Accuracy and Trending Ability of Electrical Biosensing Technology for Non-invasive Cardiac Output Monitoring in Neonates: A Systematic Qualitative Review

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Background: Electrical biosensing technology (EBT) is an umbrella term for non-invasive technology utilizing the body’s fluctuating resistance to electrical current flow to estimate cardiac output. Monitoring cardiac output in neonates may allow for timely recognition of hemodynamic compromise and allow for prompt therapy, thereby mitigating adverse outcomes. For a new technology to be safely used in the clinical environment for therapeutic decisions, it must be proven to be accurate, precise and be able to track temporal changes. The aim of this systematic review was to identify and analyze studies that describe the accuracy, precision, and trending ability of EBT to non-invasively monitor Left ventricular cardiac output and/or stroke volume in neonates.

Methods: A qualitative systematic review was performed. Studies were identified from PubMed NCBI, SCOPUS, and EBSCOHost up to November 2021, where EBT technologies were analyzed in neonates, in comparison to a reference technology. Outcome measures were bias, limits of agreement, percentage error for agreement studies and data from 4-quadrant and polar plots for trending studies. Effect direction plots were used to present results.

Results: Fifteen neonatal studies were identified, 14 for agreement and 1 for trending analysis. Only thoracic electrical biosensing technology (TEBT), with transthoracic echocardiography (TTE) as the comparator, studies were available for analyzes. High heterogeneity existed between studies. An equal number of studies showed over- and underestimation of left ventricular output parameters. All studies showed small bias, wide limits of agreement, with most studies having a percentage error >30%. Sub-analyses for respiratory support mode, cardiac anomalies and type of technology showed similar results. The single trending study showed poor concordance, high angular bias, and poor angular concordance.

Discussion: Overall, TEBT shows reasonable accuracy, poor precision, and non-interchangeability with TTE. However, high heterogeneity hampered proper analysis.
INTRODUCTION

Adequate systemic perfusion is dependent on cardiac output (CO), as determined by heart rate (HR) and stroke volume (SV) and influenced by systemic vascular resistance (SVR). A complex interaction exists between HR, blood flow, SVR and blood pressure (BP) to ensure that cellular metabolic oxygen demand is met (1). The circulatory system of neonates is significantly different from that of adults or children, as the neonatal population is a heterogeneous mix of gestational and postconceptional ages, with different degrees of cardiovascular maturation (2). Indirect measures of CO, such as HR and BP, are inadequate for the assessment of neonatal hemodynamic status (3).

CO is considered a fundamental physiological parameter for diagnosis and guidance of therapy in various neonatal conditions (4). Maintaining optimal perfusion and oxygenation is of prime concern in neonatal intensive care units (NICU). Comprehensive monitoring of various physiological variables is required, as low CO has been associated with increased morbidity, adverse neurodevelopmental outcome, and increased mortality (5).

CO measurement, via invasive techniques (e.g., intermittent pulmonary artery thermodilution and Fick’s method) are considered the gold standards for accurately determining CO (6). However, in neonates these methods are inappropriate (7) as catheters are often too big and the invasiveness of these methods have been questioned, also in adult medicine (8). Minimally invasive cardiac output monitoring technologies encompass devices not requiring the insertion of a pulmonary artery catheter, e.g., pulse contour, pulse power analysis, partial gas re-breathing and transpulmonary ultrasound dilution (9). Some of these technologies require the placement of an arterial line (pulse contour and pulse power analysis) and may need placement of a central venous line for calibration purposes (7). These technologies have been poorly studied in the neonatal population whilst others are still under development (transpulmonary ultrasound dilution) (10). Most other CO measurement technologies in neonates offer only intermittent measurement values as they are labor, skill or technology intensive [transthoracic echocardiography (TTE) and cardiac magnetic resonance imaging (cMRI)] (7).

Non-invasive cardiac output monitoring technologies were therefore developed, offering fully non-invasive methods of monitoring SV and CO. These included intermittent measurements via Doppler ultrasound [Ultrasound Cardiac Output Monitor (USCOM) and transthoracic echocardiography (TTE)] and continuous measurements via various electrical biosensing technologies (EBT) [bioimpedance (BI) and bioreactance (BR)].

For a new technology to be safely used in the clinical environment, and to allow therapeutic decisions to be based upon it, it must be proven to be accurate and precise. A good agreement between a new technology and a gold standard reference technology is defined by a small bias (indicating a high accuracy), narrow limits of agreement (indicating a high precision) and a percentage error ≤ 30% (indicating technology interchangeability) (11, 12). Trending ability (change over time) should also be assessed to ensure that the new technology’s direction and magnitude of change is in line with that of the reference technology (13).

TECHNOLOGY BACKGROUND

The first type of non-invasive cardiac monitoring, rheocardiology, was developed in 1949 by Kedrov (14) but only found popularity in 1966 when Kubicek re-designed it for use in the aerospace industry (15). Since then, numerous iterations of this technology have become available in the healthcare industry, with methodologies measuring changes in whole body, segmental or thoracic impedance from which stroke volume (SV) and hence cardiac output (CO) are derived. Numerous nomenclatures are used – whole body electrical bioimpedance (WBEB), thoracic electrical bioimpedance (TEB), electrical velocimetry, electrical cardiometry, impedance cardiometry, impedance cardiology, thoracocardiology, bioreactance, and rheocardiology. These have subtle differences, often with proprietary algorithms and models to estimate SV and CO.

For EBT a high frequency, low amplitude electrical current is applied across the thorax (TEBT) or entire body (WBEBT). The resistance (impedance, Z0) to this electrical current varies between different tissues in the body, with the primary distribution being to the blood and extracellular fluid. This change in electrical current (∆Z0) over time (dZ0/dt) corresponds to the level of SV, from which CO can be computed.

Electrical biosensing technology is divided into 2 broad categories: (1) bioimpedance (BI) which encompasses thoracic...
Electrical velocimetry/electrical cardiometry, impedance cardiography as well as WBEBT and (2) bioreactance (BR).

**Electrical Velocimetry and Electrical Cardiometry**

Electrical cardiometry (EC) is the method of non-invasive cardiac output technology that utilizes the model of thoracic electrical velocimetry (EV) to determine SV and CO (16). These are used by Aesculon and ICON, manufactured by Osypka Medical GmbH, Germany.

In EV, the change in impedance (ΔZ0) is due to the degree of erythrocyte alignment in the aorta throughout the cardiac cycle. During diastole, as the aortic blood flow ceases, erythrocytes are randomly orientated and interfere with electrical conduction. During systole as the left ventricle contracts, the erythrocytes are forced to align parallel to aortic flow and the electrical current in the aorta passes with less impedance, hence an increased conductivity. These pulsatile changes in volume and thus in impedance, in relation to the cardiac cycle (ΔZ0(t)), are used to calculate SV.

