Review Article

Evaluation of Fluid Responsiveness: Is Photoplethysmography a Noninvasive Alternative?

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Background. Goal-directed fluid therapy reduces morbidity and mortality in various clinical settings. Respiratory variations in photoplethysmography are proposed as a noninvasive alternative to predict fluid responsiveness during mechanical ventilation. This paper aims to critically evaluate current data on the ability of photoplethysmography to predict fluid responsiveness.

Method. Primary searches were performed in PubMed, Medline, and Embase on November 10, 2011.

Results. 14 papers evaluating photoplethysmography and fluid responsiveness were found. Nine studies calculated areas under the receiver operating characteristic curves for $\Delta$POP ($>0.85$ in four, $0.75–0.85$ in one, and $<0.75$ in four studies) and seven for PVI (values ranging from $0.54$ to $0.98$). Correlations between $\Delta$POP/PVI and $\Delta$PP/other dynamic variables vary substantially.

Conclusion. Although photoplethysmography is a promising technique, predictive values and correlations with other hemodynamic variables indicating fluid responsiveness vary substantially. Presently, it is not documented that photoplethysmography is adequately valid and reliable to be included in clinical practice for evaluation of fluid responsiveness.

1. Introduction

Whether or not to administer intravenous (iv) fluid is a common, difficult, and controversial challenge in clinical practice. The main aim of fluid therapy during surgery or critical illness is to provide adequate tissue perfusion by increasing stroke volume (SV) or cardiac output (CO). Goal-directed fluid therapy aiming to increase oxygen (O$_2$) delivery reduces morbidity and mortality in various clinical settings [1–8]. Fluid therapy is guided by clinical variables, as well as static and dynamic variables. Clinical variables include blood pressure, heart rate, capillary refill time, skin turgor and diuresis, mixed venous oxygen saturation (SvO$_2$), lactate, pH, electrolytes, and creatinine/urea. Conventional static variables include central venous pressure (CVP) and pulmonary artery wedge pressure (PAWP), but these variables have proven less reliable than initially assumed to evaluate fluid responsiveness [8–10]. Dynamic variables include both SV-dependent and non-SV-dependent methods. The ideal new method should be accurate [11], easy to use, noninvasive, and widely available with minimal risk of complications. Potential clinical value also depends on reproducibility and predictive values compared to established methods.

Photoplethysmography (more specifically pulse oximetry plethysmographic waveform analysis) as a noninvasive tool in evaluation of fluid responsiveness was first described by Partridge [12] and has been extensively investigated. A pulse oximeter is a standard equipment for measuring arterial O$_2$ saturation, and further analysis of the photoplethysmographic signal can easily be implemented in clinical monitoring. This paper aims to critically evaluate current data on the ability of photoplethysmography to predict fluid responsiveness.

2. Methods

This paper is based on searches performed in PubMed, Medline, and Embase on November 10, 2011 with the following search criteria: “(pulse oximetry OR plethysmographic OR
Pleth variability index OR PVI) AND (fluid responsiveness OR (volume status))." The searches generated 217 hits. Papers were checked for relevant references and 22 [13-34] papers met the following inclusion criteria:

(1) reporting predictive values of ΔPOP and/or PVI after fluid challenges and/or reporting correlations between ΔPOP, PVI, and ΔPP,
(2) mechanically ventilated patients,
(3) written in English.

3. Results

3.1. Predictive Values of ΔPOP and PVI. 14 studies performed fluid challenges and these are summarized in Table 1. Patients were mechanically ventilated with tidal volumes of 6-10 mL/kg and investigated preoperatively (n = 6) [23, 26, 28, 30-32], perioperatively (n = 3) [21, 33, 34], postoperatively (n = 2) [25, 27], and in the intensive care unit (ICU) (n = 2) [22, 24]. One study included different groups of patients [29]. The pulse oximeter was placed on a finger in all papers, and in two papers it was also placed on the earlobe [30, 34].

