, we report breakthroughs of developing a noninvasive cardiac output measurement technology by combining the indicator dilution technique with the emerging noninvasive photoacoustic technology [11–14].

In this proposed technology, a small bolus of room temperature isotonic saline is injected quickly into the central circulation of a patient as the indicator, which induces a transient hemodilution effect. A photoacoustic sensor equipped with a single ultrasound sensor is placed downstream over a tissue bed with a superficial artery. The sensor noninvasively measures the transient hemodilution effect from an artery as the indicator travels across the sensor site. As the indicator is washed out in the circulation system, a hemodilution (indicator dilution) curve is noninvasively measured, which is used to derive hemodynamic parameters, such as CO.

2. Theory

A photoacoustic signal is a linear function of a hemoglobin concentration in the blood when a tissue bed containing a superficial blood vessel is illuminated by a light source with a wavelength within the range of 600-1000 nm [15], where the product of the blood absorption coefficient and the vessel diameter is much less than 1 [16]. As a result, when a bolus of isotonic saline is injected into the blood stream, the hemoglobin concentration of the blood will be diluted, which induces photoacoustic signal variation, consequently resulting in an indicator dilution curve. When a bolus of an isotonic saline with amount of $V_I$ is instantaneously injected at $t = 0$ (i.e., the time of starting the injection is set to zero for convenience without losing the generality) into a circulation system, as the indicator is being washed out of the circulation system, CO can be calculated from the resultant indicator dilution curve by [3],

$$CO = \frac{V_I}{\int_0^{\Delta V} \frac{\Delta V_I(t)}{\Delta V} dt},$$

where $\Delta V$ and $\Delta V_I(t)$ are blood and isotonic saline volumes passing through the measurement point of the blood vessel during the unit time interval, $\Delta t$, respectively. Note that $\Delta V$ during $\Delta t$ is always constant for constant CO, which contains $\Delta V_I(t)$ for the case of indicator dilution. Equation (1) indicates that the whole isotonic saline injectate is
completely washed out of the circulation system as time goes to infinity assuming that no isotonic molecules interact with tissues outside the cardiovascular system.

As above mentioned, for a fixed amount of incident beam onto the measurement site, the photoacoustic signal is proportional to the total hemoglobin concentration in the unit blood volume \( \Delta V \). Therefore, for the stationary state before indicator injection, the photoacoustic signal measured from the targeted arterial vessel could be described by

\[
PA_b = K \frac{tHb_b}{\Delta V_b} + PA_b^0, \tag{2}
\]

where \( tHb_b \) indicates the total hemoglobin in the blood volume \( \Delta V_b \) associated with \( \Delta t \). The subscript \( b \) indicates the background that means before the indicator injection. The term \( K \) is the conversion factor from \( tHb_b/\Delta V_b \) to the photoacoustic signal, which is assumed to be a constant during the measurement of photoacoustic indicator dilution curves. The conversion factor accommodates systematic effects, such as a sensitivity of an ultrasound transducer and/or unknown blood vessel wall effect. If this measurement happens in a clinical situation, the factor might also include an effect of an unknown optical property of a background tissue bed. The term \( PA_b^0 \) represents the photoacoustic signal from all other photoacoustic sources insensitive to hemoglobin concentration variation induced by the injected indicator. At the measurement site after the injection, the amount of hemoglobin in \( \Delta V \) is decreased due to the added portion of the isotonic solution. Therefore, the PA signal variation due to injected isotonic solution can be described as

\[
PA(t) = K \frac{tHb_n(t)}{\Delta V_n(t) + \Delta V_i(t)} + PA_b^0, \tag{3}
\]

where \( \Delta V_n(t) + \Delta V_i(t) = \Delta V \) and \( \Delta V_n(t) \) is the volume of the blood not diluted by any portion of an injected isotonic solution. Because the hemoglobin concentration in pure blood is not changed before and after the indicator injection, it is valid to write

\[
\frac{tHb_b}{\Delta V_b} = \frac{tHb_n(t)}{\Delta V_n(t)}. \tag{4}
\]

Although the amount of volume of \( \Delta V_b \) is the same as that of \( \Delta V \) for the same blood vessel and constant CO, they contain different amount of hemoglobin. By substituting Eq. (4) to Eq. (3), the photoacoustic signal variation during the dilution process, \( PA(t) \) is

\[
PA(t) = K \frac{\Delta V - \Delta V_i(t)}{\Delta V \cdot \Delta V_b} \cdot tHb_b + PA_b^0. \tag{5}
\]

Considering Eq. (2), Eq. (5) is further developed to

\[
PA(t) = PA_b \left[ 1 - \alpha \frac{\Delta V_i(t)}{\Delta V} \right], \tag{6}
\]

where

\[
\alpha = \frac{PA_b - PA_b^0}{PA_b}. \tag{7}
\]