EV estimates SV by means of the following equation:

\[ SV_{TEB} = CP \times v_f \times FT \] (1)

where \( SV_{TEB} \) is SV estimated by thoracic electrical bioimpedance, \( CP \) is the patient constant (in ml), \( v_f \) is the mean blood velocity index (in s\(^{-1}\)) during flow time (FT; measured in s). The EV model estimates SV based on the input of the patient's body mass, an empiric mean velocity index derived from a peak amplitude measurement assumed to be the peak aortic blood flow acceleration and a measurement of flow time.

**Impedance Cardiography**

Impedance cardiography (ICG) and electrical cardiometry (EC) are similar as both rely on periodical volumetric changes in the aorta to determine SV and CO. However, ICG and EC differ in the model applied to determine impedance measurements, specifically as to how the change in impedance is calculated. In ICG the change in impedance (conductivity) (ΔZ(t)) is solely attributed to the volumetric expansion of the ascending aorta due to the increase of volume within the aorta or due to its wall motion. The index of peak velocity of the volumetric change is used in ICG as compared to the index of peak acceleration in EV. EV includes direction of flow whereas ICG does not. In EV, volume changes also incorporate the alignment of erythrocytes.

**Bioreactance**

In Bioreactance (BR), another thoracic EBT method, it is assumed that blood flow changes are not only related to changes in impedance but also changes in capacitance (ability of biological tissue to store an electrical current) and inductance (biological tissue's ability to store energy in a non-electrical form). BR therefore measures phase shift (\( \phi \)) of an oscillating current as it traverses the thorax. Four pairs of sensors, one electrode acting as a high frequency generator and the other as a receiver, are placed on either side of the thorax. CO measurements are determined separately from each side of the body and the final CO is the average of the measurements. BR uses the following formula to estimate SV:

\[ SV = C \times VET \times d\phi/dt_{max} \] (2)

where \( C \) is a constant of proportionality, \( VET \) is ventricular ejection time, and \( d\phi/dt_{max} \) is the peak rate of change of the phase shift (\( \phi \)). BR is used by the Reliant and its newer version, Starling, manufactured by Baxter, United States.

Significant differences exist between BI and BR (Table 1).

The aim of this systematic review was to identify and analyze studies that describe the accuracy, precision, and trending ability of EBT to non-invasively monitor cardiac output and/or stroke volume in newborn infants.

**METHODS**

A systematic search was performed on PubMed NCBI, SCOPUS, and EBSCOHost to identify English language studies published up until 30 November 2021. Search terms included: non-invasive cardiac output, thoracic impedance, bioreactance, whole body bioimpedance, electrical velocimetry, electrical cardiometry, impedance cardiometry, impedance cardiography, cardiac output, stroke volume, neonate, newborn, and infant.

Studies on human neonates, term or preterm of any gestational and postconceptional age, were eligible for inclusion. The search strategy did not explicitly exclude animal studies to ensure that studies reporting animal and human research would be identified. However, studies reporting pure animal data were excluded upon screening of the title and abstract. Studies describing accuracy, precision, agreement, or trending data for non-invasive cardiac output monitors, as compared to a standard reference technique were eligible for inclusion. Thoracic and whole body electrical biosensing technology, encompassing bioimpedance and bioreactance technologies, were eligible for inclusion. Any standard comparative technology [transthoracic echocardiography (TTE), thermodilution techniques, Fick principle derived methods, or cardiac MRI (cMRI)] to compare to the investigational technology were eligible for inclusion. Studies investigating cardiac output (CO), or stroke volume (SV) were eligible. Studies that did not compare an investigational technology to a reference technology were excluded.

Any study reporting outcome measures allowing validation of EBT technologies were included. All published and pre-print manuscripts pending publication were eligible for inclusion.

Studies were selected for inclusion by screening titles and abstracts against selection criteria by 2 independent reviewers (LVW and WPdB) and conflicts were resolved through discussion, arbitrated by a third reviewer (SG), as required. Full text articles of the included studies were reviewed by the same reviewers to confirm eligibility and perform data extraction.

Extracted data included study details (first author, year of publication and study population details), investigational technology device name, comparator technology type and outcome data. If outcome data were not specifically reported, but were calculatable from provided data, the missing outcome
TABLE 1 | Comparison of technological differences between bioimpedance and bioreactance.

|               | Bioimpedance                                                                 | Bioreactance                                                                 |
|---------------|-------------------------------------------------------------------------------|------------------------------------------------------------------------------|
| Assumptions   | Related to change in body's resistance only                                     | Related to change in body's resistance and capacitance                        |
| Flow-related  | Fluid-filled syringe or truncated cone                                          | Throat is an electrical circuit with resistor and capacitor                   |
| Electrode     | Calculated from estimated changes in bioimpedance/impedance (Z0) – electrical | Peak rate of change over time (dZ0/dt)                                          |
| placement     | current flow over time due to aortic blood flow changes                         | VET is determined by the distance between QRS complexes                       |
| SV estimation | Calculated from phase shift, determined from resistance and capacitance (Z0)  | Applied at the base of the neck (thoracic inlet) and ospharyngeal constriction |
| VET determination | VET is determined by the distance between changes in phase shift               |                                                                                   |

Outcome measures were defined as follows:
For agreement (accuracy and precision) (12):
1. a. Bias CO or SV, mean difference (investigative technology – comparator).
2. b. Mean CO or SV ((investigative technology + comparator)/2).
3. c. Percentage error (1.96 × SD of bias/mean × 100%).
4. d. Limits of agreement (bias ± 1.96 × SD).
5. e. True precision (TP) of comparator correcting for the true precision of TTE's precision of ±30%:

\[ TP = \sqrt{(PE)^2 - (0.3)^2} \]

For trending ability (19):
1. Concordance rate: number of sequential changes in CO (ΔCO) or SV (ΔSV) (data points in concordant quadrants/all data points) × 100%.
2. Mean angular bias: average angle between all polar data points and polar axis.
3. Radial limits of agreement (LOA): defined as the radial sector containing 95% of data points (mean angular bias ± 1.96 × SD).
4. Angular concordance rate: calculated as the percentage of data points in the ±30° radial zone.

STRATEGY FOR DATA SYNTHESIS

Data synthesis strategy was defined by the extracted data. Although included studies were quantitative in nature and data were descriptively summarized, data gathered in this review were considered to be too heterogeneous to allow statistical pooling for meta-analysis. There was variable reporting of measured hemodynamic parameters (CO or SV), hemodynamic parameter measurement unit (weight indexed vs. non-weight indexed) as well as data reporting in a wide variety of clinical situations. Studies were thus grouped primarily according to hemodynamic parameter, type of measurement unit, and according to clinical variables and interventions.

Heterogeneity was visually assessed by tabulation of the primary author, year of publication, hemodynamic variable, measurement unit, investigational technology, study population (gestational age, postnatal age, and birth weight) and interventions. Data were analyzed in line with these factors.

Effect direction plots were utilized to visually synthesize the diverse outcome measures, providing a link between the data and the narrative (20). Size and direction of arrows were used to indicate sample size, bias direction, precision, and percentage error (above or below 30% benchmark). Results were presented for overall results as well as for sub-analyses.
This review was performed according to the synthesis without meta-analysis (SWiM) guideline (21).