Different types of pulse oximeters were used, summarized in Table 2. The number of patients included in each study varied substantially (n = 8-43). Registration periods were, with some exceptions [22, 23, 32], short (<1 min). Six studies did not indicate duration of the registration period [24, 28-31, 34]. Patients with arrhythmias were excluded in all papers except one, in which this was not explicitly stated [34]. Patients receiving vasoactive medication were included in four papers [24, 27, 29, 33] and excluded in four [23, 26, 31, 32]. Six papers did not indicate use of vasoactive medication [21, 22, 25, 28, 30, 34].

These data are summarized in Table 3. Fluid challenges were given as HES 6% (n = 9) [22-26, 28-31], "colloid fluid" (n = 3) [21, 33, 34], and NaCl 0.9% (n = 2) [27, 32]. Fluid volumes ranged from 250 mL to 1000 mL. Fluid responsiveness was defined as increased CO > 15% (n = 2) [25, 29], increased CI > 15% (n = 5) [22-24, 26, 30], ΔPP > 13% (n = 1) [27], increased SVI > 10% (n = 1) [21], increased SVI > 15% (n = 2) [28, 31], increased SV > 10% (n = 1) [34], increased SV > 15% (n = 1) [33], and aortic velocity-time integral (AVTI) > 15% (n = 1) [32].

Best cut-off values ranged from 8.8 to 15% for ΔAPP, 9.5 to 15% for ΔPOP, and 9.5 to 17% for PVI. CO was measured with thermodilution (n = 5) [21-23, 26, 30], echo Doppler (n = 6) [24, 29, 31-34], FloTrac/Vigileo (n = 1) [28], and intermittent thermodilution by pulmonary artery catheter (Vigilance monitor) (n = 1) [25].

Nine studies calculated areas under receiver operating characteristics curves (ROC curves) for ΔPOP [21-27, 32, 33]. It was calculated to >0.85 in four studies [24-27], 0.75-0.85 in one [23], and <0.75 in four [21, 22, 32, 33]. In five studies values for ΔPOP were as good as, or better than, values for ΔPP [23, 24, 26, 27, 33]. In one of these studies, predictive value of ΔPOP was defined as a certain change in ΔPP, thus presuming that ΔPP is a good indicator [27]. One study found poor values for both ΔPP and ΔPOP [33].

3.2. Correlations between ΔPOP, PVI, and ΔPP. ΔPP is considered to be a good predictor of fluid responsiveness [6]. Thus, other variables should correlate with ΔPP. 11 of the included papers reported correlations between ΔAPP and ΔPP. Six of these papers reported relatively good correlations (r > 0.84) [13, 15, 17, 23, 24, 27]. However, five papers reported relatively poor correlations (r < 0.78) [14, 16, 18, 32, 33]. One of these investigated children preoperatively [32]. Landsverk et al. [16] concluded that there are poor correlations between ΔPOP and ΔPP in ICU patients due to sympathetic oscillations in skin circulation, which lead to larger variation in ΔPOP than in ΔPP during registrations over longer time periods. These findings are supported by Hoiseth et al. [33] who also found larger variation in ΔPOP than in ΔPP during ongoing open major abdominal surgery. Four papers examined correlations between PVI and ΔPP. Three of them found relatively poor correlations (r = 0.72, 0.46 and 0.78) [17, 20], whereas one reported better correlations (r = 0.85) [29]. Three papers investigated correlations between PVI and ΔPOP [17, 26, 32]. One study reported poor correlations (r = 0.39) [32], whereas two studies reported relatively good correlations (r = 0.92) [17, 26]. These data are presented in Table 2.

4. Discussion

Photoplethysmography is applicable on most patient categories and is noninvasive, simple, widely available, and without risk of complications. Several physiological, clinical, and practical factors must be taken into account when evaluating whether or not it is a noninvasive alternative to evaluate fluid responsiveness.

Firstly, there are several physiological prerequisites for using dynamic variables.