Integrating both sides of Eq. (6) in time with the consideration of Eq. (1) derives the blood flow rate as
In the derivations from Eqs. (5) to (8), it is assumed that \( PA_h \) and \( PA_b \) are stationary in time during photoacoustic indicator dilution measurements. In Eq. (8), both \( PA(t) \) and \( PA_b \) are measurable and the amount of isotonic injection \( V_i \) is also known. The term \( \alpha \) is the only unknown factor on the estimation of CO since it is very difficult to measure \( PA_h \), especially in a clinical situation. However, it is reasonable to assume that typically, the \( PA_h \) from the targeted blood vessel is negligible compared to the main photoacoustic signal with an optical wavelength (600-1000nm) where the dominant absorber is the hemoglobin. Therefore, it would be generally accepted that \( PA_h >> PA_b \), which means \( \alpha \approx 1 \). Therefore, we set \( \alpha \) to 1 for the estimation of CO from ex-vivo experimental data in this paper.

3. Experimental setup

Ex-vivo experiments were performed to demonstrate the concept described in the previous section. Figure 1 shows the schematic of the experimental setup for the measurement of a flow rate (i.e., cardiac output in a human cardiovascular circulatory system).

In these experiments, a high power pulsed laser diode at 905 nm (OSI Laser Diode Inc.: CVN 5S63) was used as the light source. The laser diode was driven with a pulse laser diode driver (Laser Component: LSP-40), running at the repetition rate of 3.2 KHz and the pulse width of 100 ns, which delivers a pulse energy of about 6.5uJ to the pig blood in a tube. A function generator (Agilent: 33522A) was used to generate 100 ns pulses to trigger the laser as well as to synchronize the data acquisition system. A photoacoustic sensor equipped with a commercial ultrasound transducer (Olympus Panametrics: NDT C382, 3.5 MHz) and fiber optics was integrated with a home-made spring loaded plastic sensor holder as shown in Fig. 1. In this sensor design, the transducer was resting on a spring, and its focus can be adjusted by turning the screw at the top of the sensor holder. A thin Rexolite disk with low ultrasound attenuation was used to seal the bottom of the plastic sensor holder. This special sensor design
not only allows the sensor holder to be filled with water as an ultrasound coupling medium, but also provides a solid ultrasound interface, which makes the sensor portable and easy to use for in-vivo testing with an ultrasound gel. A short plastic fiber was inserted through a hole in the wall of the plastic housing to deliver light to the phantom. A picture of the PA sensor for the bench-top experiment is shown in the inset of Fig. 1. Once the photoacoustic signal was detected by the PA detector, the signal was amplified with a pre-amplifier (Olympus: 5662) and digitized with a high speed digitizer (Gage: CS1642-128M), and acquired into a laptop with the LabVIEW program developed in-house. The acquired PA signals were averaged 800 times in real-time, and digitally band-pass filtered (0.5-5MHz) with the Labview program. The peak-to-peak voltages of the photoacoustic signals were calculated to derive the hemodilution curve.

In order to generate a reliable reference to validate the algorithm, the pump speed was calibrated for the selected flow rates of the pig blood. To obtain the calibration coefficient, the pig blood was first weighted with a digital scale (METTLER PM2000, resolution = 0.01 g) from 1.00 ml to 8.00 ml with an increment of 1.00 ml to obtain its density of (1.036 ± 0.003 g/ml) from the linear regression with the Pearson's correlation coefficient of 0.99998. The same blood was then pumped at discrete pump speed settings (12 to 28 with an increment of 4) to a beaker on the digital scale for 60 seconds and its whole weight was measured. The weight was converted to the volume with the previously measured blood density. Figure 2 shows the calibrated flow rates of the pig blood for the pump with pump speeds setting from 12 to 28.

In the experiment, the total volume of the pig blood was 27 ml (10 ml in the beaker plus 17 ml in the tubing). An indicator of 1.00 ml isotonic saline was rapidly injected into the beaker through the blood injection port. In clinical indicator dilution measurements for an adult, the amount of indicator injection could be up to 15ml and the blood volume contained in a heart is 250~300ml. Therefore, the indicator injection ratio in this ex-vivo experiment is similar with that in the clinical situation. When the 1.00 ml isotonic saline indicator was washing through the detecting site at the downstream tube, a photoacoustic hemodilution curve was obtained, which was used to estimate the flow rate. A 0.5% Intralipid solution (Liposym III 20%) was used to mimic skin tissue bed scattering and the tube was positioned about 3 mm below the surface of the Intralipid solution.