RESULTS

From an initial search, 295 studies were identified, which were assessed for eligibility. After full-text review, 15 studies were included, of which 14 were agreement studies (22–35) and 1 was a trending analysis study (36) (Figure 1 and Table 2).

All studies were observational, prospective, method comparison studies. No study utilized whole body electrical biosensing technology (WBEBT), therefore only thoracic electrical biosensing technologies (TEBT) were included. The only reference method utilized in the included studies was transthoracic echocardiography (TTE). Investigational technologies comprised bioreactance (BR) (3 studies) and bioimpedance (11 studies) (Table 2). BI included: NCCOM³ Reliant (3 studies) (Table 2).

Most studies (n = 7) reported CO measurements only, 2 studies reported CO and SV, 2 studies reported SV only and 1 study reported TTE-VTI (velocity time integral) as well as TTE-MM (M-mode). Most studies (n = 7) reported weight-indexed measurements, 6 studies reported non-weight indexed measurements, and 1 study reported both (Table 2).

Thirteen studies reporting accuracy and precision outcome measures could be identified with only 1 neonatal study reporting trending data. Descriptive tables were ordered by hemodynamic parameter, weight-indexed or non-weight-indexed measurement and sample size.
TABLE 2 | Study characteristics of included neonatal studies comparing TEBT and a reference technology.

| References | Patient characteristics | Gestational age (weeks) (mean ± SD) | Birth weight (kg) (mean ± SD) | Postnatal age (days) [median (range)] | Investigative method (model used) | Reference method | Sample size [patients (measurements)] | Variable measured (unit) |
|------------|-------------------------|------------------------------------|-----------------------------|--------------------------------------|---------------------------------|-----------------|--------------------------------------|------------------------|
| Tibballs (22) | Post cardiac surgery and RDS 100% IV Inotropes – NS | Prem and term | 0.75–4.95 | 17 (0–121) | Bi (NCCOM3) | TTE | 26 (81) | CO (ml/kg/min) |
| Grollmuss (23) | TGA switch procedure 100% IV Inotropes: NS | NS | 3.3 ± 0.5 | 10 (3–29) | Bi (Aesculon) | TTE | 24 (240) | SV (ml) |
| Noori (24) | 40% PDA | 39.2 ± 1.1 | 3.09 ± 0.33 | 4 (1–13) | Bi (Aesculon) | TTE | 20 (115) | CO (ml/min) |
| Weisz (25) | 10% IV and 20% NIV | 37 ± 6.6 | 2.72 ± 1.23 | NS | BR (Reliant) | TTE | 10 (97) | CO (ml/min) SV (ml) |
| Blohm (26) | 4.9% IV 38.3% NIV 5% PDA | 25–34 | 1.66 | NS | Bi (Aesculon) | TTE | 26 (40) | |
| Grollmuss (27) | 67% IV | 31.7 ± 3.1 | 3.1 ± 1.61 | 15 (1–48) | Bi (ICON) | TTE | 28 (228) | SV (ml/kg) |
| Song (28) | 50.4% IV 49.4% NIV 64% PDA 42.5% inotropes | 27 ± 2.96 | 1.07 ± 0.78 | 0–1.5 | Bi (NS) | TTE | 40 (109) | CO (ml/kg/min) |
| Weisz (29) | PDA ligation 88% IV 100% IV 52% inotropes | 30.6 ± 2.51 | 0.7 ± 0.11 | NS | BR (Reliant) | TTE | 25 (78) | CO (ml/kg/min) |
| Torigoe (30) | 29.6% IV 45.6% NIV 100% PDA | 32 ± 2.9 | 1.63 ± 0.53 | NS | Bi (NS) | TTE | 28 (81) | CO (ml/min) |
| Blohm (31) | 4% PDA 42% PFO 21% PDA and PFO | Preterm and term | 3.3 ± 2.51 | 1.9 (0.16–240) | Bi (Aesculon) | TTE | 99 (291) | CO (ml/min) |
| Boet (32) | 31.6% IV 41.8% NIV 4% PDA | 31 ± 3.2 | 1.11 ± 0.53 | 0–7 | Bi (NS) | TTE | 79 (451) | CO (ml/kg/min) SV (ml) |
| Hsu (33) | 67% IV 35% NIV 100% PDA | 27.2 ± 6.6 | 1.01 ± 1.00 | 6 (2–22) | Bi (Aesculon) | TTE | 36 (905) | CO (ml/kg/min) |
| Van Wyk (34) | 70% NIV | 31.3 ± 2.7 | 1.56 ± 0.41 | 0–3 | BR (Reliant) | TTE | 63 (674) | CO (ml/kg/min) SV (ml/kg) |
| Hassan (35) | 8% HFO 46% SIMV 13% NIMV 25%CPAP 16% PDA | 25.2 (22.3–31.6) | 668 ± 157 | 24 (9–80) | Bi (ICON) | TTE | 38 (685) | CO (ml/kg) |

Agreement studies (accuracy and precision)

Trending analysis studies

Bi, bi impedance; BR, bi reactivity; CO, cardiac output; ICON, index of contractility monitor; IV, invasive ventilation; NCCOM3, non-invasive computerized cardiac output monitor; NIV, non-invasive ventilation; NS, not specified; PDA, patent ductus arteriosus; PFO, patent foramen ovale; SD, standard deviation; TTE, trans thoracic echocardiography.
The 14 eligible agreement studies involved a total number of 504 patients, encompassing 2668 paired measurements. Study sample sizes were generally small [average number of patients per study 38.7 (range 10–99)] with only 3 studies recruiting more than 50 patients. The average number of paired measurements per patient was 5.4 (range 1.5–11.9) with only 5 studies performing more than 5 paired measurements per patient (Table 2).

Studies varied widely in gestational ages, birth weight and chronological ages of enrolled neonates. Disease states, ventilation requirements and surgical intervention also varied widely between studies (Table 2), thereby illustrating the high heterogeneity between studies.

**Agreement Analysis**
To determine the overall agreement (i.e., accuracy and precision) of TEBT vs. TTE, all studies were included in the analysis (Table 3). Most studies (8 out of 14) were large (>100) in sample size (number of measurements). An equal number of studies (7 of 14) showed that mean bias was positive or negative, indicating an overestimation and underestimation, respectively, of TTE measurements. The effect direction plots show a visual representation of sample sizes as well as the overestimation and underestimation of measurements (Table 4).

Non-weight indexed SV mean bias was reported in 4 studies and varied between 0.6 to 1.1 ml. Limits of agreement (LOA) varied between ±0.75 and ±2.35 ml. Percentage error (PE), reported in only 3 studies, varied between 29 and 58% (Tables 3, 4).

CO mean bias measurement was reported in 9 studies and ranged between 8.9 to −18.5 ml/kg/min for weight indexed and 6 to −153 ml/min for non-weight-indexed CO. LOA varied between ±13.5–132.7 ml/kg/min and ±66.5–233 ml/min, respectively. PE ranged between 5.3 to 71.6%, with only 5 of the 14 included studies reporting a PE ≤30% (Tables 3, 4).