Mechanical ventilation provides the stable and predictable variations in intrathoracic pressure required for photoplethysmography to be accurate. A large mechanical tidal volume will influence intrathoracic pressure to a greater extent than a small tidal volume. It is presumed that the influence of tidal volume reaches significance at >8 mL/kg. It is a challenge that the accuracy of photoplethysmography increases with larger tidal volumes, whereas it is clinically desirable to minimize the tidal volume. The accuracy of
Table 1: Papers in which ΔPOP and/or PVI have been evaluated and fluid challenges performed.

| Author, Ref.          | Fluid challenge | CO/Cl/SVI measurement | Responder | ROC | Threshold value | Sens/spec |
|-----------------------|-----------------|------------------------|-----------|-----|-----------------|-----------|
| Solus-Biguenet et al. [21] | 250 mL colloid | Thermodilution          | SVI: 10%  | 0.81 | PPV<sub>fin</sub>: 14% | No data   |
|                       |                 |                        |           | 0.68 | ΔPOP: 9.5%      | No data   |
|                       |                 |                        |           | 0.79 | PPV<sub>art</sub>: 12.5% | No data   |
| Natalini et al. [22]  | 500 mL HES 6%   | Thermodilution          | CI: 15%   | 0.72 | ΔPOP: 15%       | 56/86 (PPV/NPV) |
|                       |                 |                        |           | 0.74 | ΔPP: 15%        | 55/100 (PPV/NPV) |
| Cannesson et al. [23] | 500 mL HES 6%   | Thermodilution          | CI: 15%   | 0.847| ΔPOP: 13%       | 93/90     |
|                       |                 |                        |           | 0.847| ΔPP: 11%        | 80/90     |
| Feissel et al. [24]   | 8 mL/kg HES 6%  | Echo-Doppler            | CI: 15%   | 0.94 | ΔPP: 12%        | 100/70    |
|                       |                 |                        |           | 0.94 | ΔPOP: 14%       | 94/80     |
| Wyffels et al. [25]   | 500 mL HES 6%   | Intermittent thermodilution by pulm. artery catheter | CO: 15%   | 0.94 | PPV: 11.3%      | 95/91.7   |
|                       |                 |                        |           | 0.89 | ΔPOP: 11.3%     | 90/83.3   |
| Cannesson et al. [26] | 500 mL HES 6%   | Thermodilution          | CI: 15%   | 0.94 | ΔPP: 12.5%      | 87/89     |
|                       |                 |                        |           | 0.94 | ΔPOP: 12%       | 87/89     |
| Westphal et al. [27]  | 500–1000 mL NaCl| Not measured            | ΔPP > 13% | 0.95 | ΔPOP: 11%       | 91/100    |
| Zimmermann et al. [28]| 7 mL/kg HES 6%  | FloTrac                 | SVI: 15%  | 0.97 | PVI: 9.5%       | 93/100    |
| Loupec et al. [29]    | 500 mL HES or PLR if ΔPP < 13% | Echocardiography | CO: 15%   | 0.88 | PVI: 17%        | 95/91     |
| Author, Ref. | Fluid challenge | CO/CI/SVI measurement       | Responder | ROC  | Threshold value | Sens/spec |
|-------------|-----------------|----------------------------|-----------|------|-----------------|-----------|
| Desgranges et al. [30] | 500 mL HES | Thermodilution | CI: 15% | 0.91 | PVI<sub>forehead</sub>: 15% | 89/78     |
|              |                 |                           |           | 0.88 | PVI<sub>forehead</sub>: 16% | 74/74     |
|              |                 |                           |           | 0.84 | PVI<sub>finger</sub>: 12% | 74/67     |
|              |                 |                           |           | 0.84 | PPV > 11%       | 74/89     |
|              |                 |                           |           |      | PVI<sub>forehead</sub>: 15% and PI<sub>forehead</sub>: 1.37 | 89/100    |
| Renner et al. [31] | HES 10 mL kg<sup>-1</sup> | Transoesophageal echocardiography | SVI: 15% | 0.79 | PVI > 13%       | 84/64     |
| De Souza Neto et al. [32] | Saline, 20 mL/kg | Transthoracic echography (aortic velocity-time integral) | AVTI: 15% | 0.51 | 0–6 yr: ΔPOP | No data   |
|              |                 |                           |           | 0.63 | 0–6 yr: PVI     | No data   |
|              |                 |                           |           | 0.71 | 0–6 yr: ΔPP    | No data   |
|              |                 |                           |           | 0.52 | 0–6 yr: PPV    | No data   |
|              |                 |                           |           | 0.57 | 6–14 yr: ΔPOP  | No data   |
|              |                 |                           |           | 0.54 | 6–14 yr: PVI   | No data   |
|              |                 |                           |           | 0.60 | 6–14 yr: ΔPP   | No data   |
|              |                 |                           |           | 0.60 | 6–14 yr: PPV   | No data   |
| Hoiseth et al. [33] | 250 mL colloid | Esophageal doppler | SV ≥ 15% | 0.67 | ΔPP: 8.8%       | 82/67     |
|              |                 |                           |           | 0.72 | ΔPOP: 11.4%     | 86/67     |
| Hood and wilson [34] | 500 mL colloid | Esophageal doppler | SV ≥ 10% | 0.96 | PVI<sub>finger</sub> (baseline): 10% | 86/100    |
|              |                 |                           |           | 0.98 | PVI<sub>earlobe</sub> (baseline): 9.5% | 95/100    |
|              |                 |                           |           | 0.71 | PVI<sub>finger</sub> (during surgery): 10% | 65/67     |
|              |                 |                           |           | 0.54 | PVI<sub>earlobe</sub> (during surgery) | No data |