4. Results

Figure 3(a) shows an example of a photoacoustic signal measured by the ultrasound transducer focusing on the pig blood tube in the experimental setup of Fig. 1. Temporal variation of the amount of the signal becomes a photoacoustic indicator dilution curve. We averaged 400 photoacoustic signals for a single data point of indicator dilution curves, which means the acquisition rate for measuring the dilution curve is 8Hz for the 3.2kHz pulsed diode
laser. For measuring the photoacoustic signal variation, both peak-to-peak and the area under the signal were considered, the results of which are shown in Fig. 3(b). These dilution curves are normalized to 1.0 as the baseline for a comparison. As shown in Fig. 3(b), there is no much difference between two methods of peak-to-peak and area under the signal, so we consider peak-to-peak photoacoustic dilution curves for the estimation of CO.

![Image](image.png)

Fig. 3. (a) one example of photoacoustic signals from the pig blood tube in the experimental setup. (b) A normalized indicator dilution curves extracted from consecutively measured photoacoustic signals.

In the measured indicator dilution curves, photoacoustic signals are decreased when the diluted blood passes at the measurement site. The second valleys in Fig. 3(b) are caused by the recirculation of the injected isotonic saline indicator, as typically shown in a clinical situation. Because of the recirculation, it is impossible to completely measure the tail of the photoacoustic dilution curve for the first circulation. A typical approach for finding the complete dilution curve is to remove the recirculation portion of the curve, and curve-fit the first circulation curve using an exponential model [17], where it is assumed that the tail of dilution curves exponentially changes based on an appropriate model of vascular structures. For the exponential model to estimate CO in this paper, we considered a single blood container mimicking a heart chamber in a cardiovascular structure. Since an indicator dilution decreases the measured photoacoustic signals, the denominator of Eq. (8) indicates the area between the scaled photoacoustic dilution curve, $1 - PA(t)/PA_0$, and the normalized baseline 0, as shown in Fig. 4. Different from other previously known indicator dilution methods [6] for estimating CO, the normalization factor $PA_0$ in the integration of Eq. (8) is naturally obtained during the indicator dilution measurement process.

Figure 5 shows the estimated flow rates calculated using Eq. (8), where 5 different flow rates were tested. As mentioned before, we performed experiments with water and 0.5% Intralipid as the intermediate medium between the photoacoustic sensor and the pig blood tube, which imitate an ideal no-scattering medium and photon-scattering tissue bed, respectively. Although the experiment with Intralipid is more realistic, we conducted the experiment with water to validate the algorithm of Eq. (8) because the photoacoustic signal is greatly increased with almost no absorption in the water tissue bed, which minimizes the effect of α. As indicated in Fig. 5, the mean and standard deviation for differences between the estimated and actual values are $-0.61\text{ml/min}$ and $2.54\text{ml/min}$ for water, and $5.63\text{ml/min}$ and $3.87\text{ml/min}$ for Intralipid. The estimated flow rates for water are highly consistent with the actual values, but the results for 0.5% Intralipid are slightly over estimated. It is conjectured that the over-estimation may be caused by the Intralipid absorption, as observed during the experiments, that indicates $\alpha < 1$ for the actual situation, but we assumed $\alpha = 1$ in Eq. (8) for the estimation of flow rates. Because there is no clear boundary between $PA_0$ and
in Eq. (7) and the overall photoacoustic signal is the mixed one, as shown in Fig. 3(a), the contribution of $PA_0$ to $PA_b$ could be either negative or positive, which means $\alpha$ could be more or less than 1, especially, in a clinical measurement.

![Image](image_url)

**Fig. 4.** The red curve indicates the smoothed dilution signal after applying an averaging filter. The exponential method estimates the tail of the dilution curve that is shown as the blue curve. The integration in the denominator of Eq. (8) indicates the area under the scaled dilution curve.

![Image](image_url)

**Fig. 5.** Estimated flow rates from the theory of photoacoustic indicator dilution.

### 5. Conclusion and discussion

We presented a new concept to noninvasively measure cardiac output by combining a photoacoustic technique and indicator dilution method. The theory was developed to calculate cardiac output from the indicator dilution curves measured by the photoacoustic system. The estimated flow rates from the *ex-vivo* bench-top experimental data are consistent with actual values with acceptable errors, which validates the feasibility of this technology.
The measurement accuracy of this method and the impact of $\alpha$ have also been studied with \textit{in-vivo} swine model, and compared with transitional transpulmonary thermo-dilution technique. The results, which will be published elsewhere, demonstrated that this new technology is capable of achieving comparable accuracy with those of the traditional invasive methods.

To further improve the measurement accuracy, different indicators, such as an ICG dye, may be used to strengthen the dilution signal, reducing the adverse effect of the background noise signals. Alternatively, the system imaging resolution may be optimized to minimize the interference of the background signals. In addition, advanced methods for actively reducing or quantifying the amount of $\alpha$, thus improving the overall measurement performance could be developed in terms of hardware and software. Our future study will be focused on the pre-clinical experiments for this photoacoustic indicator dilution method considering the improvement of clinical measurement accuracy.

**Acknowledgment**

The authors thank Mr. Charles Haisley for his help on designing the PA sensor holder.