One study used VTI as hemodynamic reference parameter. The bias percentage was 39% and PE was 46%.

In order to determine whether there were differences in accuracy and precision when studies were performed with different ventilation modes, the 5 studies reporting outcome measures for various respiratory support methods were analyzed (Table 5). Five studies reported outcome measures for non-invasively ventilated (NIV) infants and 4 studies for invasively ventilated (IV) infants.

In the NIV group, most studies reported outcomes for CPAP whereas 1 study reported non-invasive intermittent ventilation outcome measures. Sample sizes were small (8–54 measurements) with one large study (>100 measurements). TEBT underestimated TTE in 4 out of 5 studies. Weight-indexed CO mean bias varied between −2.8 and −23.0 ml/kg/min with wide LOA (±40.7 to 124.9 ml/kg/min). Non-weight-indexed CO mean bias varied between 3.6 and −178 ml/min with LOA between ±70.1 to ±94.6 ml/kg/min and ±LOAs available. PE for NIV studies ranged between 17.3 to 78.5% with only 2 out of 5 NIV studies having a PE ≤30% (Table 5).

For the IV group, outcome measures were reported in 4 studies utilizing various intermittent mandatory ventilation (IMV) modes [synchronized IMV, assist control ventilation, 4 studies utilizing high frequency ventilation (HFV) and 1 study utilizing high frequency jet ventilation (HFJV)]. In all studies, sample sizes were small (<50 measurements). TEBT underestimated TTE in most studies (8 out of 9). For IMV modes, the weight-indexed CO mean bias was −1.4 to −30.2 ml/kg/min with wide LOA (±70.1 to ±94.6 ml/kg/min) and non-weight-indexed CO mean bias of −29.6 to −115 ml/min with only a single study providing LOA (±97.8 ml/min). Three studies provided PE, ranging from 27.4 to 69.7%, with 3 studies providing PE with only 1 of the 3 of studies reporting a PE ≤30%. For HFV studies, CO mean bias was −16.2 to 38.2 ml/kg/min and −12 ml/min for weight-indexed and non-weight indexed measurements, respectively. LOA ranged between ±79 to ±179.1 ml/kg/min and only 1 non-weight-indexed study proving a LOA of ±105.9/ml. For HFV studies, PE ranged between 33.2–85.5% with no studies reporting a PE ≤30%. The single HFJV study reported a CO mean bias of −10.9 ml/kg/min and PE of 76.4% (Table 5).

In a cross comparison of all respiratory modes, mean bias, LOA and PE seemed to show an increase as the complexity of respiratory support increased (Table 6).

The effect direction plots (Table 4) show a visual summary of the sample sizes, bias and PE for all respiratory support studies. Numerous other studies included patients on CPAP but did not specifically report outcome measures for that subset of data (23, 25, 26, 32).

For NIV, Van Wyk et al. (34) showed a significant difference in weight-indexed CO bias between CPAP and no respiratory support (78.0 vs. 74.4%, p = 0.026) but not for weight-indexed SV bias (p = 0.113). Blohm et al. (26) showed a significant effect of CPAP on the bias between TEBT and TTE-VTI (p = 0.022) but not TEBT and TTE-MM (m-mode).

Song et al. (28) showed no significant difference in PE between CPAP and SIMV (57 vs. 69.7%, p = 0.160) nor SIMV and HFV (69.7 and 85.5%, p = 0.729). Hsu et al. (33) showed minimal change in bias between CPAP and IMV but a very large increase in bias between IMV and HFV, with incremental increases in PE from CPAP to IMV to HFV. Song et al. (28) reported a worsening of weight-indexed CO bias between neonates on CPAP and SIMV (−18.2 ml/kg/min and −30.2 ml/kg/min, respectively). Increased complexity of respiratory support intervention caused an increase in PE. Hassan et al. (35) showed that HFV only had a significantly lower bias (p = 0.002) as compared to IMV and CPAP.

Other studies reported no effect of respiratory support mode on CO bias. Torigoe et al. (30) showed no effect of mechanical ventilation on bias (estimated mean bias of 60 ml/min for no respiratory support and SIMV and approximately 25 ml/min for nCPAP and HFV, p = 0.14). Grollmuss et al. (23) reported that method interchangeability was not affected by respiratory support mode although no data was provided.

To determine whether there were differences in accuracy and precision when cardiac lesions were present, studies reporting outcome measures for different congenital cardiac lesions (pathological and physiological) were analyzed. Six studies were included. Most studies reported small sample sizes with only
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**TABLE 3 | Overall data regarding accuracy and precision of TEBT and reference technology.**

| Authors/Year | Unit of measurement | Number Patients (measurements) | Mean* | Bias** Mean ± SD | LOA (precision) | Overall PE |
|--------------|---------------------|--------------------------------|-------|------------------|-----------------|------------|
| Stroke Volume |                     |                                |       |                  |                 |            |
| Grollmuss (23) | ml                  | 24 (240)                       | 3.7   | 0.28             | ±2.3            | 29         |
| Weisz (29)    | ml                  | 25 (78)                        | 1.25  | −0.6 (39%)       | ±0.75           | 58.0       |
| Blohm (31)    | ml                  | 99 (291)                       | 5.2   | 0.7              | ±2.35           | 44.9       |
| Boet (32)     | ml                  | 79 (451)                       | NS    | 1.1              | ±1.85           | NS         |

**Cardiac Output |**

| Authors/Year | Unit of measurement | Number Patients (measurements) | Mean* | Bias** Mean ± SD | LOA (precision) | Overall PE |
|--------------|---------------------|--------------------------------|-------|------------------|-----------------|------------|
| Tibballs (22) | ml/kg/min            | 26 (78)                        | 239   | 0.23 ± 6.5       | ±13.50          | 5.3        |
| Grollmuss (27) | ml/kg/min            | 28 (228)                       | 256.4 | 8.9 ± 3.19       | ±82.7           | 24         |
| Song (28)     | ml/kg/min            | 40 (109)                       | 209.5 | −18.8 ± 67.7     | ±132.7          | 60.2       |
| Hsu (33)      | ml/kg/min            | 38 (105)                       | 258   | −5.3 ± 37.2      | ±72.9           | 28.2       |
| Van Wyk (34)  | ml/kg/min            | 63 (754)                       | 124.4 | −18.5            | ±87.6           | 71.6       |
| Noori (24)    | ml/min               | 20 (115)                       | 536   | −4               | ±233            | 43.6       |
| Weisz (25)    | ml/min               | 10 (97)                        | 417   | −153 ± 56        | ±152.5          | 48.3       |
| Torgoe (30)   | ml/min               | 28 (81)                        | 314   | 6 ± 46.9         | ±66.5           | 21         |
| Hassan (35)   | ml/min               | 38 (85)                        | 271   | −126             | ±178.5          | 66         |
| VTI           |                     |                                |       |                  |                 |            |
| Blohm (26)    | m                   | 26 (41)                        | NS    | 39%              | ±66.5           | 46.2       |

*Mean = (mean TEBT + mean TTE)/2; **Bias = TEBT – TTE; LOA, limits of agreement; NS, not stated; PE, percentage error; SD, standard deviation; TEBT, thoracic electrical biosensing technology; TTE, transthoracic echocardiography; VT, velocity time integral. Bold values have been calculated from available data or determined from provided from graphs.