ΔPOP: pulse oximetry plethysmography; ΔPP: pulse pressure; PVI: Pleth variability index; CI: cardiac index; SV: stroke volume; SVI: stroke volume index; SVV: stroke volume variation; CO: cardiac output; PPV<sub>fina</sub>: pulse pressure variation obtained with Finapres; PPV<sub>art</sub>: pulse pressure variation obtained with intraarterial equipment; PPV/PPV: positive predictive value/negative predictive value.
Table 2: Papers in which correlations between ΔPOP, PVI, and ΔPP have been investigated.

| Author, Ref. | Relation | Correlation | Pulse oximeter/monitor |
|--------------|----------|-------------|------------------------|
| Cannesson et al. [13] | ΔPOP-ΔPP | $r = 0.91, P < 0.001$ | M1190A, Philips, Suresnes, France |
| Natalini et al. [14] | ΔPOP-ΔPP | $r = 0.62, P < 0.001$ | Datex-Engstrom CS/3 Critical Care Monitor, Instrumentarium, Helsinki, Finland |
| Cannesson et al. [15] | ΔPOP-ΔPP | $r = 0.89, P < 0.01$ | Oxymax Tyco Healthcare Group LP, Pleasanton, CA, USA |
|                        | ΔPOP-PVI | $r = 0.92, P < 0.05$ | Intellivue MP70, Philips Medical Systems, Suresnes, France |
| Landsverk et al. [16] | ΔPOP-ΔPP | $r = 0.05, P = 0.15$ | Oximax 451N5, Nellcor, Boulder, CO, USA |
| Cannesson et al. [17] | PVI-ΔPP  | $r = 0.72, P < 0.05$ | LNOP Adt, Masimo Corp., Irvine, CA, USA |
|                        | ΔPOP-ΔPP | $r = 0.86, P < 0.05$ | Intellivue MP70, Philips Medical Systems, Suresnes, France |
| Pizov et al. [18]     | ΔPOP-ΔPP | $r = 0.75$ | Datex-Ohmeda AS-3, Datex, Helsinki, Finland |
| Desebbe et al. [19]   | PVI_{VT=6}−PVI_{VT=10} | 9%–12%, $P = 0.001$ | LNOP Adt, Masimo Corp., Irvine, CA, USA |
|                        | PVI_{PEEP}−PVI_{P_{0.01}−PEEP} | (Significant change) | |
| Biais et al. [20]     | PVI-ΔPP  | $r = 0.46, P = 0.001$ | LNOP Adt, Masimo Corp., Irvine, CA, USA |
|                        | PVI_{N_{E=1}−1}−ΔPP | $r = 0.20, P > 0.05$ | Masimo Radical 7 monitor, Masimo SET, Masimo Corp., Irvine, CA, USA |
|                        | PVI_{N_{E=1}−1}−ΔPP | $r = 0.72, P < 0.001$ | |
| Solus-Biguenet et al. [21] | No data | | |
| Natalini et al. [22]  | No data | | Datex-Engstrom CS/3 Critical Care Monitor, Instrumentarium, Helsinki, Finland |
| Cannesson et al. [23] | ΔPOP-ΔPP | $r = 0.90, P < 0.01$ | Oxymax Tyco Healthcare Group LP, Pleasanton, CA, USA |
|                        |          |             | Intellivue MP70, Philips Medical Systems, Suresnes, France |
| Author, Ref. | Relation | Correlation | Pulse oximeter/monitor |
|-------------|----------|-------------|-----------------------|
| Feissel et al. [24] | ΔPOP-ΔPP | \( r = 0.84, P < 0.001 \) | SpO2/Pleth, M3150A technology, Philips Medical Systems, Andover, MA, USA |
| Wyffels et al. [25] | ΔPOP-ΔPP | \( r = 0.90, P < 0.001 \) | Monitor Hewlett Packard M1166A model G65 |
| Cannesson et al. [26] | ΔPOP-PVI | \( r = 0.92, P < 0.01 \) | LNOP Adt, Masimo Corp., Irvine, CA, USA, with Masimo Radical 7, 7.0.3.3 Oxymax, Tyco Healthcare Group LP, Pleasanton, CA, USA |
| Westphal et al. [27] | ΔPOP-ΔPP | \( r = 0.90, P < 0.001 \) | S/5, Datex-Ohmeda, Helsinki, Finland |
| Zimmermann et al. [28] | ΔPOP-ΔPP | \( r = 0.85, P < 0.0001 \) | LNCS, Masimo Corp., Irvine, CA, USA, Masimo Radical-7 monitor, 7.0.3.3 |
| Loupec et al. [29] | PVI_{baseline}-ΔPP_{baseline} | \( r = 0.85, P < 0.0001 \) | LNCS Adtx, Masimo corp., Irvine, CA, USA |
| Desgranges et al. [30] | ΔPOP-PVI | \( r = 0.39, P < 0.05 \) | LNOP Adt, Masimo Corp., Irvine, CA, USA Oximax, Tyco Healthcare Group LP, Pleasanton, CA, USA |
| | ΔPOP-ΔPP | \( r = 0.48, P < 0.001 \) | LNOP TC-I, Masimo Corp., Irvine, CA, USA LNOP TF-I, Masimo Corp., Irvine, CA, USA Masimo Radical 7, Masimo SET, Masimo Corp., version 7.1.1.5 |
| | PVI-PPV | \( r = 0.78, P < 0.001 \) | Masimo Rainbow SET, Masimo Corp., Radical 7, V7.6.2.2 |
| Renner et al. [31] | | | Masimo Rainbow SET, Masimo Corp., Radical 7, V7.6.2.2 |
| De Souza et al. [32] | ΔPOP-PVI | \( r = 0.39, P < 0.05 \) | Oximax, Tyco Healthcare Group LP, Pleasanton, CA, USA |
| | ΔPOP-ΔPP | \( r = 0.48, P < 0.001 \) | LNOP, Masimo Corp., Irvine, CA, USA |
| | PVI-PPV | \( r = 0.78, P < 0.001 \) | Oximax 451N5, Nellcor, Boulder, CO, USA |
| Hoiseth et al. [33] | ΔPOP-ΔPP | \( r = 0.78, P < 0.001 \) | Masimo Rainbow SET, Masimo Corp., Irvine, CA, USA |
| Hood and Wilson [34] | | | Masimo Rainbow SET, Masimo Corp., Irvine, CA, USA |