1 study consisting of more than 100 measurements (Table 5). Most studies (6 out of 8) showed that TEBT underestimated TTE (Table 7).

Two studies reported outcome measures in cardiac surgical interventions. The TGA switch study (23) showed a non-weight-indexed SV mean bias of 0.27 ml and a PE of 29%. The PDA ligation study (29) reported a non-weight-indexed SV mean bias of −0.6 ml and a PE of 58% (Table 7).

In studies (n = 4) reporting outcome measures for patients with a PDA, SV and CO mean bias were reported for both weight- and non-weight-indexed measurements. PE in the 4 studies varied between 21 to 74.4%. Hemodynamic significant PDA was defined in only 2 studies (30, 35) (Table 7).

In studies (n = 2) reporting outcome measures for patients with a PFO, only 1 study reported data for PFO only and one for PDA combined with PFO. Both studies showed a high PE (40.2 and 56.3%, respectively) (Table 7).

The effect direction plots (Table 4) shows a visual summary of the sample sizes, bias and PE for all studies reporting data for cardiac lesions.

Other studies included patients with a PDA but did not specifically report outcome measures for that subset of data (28, 32). Various studies reported associations between PDA and mean bias without reporting specific outcome measures. Noori et al. (24) reported no statistically significant difference (p = 0.800) in bias (12 vs. 2 ml/min) or precision (±296 vs. ±218 ml/min) between neonates with a hemodynamically significant (dual diameter > 2 mm) PDA as compared to those without. Van Wyk et al. (34) reported a significantly higher mean bias for infants with an open PDA compared to a closed PDA (−28.7 ml/kg/min vs. −12.5 ml/kg/min, p < 0.001). Blohm et al. (26) showed that PDA status showed a trend toward significance when TEBT was compared to TTE measured VTI mean bias (p = 0.077) but not when TEBT was compared to TTE m-mode measurements. Hassan et al. (35) showed a near doubling in mean CO bias between PDA and non-PDA measurements (−167 ml/min and −89 ml/min, respectively, p < 0.001).

To determine if accuracy and precision was related to the type of TEBT technology utilized, studies were analyzed according to whether bi impedance (BI) (n = 11) or bi reactorance (BR) (n = 3) was used (Table 8).

In studies utilizing BI, 5 out of the 11 studies showed overestimation and 6 out of 11 showed underestimation of TTE hemodynamic parameters. SV mean bias varied between 0.28 and 1.1 ml with LOA between ±1.85 and ±2.35 ml. CO mean bias varied between −18.8 and 0.23 ml/kg/min and −4 to 6 ml/min with LOA between ±13.5 to ±132.7 ml/kg/min and ±66.5 to ±233 ml/min for weight-indexed and non-weight-indexed CO bias, respectively. PE varied between 5.3–46.2%, with 5 out of 10 studies meeting the PE < 30% benchmark (1 study did not provide PE data) (Table 8).

In studies utilizing bi reactorance, all studies showed an underestimation of TTE hemodynamic parameters (Table 8). PE ranged between 48.3–71.6% with all studies exceeding the PE < 30% benchmark.

**Trending Analysis**

The single neonatal study assessing trending parameters utilized bi reactorance (36). The study showed a poor concordance rate (77.2%), high mean angular bias (28.6°) with wide angular LOA and poor angular concordance (17.4%). Exclusion zone size affected trending parameters. PDA and CPAP showed no association with trending parameters, but CO level was significantly associated with trending parameters.
### TABLE 4 | Effect direction plots for TEBT studies: overall accuracy and sub-analyses of clinical conditions.

| Study          | Overall | CPAP | IMV | HFO/HJV | Cardiac surgical intervention | Physiological shunts | Bioimpedance | Bioreactance |
|----------------|---------|------|-----|---------|-------------------------------|----------------------|--------------|--------------|
|                | Sample  | Mean Bias | PE | Sample  | Bias | PE | Sample  | Bias | PE | Sample  | Bias | PE | Sample  | Bias | PE |
| Tibbals 1989   | ▲ ▼ ▼   | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ |
| Grollmuss 2012 | ▲ ▼ ▼   | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ |
| Noon 2012      | ▲ ▼ ▼   | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ |
| Weisz 2012     | ▲ ▼ ▼   | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ |
| Blohm 2014     | ▲ ▼ ▼   | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ |
| Weisz 2014     | ▲ ▼ ▼   | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ |
| Grollmuss 2014 | ▲ ▼ ▼   | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ |
| Song 2014      | ▲ ▼ ▼   | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ |
| Torigoe 2015   | ▲ ▼ ▼   | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ |
| Blohm 2016     | ▲ ▼ ▼   | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ |
| Boet 2016      | ▲ ▼ ▼   | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ |
| Hsu 2017       | ▲ ▼ ▼   | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ |
| Van Wyk 2020   | ▲ ▼ ▼   | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ |
| Hassan 2020    | ▲ ▼ ▼   | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ | ▲ ▼ ▼ |

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*HFO: **HFV: *PDA only; **PFO only; **PDA < 1.5 mm; **PDA ≥ 1.5 mm; CPAP, continuous positive airway pressure; HFO, high frequency oscillation; HFV, high frequency jet ventilation; IMV, intermittent mandatory ventilation; NS, not specified; PDA, patent ductus arteriosus; PFO, patent foramen ovale; PE, percentage error; VTI, velocity time integral. Legend: Sample size (based on number of measurements: large arrow – large sample size > 100, medium arrow 50–100, and small arrow <50). Bias effect: Downward green arrow – TEBT underestimates reference method, upward red arrow – TEBT overestimates reference method; size of arrow indicates degree. Percentage error: downward green arrow PE < 30%, upward red medium arrow 30–50%, upward large red arrow > 50.
### TABLE 5 | Accuracy and precision in studies with different respiratory support modes.