ΔPOP: pulse oximetry plethysmography; ΔPP: pulse pressure; PVI: Pleth variability index; PEEP: positive end expiratory pressure; NE: norepinephrine; VT: tidal volume.
| Author, Ref. | Year | n  | Patient category | Ventilation | Site of meas. | Reg. period | Vasoact. |
|-------------|------|----|------------------|-------------|--------------|------------|----------|
| Cannesson et al. [13] | 2005 | 22 | ICU              | Mech. vent. 6–10 mL/kg, volume | Finger | 3 respiratory cycl. | Incl. |
| Natalini et al. [14] | 2006 | 49 | OR/ICU           | Mech. vent. 6–9 mL/kg, volume | Finger/toe | 5 respiratory cycl. | No data |
| Cannesson et al. [15] | 2007 | 25 | Preop. CABG/AAA  | Mech. vent. 8–10 mL/kg, volume | Finger | 3 respiratory cycl. | Excl. |
| Cannesson et al. [16] | 2008 | 14 | ICU              | Mech. vent. 8 mL/kg, volume/pressure | Finger | 15 min | Incl. |
| Cannesson et al. [17] | 2008 | 25 | Preop.CABG       | Mech. vent. 8–10 mL/kg | Finger | 3 respiratory cycl. | Excl. |
| Pizov et al. [18] | 2010 | 33 | Preop. surgery   | Mech. vent. 8–10 mL/kg | Finger | 3 min | Incl. |
| Desebbe et al. [19] | 2010 | 21 | Postop. CABG and ICU | Mech. vent. 6–10 mL/kg, volume | Finger | 3 respiratory cycl. | Excl. |
| Biais et al. [20] | 2011 | 67 | ICU              | Mech. vent. 8 mL/kg, volume | Finger | 3 respiratory cycl. | Incl. |
| Solus-Biguenet et al. [21] | 2006 | 8  | During hepatic surgery | Mech. vent. 8–10 mL/kg | Finger | 3 respiratory cycl. | No data |
| Natalini et al. [22] | 2006 | 22 | ICU              | Mech. vent. 6–10 mL/kg, volume | Finger | 2 min | No data |
| Cannesson et al. [23] | 2007 | 25 | Preop. CABG       | Mech. vent. 8–10 mL/kg, volume | Finger | 3 respiratory cycl. | Excl. |
| Feissel et al. [24] | 2007 | 23 | ICU              | Mech. vent. 8 mL/kg, pressure | Finger | No data | Incl. |
| Wyffels et al. [25] | 2007 | 32 | Postop. heart surgery | Mech. vent. 8–10 mL/kg | Finger | 3 respiratory cycl. | No data |
| Cannesson et al. [26] | 2008 | 25 | Preop.CABG       | Mech. vent. 8–10 mL/kg, volume | Finger | 3 respiratory cycl. | Excl. |
| Westphal et al. [27] | 2009 | 43 | Postop. heart surgery | Mech. vent. 8–10 mL/kg, volume | Finger | 1 min | Incl. |
| Zimmermann et al. [28] | 2010 | 20 | Preop. abd. surgery | Mech. vent. 7 mL/kg, volume | Finger | No data | No data |
| Loupec et al. [29] | 2011 | 40 | Several categories | Mech. vent. 8 mL/kg, volume | Finger | No data | Incl. |
| Desgranges et al. [30] | 2011 | 28 | Preop. cardiac surgery | Mech. vent. 8 mL/kg, volume | Finger/earlobe | No data | No data |
| Renner et al. [31] | 2011 | 27 | Infants preop. cardiac surgery | Mech. vent. 10 mL/kg, volume | No data | No data | Excl. |
| Pereira de Souza et al. [32] | 2011 | 30 | Children preop neurosurgery | Mech. vent. 10 mL/kg, volume | Finger | 3 respiratory cycl. | Excl. |
| Hoiseth et al. [33] | 2011 | 25 | During abd. surgery | Mech. vent. 8 mL/kg, volume | Finger | App. 5 min | Incl. |
| Hood and wilson [34] | 2011 | 25 | During colorectal surgery | Mech. vent. 8–10 mL/kg, volume | Finger/earlobe | No data | No data |

ICU: intensive care unit; OR: operating room; CABG: coronary artery bypass grafting; AAA: abdominal aortic aneurysm.
photoplethysmography relies on a continuous beat-to-beat-analysis. Thus, patients need to have stable heart rate. Additionally, decreased RV ejection fraction can lead to false-positive variations in pulse pressure [35]. These requisitions also apply for other dynamic variables [36–39].