| Authors          | Unit of measurement | n Measurements | Type of respiratory support | Mean | Bias** Mean ± SD | LOA (precision) | PE |
|------------------|---------------------|----------------|----------------------------|------|------------------|-----------------|----|
| **Non-invasive ventilation** |                     |                |                             |      |                  |                 |    |
| Song (28)        | ml/kg/min           | 54             | CPAP                       | 209  | -18.2 ± 63.7     | ±124.9          | 57 |
| Hsu (33)         | ml/kg/min           | 37             | CPAP                       | 258  | -2.8 ± 20.8      | ±40.7           | 17.3 |
| Van Wyk (34)     | ml/kg/min           | 335            | CPAP                       | 124.4| -23.0 ± 45.8     | ±84.8           | 78.5 |
| Torigoe (30)     | ml/min              | 37             | HFOV                       | 314  | 3.6              | ±73.1           | 25.0 |
| Hassan (35)      | ml/min              | 21             | CPAP                       | 268  | -178             | NS              | NS |
|                  |                     |                | NIMV                       | 291  | -138             | NS              | NS |
| **Invasive ventilation** |                     |                |                             |      |                  |                 |    |
| Song (28)        | ml/kg/min           | 39             | SIMV                       | 209  | -30.2 ± 73.8     | ±94.6           | 69.7 |
| Hsu (33)         | ml/min              | 10             | SIMV                       | 314  | -29.6            | ±97.8           | 31.7 |
| Song (28)        | ml/kg/min           | 8              | HFOV                       | 209.5| 38.2 ± 91.4      | ±179.1          | 85.5 |
| Hsu (33)         | ml/kg/min           | 24             | HFOV                       | 258  | -16.2 ± 40.4     | ±79             | 37.8 |
| Hassan (35)      | ml/min              | 7              | HFOV                       | 177  | -21.5            | NS              | NS |

*Mean = (mean TEBT + mean TTE)/2; **Bias = TEBT – TTE. CPAP: continuous positive airway pressure; HFOV: high frequency oscillatory ventilation; HFJV: high frequency jet ventilation; ACV: assist control ventilation; LOA, limits of agreement; NS, not stated; PE, percentage error; SD, standard deviation; SIMV: synchronized intermittent mandatory ventilation; TEBT, thoracic electrical biosensing technology; TTE, transthoracic echocardiography. Bold values have been calculated from available data or determined from provided graphs.

No other studies performed formal trending analysis. However, 2 studies showed changing agreement parameters over time. The TGA switch study (23) also showed varying bias and PE dependent on timing of measurements after surgery: 24, 35, and 28% within 0–36, 36–72 and after 72 h of surgery, respectively. The PDA ligation study (29) also showed an increase in bias over time: 7.9% (6–8 h post-ligation) and 9.7% (16–18 h post-ligation) as compared to measurements 1-h post-ligation.

**DISCUSSION**

In this review of TEBT technology in neonates, 14 studies were found describing the agreement between bioimpedance (n = 11) and bioreactance (n = 3) technology and transthoracic echocardiography (TTE). A total of 542 neonates were enrolled in these studies, comprising 2755 paired measurements. Only 1 study was found describing trending analysis, comparing bioreactance to TTE, which comprised 63 neonates and 691 paired measurements.

**Thoracic Electrical Biosensing Technology Accuracy and Precision**

In the current review, the mean bias (difference between TEBT technology and the TTE reference) was small in many studies but limits of agreement were wide, indicating acceptable accuracy but a lack of precision. This was similar to 2 adult TEBT systematic reviews, where bias was determined to be small [−0.22 l/min (37) and 0.03 l/min (38)] with wide LOA [−2.43; 1.99 l/min (37) and −2.78; 2.84 l/min (38)]. A pediatric systematic review showed similar small bias [−0.02 to −0.01 l/min (38, 39) and 3.4 ml/kg/min (39) and wide LOA −1.22; 1.18 l/min (38) and ±75.17 ml/kg/min (39). However, in the most recent pediatric systematic review (39), only 5 neonatal studies were included.

Most studies providing PE data (8 out of 13) in this review did not meet the clinically accepted percentage error benchmark of less than 30%, thereby indicating that TEBT is not interchangeable with the reference technology, TTE. PE represents the LOA adjusted for the mean of both methods and therefore represents the random error between the two methods. It depicts the intrinsic variations in the assessed hemodynamic parameter that are not linked to true changes of that parameter (CO or SV), but rather to the environment and random precision error of the investigated or reference technology (38). The commonly accepted 30% arises from the original cardiac output method comparison studies using thermodilution as the reference technology, which has an inherent precision of 20% or less. Thus, if a new technology has a similar precision to thermodilution (i.e., ±20%), the combination will lead to a total error of ±28.3%, which is commonly rounded off to 30%. Therefore, if a new technology has a percentage error of <30%, the technology has a similar percentage error to the reference technology and is therefore an acceptable alternative (11). However, this PE only holds true when the reference technology has the same PE as thermodilution, i.e., 20%. The inherent percentage error for other technologies is often higher. TTE has been shown to have a much higher PE (39–53%) than thermodilution and cannot be considered as accurate as thermodilution. For this reason, it has been suggested that the PE threshold should be increased to 45% to conclude agreement between the two methods, to compensate for the variability of the reference method (40). If this argument were to be followed in this review, it would only increase the
| Authors | Unit of measurement | n | Mean bias | Precision | PE |
|---------|---------------------|---|-----------|-----------|----|
| Song (28) | ml/kg/min | 54 | 18.2 | ±34.6 | 69.7 |
| Hsu (33) | ml/kg/min | 37 | 2.8 | ±4.0 | 17.3 |
| Torigoe (30) | ml/min | 37 | 3.6 | ±3.1 | 25.0 |
| Van Wyk (34) | ml/kg/min | 235 | 2.3 | ±4.5 | 18.0 |
| Hassan (35) | ml/min | 21 | 2.5 | ±1.5 | 30.0 |

**Bias = TEBT – TTE.** CPAP, continuous positive airway pressure; SIMV, synchronized intermittent mandatory ventilation; HFOV, high frequency oscillatory ventilation; HFJV, high frequency jet ventilation; LOA, limits of agreement; PE, percentage error; TEBT, thoracic electrical biosensing technology; TTE, transthoracic echocardiography.

The current review showed significant heterogeneity amongst studies regarding measured hemodynamic parameter, unit of measurement, gestational and postnatal age of included neonates as well as different management strategies (respiratory support modes, inotropic support) as well as presence of physiological shunts or congenital cardiac disease. This is similar to the adult and pediatric systematic reviews showing high heterogenic indices [79.2–93% (37–39)].

Sub-analyses for respiratory support mode, presence of physiological shunts or cardiac disease/cardiac surgery as well as type of EBT technology, showed persistence of small bias, wide LOA, and high PE. Type of EBT technology assessment also showed variable effect of bias directionality: BI studies showed variable under- and overestimation whilst BR showed consistent underestimation of hemodynamic parameters. These differences may be due to sample sizes and population heterogeneity of the included studies, or may be due to the varying underlying mathematical models of the different technologies.

Most studies were performed in preterm infants, confirming the interest by clinicians to determine CO in this vulnerable population.

Percentage error differed between studies depending on respiratory support mode. In general, mean bias and PE increased with complexity and intensity of respiratory support mode, with invasive ventilation having higher PE than non-invasive ventilation methods. Most studies reported PE exceeding the benchmark, suggesting that respiratory support modes made TEBT less interchangeable with TTE. In most studies, the degree of respiratory support was not stated, i.e., mean airway pressure, which may affect bias, due to the amount of air between the sensors and aorta interfering with the sensors ability to measure cardiac outflow.