Secondly, the complex network of correlations between ΔPOP/ΔPI and other hemodynamic variables varies greatly between different studies. The best correlations are found in studies where short registration periods (3–5 respiratory cycles) have been used and in patients under stable pre- and postoperative conditions. These conditions do not reflect genuine intraoperative instability, the setting where precise guidance of fluid therapy is perhaps most important. The correlations are poorer with longer periods of registration [16], in heterogeneous patient groups in ICUs [16], and during ongoing open abdominal surgery [21, 33]. The best predictive values for ΔPOP and ΔPI were found in papers in which patients were investigated preoperatively [26, 28]. The poorest predictive values (0.51–0.72) were found during ongoing open major abdominal surgery [21, 33], on sedated patients in ICU [22], and on children preoperatively [32]. In one paper, the predictive value of PVI decreased from 0.96 at baseline to 0.71 during surgery [34]. This indicates that photoplethysmography shows best results in standardized conditions, during short registration periods, and in homogenous groups of pre- and postoperative patients. Importantly, it has been demonstrated that PVI reduces both lactate levels and volumes of fluid administered in surgical patients [40]. This is interesting evidence. However, the study does not report improvement in terms of the number of complications. Further studies are needed to clarify the very important aspect of improved outcome.

Finally, a number of additional factors must be considered. Variations in total peripheral resistance and vasomotor tone increase under the influence of general anesthesia [41, 42], with vasoactive drugs, with site of measurement, and with physiological responses such as inflammation, pain, fear, and body temperature. This may lead to inaccuracy of the photoplethysmography signal. The papers included suggest that ΔPOP is less reliable in ICU patients. This may be explained by the above-mentioned factors. Hemodynamics of patients in the OR or in ICUs changes rapidly and continuously. In most papers which good predictive values for photoplethysmography have been found, short registration periods are used. In papers with longer registration periods, poorer predictive values have been reported.

A threshold value refers to a value of ΔPOP, ΔPP, or PVI that separates responders from nonresponders. Failure to agree upon a threshold value in clinical settings does not necessarily make the parameters (i.e., PVI or POP) less valuable. Different patient groups may well present with different threshold values. A septic patient may have a threshold value different from that of a hemodynamically stable patient undergoing surgery. In the same way, threshold values may also change pre-, peri-, and postoperatively. Cannesson et al. [43] discussed the very interesting notion of a gray-zone approach to fluid responsiveness and found that an intermediate zone of pulse pressure variation could not predict fluid responsiveness. Future studies should grade responses instead of dividing responses in two categories.

Cut-off values for increases in SV/CO/CI are defined to separate responders and nonresponders. These thresholds are based on the variability and errors in the chosen measuring technique as well as what change is believed to be clinically important. These thresholds may be more or less arbitrarily chosen and differ between the studies.

Level of intra-abdominal pressure may influence ΔPP and ΔPOP and is relevant in three of the articles included [21, 28, 33]. Results are not coherent. Animal studies have shown that increased intra-abdominal pressure leads to an increase in ΔPP [44]. Studies investigating the influence of these fluctuations during laparoscopic surgery are currently running.

In theory, a number of potentially confounding factors exist. Different pulse oximeter-technology, errors due to software autogain features which filter and amplify the raw signal (thus making it unreliable for quantitative analysis), atherosclerosis, type of fluid, skin pigmentation, saturation, movement artefacts, statistical weaknesses, variations in pleural and transpulmonary pressures, and venous components of the pulsatile signal may affect measurements.

5. Conclusion

We conclude that although photoplethysmography is a promising technique, predictive values and correlations with other hemodynamic variables indicating fluid responsiveness vary substantially. Based on studies using short registration periods photoplethysmography might seem promising for evaluation of volume status. However, in studies using longer registration periods it has been shown that intra- and interindividual variability for ΔPOP is greater than for ΔPP, leading to poor agreement between ΔPOP and ΔPP. Thus, it is not presently evident that photoplethysmography is adequately accurate, valid, and reliable to be included in clinical practice for evaluation of volume status. In future studies it is important to evaluate new hemodynamic methods in clinically relevant settings and to test their reproducibility in clinically relevant time frames. Relatively poor predictive values during ongoing major surgery further underscore this point and results vary in different patient groups. The greatest potential for photoplethysmography in evaluation of volume status might be in settings where invasive monitoring is not indicated.

Conflict of Interests

There is no conflict of interests for any of the authors.

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