In studies reporting accuracy and precision for cardiac anomalies, accuracy was reasonable, but precision was poor. PE exceeded the benchmark in most studies. However, different definitions were used to define PDA (size and flow velocity in the pulmonary artery) and the size of PFO was not defined. Two studies were performed in neonates undergoing cardiac surgical intervention, increasing the heterogenicity of results.

Heteroscedasticity implies proportionality of bias, or variability of changes with the magnitude of measurement (57), which is a problem with healthcare method comparison studies (40). In TEBT method comparison studies, this implies increasing bias (difference between TEBT and reference method) with increasing CO or SV. Three studies reported heteroscedasticity in CO or SV bias. Boet et al. (32) showed that TEBT overestimated TTE when SV > 2 ml and CO > 0.4 l/min. Van Wyk et al. (34) showed an increasing bias when CO ≥ 150 ml/kg/min as compared to <150 ml/kg/min (p < 0.001). Hsu et al. (33) showed a statistically significant increase in bias (p = 0.001) but no statistically significant difference in PE when comparing a CO ≥ 280 ml/kg/min as compared to a CO < 280 ml/kg/min. This may be clinically relevant in larger neonates or neonates with high
TABLE 7 | Accuracy and precision in studies with cardiac shunts.

| Authors/Year | Unit of measurement | n measurements | Cardiac defect | Mean* | Bias ** Mean ± SD | LOA (precision) | PE |
|--------------|---------------------|----------------|---------------|-------|-------------------|-----------------|----|
| **Cardiac surgical intervention** | | | | | | | |
| Grollmuss (23) | ml | 240 | TGA switch surgery | 3.7 | 0.27 ±1.06 | ±0.75 | 29.0 |
| Weisz (29) | ml | 78 | PDA ligation | 1.25 | −0.6 (39%) | ±0.75 | 58.0 |
| **Physiological shunt** | | | | | | | |
| Blohm (31) | ml | 12 | PDA only | 4.1 | −0.8 ± 1.73 | ±0.98 | 72.1 |
| Weisz (29) | ml | 63 | PDA + PFO | −1.1 ± 1.58 | ±1.09 | 56.3 |
| | | 125 | PFO only | −0.6 ± 1.11 | ±0.59 | 40.2 |
| Van Wyk (34) | ml/kg/min | 304 | PDA | 124.4 | −28.7 ± 43.7 | ±78.1 | 74.4 |
| Torigoe (30) | ml/min | 23 | PDA ≥ 1.5 mm | 317 | 5.5 | ±66.2 | 21.0 |
| | | 58 | PDA < 1.5 mm | −36.1 | ±119.5 | 38.6 |
| Hassan (35) | ml/min | 14 | DFLPA > 30 cm/s | 295.5 | −167 | 203.8 | 69 |

*Mean = (TEBT + TTE)/2. **Bias = TEBT – TTE. Bold data indicates calculated data or data estimated from provided graphs. DFLPA, end diastolic flow in left pulmonary artery; LOA, limits of agreement; PDA, patent ductus arteriosus; PE, percentage error; PFO, patent foramen ovale; SD, standard deviation; TGA, transposition of great arteries.

TABLE 8 | Outcome measure for studies using bioimpedance or bioreactance technology.

| Authors year | Unit of measurement | n Patients (measurements) | Specific technology | Mean* | Bias ** Mean ± SD | LOA (precision) | Overall PE |
|--------------|---------------------|---------------------------|---------------------|-------|-------------------|-----------------|------------|
| **Bioimpedance** | | | | | | | |
| Grollmuss (23) | ml | 24 (240) | Aesculon | 3.7 | 0.28 ± 0.05 | ±2.3 | 29 |
| Blohm (31) | ml | 99 (291) | Aesculon | 5.2 | 0.7 | ±2.35 | 44.9 |
| Boet (32) | ml | 79 (451) | NS | NS | 1.1 | ±1.85 | NS |
| Hsu (33) | ml/kg/min | 36 (105) | Aesculon | 258 | −5.3 ± 37.2 | ±72.9 | 28.2 |
| Tibballs (22) | ml/kg/min | 267 (78) | Aesculon | 239 | 0.23 ± 6.5 | ±13.50 | 5.3 |
| Grollmuss (27) | ml/kg/min | 68 (228) | ICON | 256.4 | 8.9 ± 31.9 | ±62.7 | 24 |
| Song (23) | ml/kg/min | 80 (109) | NS | 209.5 | −18.8 ± 67.7 | ±132.7 | 60.2 |
| Noori (24) | ml/min | 20 (115) | Aesculon | 536 | −4 | ±233 | 43.6 |
| Torigoe (30) | ml/min | 28 (81) | NS | 314 | 6 ± 46.9 | ±66.5 | 21 |
| Blohm (26) | ml² | 26 (41) | Aesculon | NS | 39% | NS | 46.2 |
| Hassan (35) | ml/min | 38 (85) | ICON | 271 | −126 | ±178.5 | 66 |
| **Bioreactance** | | | | | | | |
| Weisz (29) | ml | 25 (78) | Reliant | 1.25 | −0.6 ± 0.37 | (39%) | ±0.75 | 58.0 |
| Van Wyk (34) | ml/kg/min | 63 (754) | Reliant | 124.4 | −18.5 | ±87.6 | 71.6 |
| Weisz (25) | ml/min | 10 (97) | Reliant | 417 | −153 ± 56 | ±152.5 | 48.3 |

*TTE VTI was hemodynamic parameter measured by TTE. *Mean = (TEBT + TTE)/2. **Bias = TEBT – TTE. Bold data indicates calculated data or data estimated from provided graphs. LOA, limits of agreement; NS, not specified; PDA, patent ductus arteriosus; PE, percentage error; PFO, patent foramen ovale; SD, standard deviation; TGA, transposition of great arteries.

output cardiac failure with high CO or SV and requires further research.

The mean bias represents the systematic error between measurements, i.e., the mean constant difference. Although bias and LOA are used to statistically define accuracy and precision, there is no consensus regarding acceptable clinical cut-off values for these factors. What, therefore, represents an acceptable bias in neonatal studies requires further research.

Thoracic Electrical Biosensing Technology Trending Ability

In trending analysis, the benchmark is a concordance rate of >92%, an angular bias <5°, angular limits of agreement ±30° and an angular concordance of >90% (19). The single trending study in this review did not meet any of these criteria. Two other agreement studies with measurements over time (23, 29), showed a temporal increase in bias and PE. TEBT is therefore not able to accurately track CO, as compared to TTE.

In trending studies, exclusion zones are required. In the single published neonatal study (36), the choice of exclusion zones significantly influenced trending data results. It is unknown what the exclusion zones should be in neonatology and requires further research.

In method comparison studies, the aim is to determine whether a new technology’s accuracy, precision and trending ability is similar to that of a gold reference technology and can therefore be used interchangeably. However, even the most
accurate technologies (thermodilution) are known to have a degree of error. However, the reference technology in all these studies was TTE, which is known to be a relatively inaccurate reference technology (11, 42). Therefore, it cannot be considered an accurate reference method. However, the lack of other reference technologies attests to the difficulties of invasive testing for CO or SV in sick and small neonates and the difficulties in performing cMRI studies. In many of the included method comparison studies, a variety of TTE methods were used to measure SV or CO – aorta measurements at the annulus (24, 32, 33, 35), valve hinge points (25, 26, 30, 34), and Darmon’s triangular technique (23, 27) or were not specified (28, 29). This inconsistency in standardization of TTE measurements may also contribute to the range of agreement and trending data found in this review.

There also remains a question as to the what the impedance signal (Z0) reflects. The classic assumption that the impedance signal (Z0) reflects. The classic assumption that the impedance signal (Z0) originates from the contractile changes of the aorta and/or the alignment of red blood cells during systole and diastole has been questioned (42). Mathematical models have not been able to confirm the origin of the Z0 signal and numerous recommendations have been made to improve these models for future research (43). The impact of these assumptions on the integrated measurement of left ventricular output, right ventricular output, in the presence of PDA and patent foramen ovale in neonates also needs to be considered carefully.

Various United States health insurers have stated that bioimpedance “continues to be reasonable and necessary” in various adult cardiac disease processes (44), despite an earlier finding by the National Institute for Health Research of inadequate evidence to support its use (45). Various concerns regarding the accuracy of these non-invasive cardiac output monitors have been raised in adult and pediatric medicine (37, 38, 46). Despite this, these monitors have been used in neonatology in research and clinical environments for diverse scenarios: monitoring transition at birth (47, 48), cardiac adaptation after birth (49), patent ductus arteriosus (PDA) diagnosis (50), PDA ligation (29), PDA medical therapy (51), monitoring congenital heart disease (52, 53), managing neonatal hemodynamic shock (54) and sepsis (55), and to predict clinical outcomes (56).

Several technological and physiological aspects have to be met prior to routine use of non-invasive cardiac output monitors in the clinical environment: (1) validation against gold reference standards, (2) accuracy along the entire spectrum of gestational age and birth weight, (3) ability to provide continuous measurements in absolute numbers, (3) be reliable, practical and non-invasive, (4) easy to apply, (5) inexpensive for widespread use, (6) feasible, (7) useful in neonates with extra- and intra-cardiac shunts as well as congenital cardiac disease and (8) continuously recordable alongside other physiological monitors (1). Although many of the technical usability aspects (continuous measurements, ease of use, non-invasive, easy to apply, recordable alongside other physiological monitors) have been proven in various studies, this review suggests more research is required regarding the accuracy in different gestational ages, cardiac shunts and congenital heart disease and the technology should be validated against a true reference method.

CONCLUSION

Thoracic electrical biosensing technology, irrespective of the type of technology, has a poor interchangeability with TTE in newborn infants for left ventricular cardiac output/stroke volume. High heterogeneity of patients and interventions in the neonatal population made direct comparisons of studies difficult. TTE, as a comparator in this review, however, cannot be considered an ideal reference method and studies evaluating TEBT against an accurate reference method are required. TEBT should be used with caution in the neonatal population for monitoring and determining therapeutic interventions.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/subplementary material, further inquiries can be directed to the corresponding authors.

AUTHOR CONTRIBUTIONS

LV, SG, and W-PB contributed to conception and design of the study, performed literature search, and extracted data for analysis. LV wrote the first draft of the article. All authors contributed to manuscript revision, read and approved the submitted version.

REFERENCES

1. Soleymani S, Borzage M, Seri I, Seri I, Borzage M, Seri I, et al. Hemodynamic monitoring in neonates: advances and challenges. J Perinatol. (2010) 30:538-45. doi: 10.1038/jp.2010.101
2. Vrancken SL, van Heijst AF, de Boode WP. Neonatal hemodynamics: from developmental physiology to comprehensive monitoring. Front Pediatr. (2018) 6:87. doi: 10.3389/fped.2018.00087
3. Azhibekov T, Noori S, Soleymani S, Seri I. Transitional cardiovascular physiology and comprehensive hemodynamic monitoring in the neonate: relevance to research and clinical care. Semin Fetal Neonatal Med. (2014) 19:45-53. doi: 10.1016/j.siny.2013.09.009
4. Marik PE. Noninvasive cardiac output monitors: a state-of-the-art review. J Cardiothorac Vasc Anesth. (2013) 27:121-34. doi: 10.1053/j.jvca.2012.03.022
5. Beausoleil TR, Janaiyic M, Barrington KJ, Lapointe A, Dehaes M. Cerebral oxygen saturation and peripheral perfusion in the extremely premature infant with intraventricular and/or pulmonary haemorrhage early in life. Sci Rep. (2018) 8:41598. doi: 10.1038/s41598-018-24836-8
6. Lee AJ, Cohn JH, Ranasinghe JS. Cardiac output assessed by invasive and minimally invasive techniques. Anesthesiol Res Pract. (2011) 2011:475151. doi: 10.1155/2011/475151
7. McGovern M, Miletin J. Cardiac output monitoring in preterm infants. Front Pediatr. (2018) 6:84. doi: 10.3389/fped.2018.00084
8. Hadian M, Pinsky MR. Evidence-based review of the use of the pulmonary artery catheter: impact data and complications. Crit Care. (2006) 10:58. doi: 10.1186/cc4834
18. Wan X, Wang W, Liu J, Tong T. Estimating the sample mean and standard deviation from the sample size, median, range and/or interquartile range. BMC Med Res Methodol. (2014) 14:135. doi:10.1186/1471-2288-14-135

19. Critchley LA, Yang XX, Lee A. Assessment of trending ability of cardiac output monitors by polar plot methodology. J Cardiothorac Vasc Anesth. (2011) 25:536–46. doi:10.1053/j.vca.2011.01.003

20. Thomson HJ, Thomas S. The effect direction plot: visual display of non-standardised effects across multiple outcome domains. Res Synth Methods. (2013) 4:95–101. doi:10.1002/jrsm.1060

21. Campbell M, McKenzie JE, Sowden A, Li T, Sowden AJ, Li T, et al. Systematic reviews without meta-analysis (SWiM) in systematic reviews: reporting guideline. BMJ. (2020) 368:l6890. doi:10.1136/bmj.l6890

22. Tibbals J. A comparative study of cardiac output data vs. neonates supported by mechanical ventilation: measurement with thoracic electrical bioimpedance and pulsed Doppler ultrasound. J Pediatr. (1989) 114:632–5. doi:10.1016/S0022-3476(89)80710-3

23. Grollmus O, Demontoux S, Capderou A, Serraf A, Belli E. Electrical velocimetry as a tool for measuring cardiac output in small infants after heart surgery. Intensive Care Med. (2012) 38:1032–9. doi:10.1007/s00134-012-2530-3

24. Noori S, Drabu B, Soleymani S, Seri I, Drabu B, et al. Continuous non-invasive cardiac output measurements in the neonate by electrical cardiometry: a comparison with echocardiography. Arch Dis Child Fetal Neonatal Ed. (2012) 97:F340–3. doi:10.1136/fgnfae2011-301090

25. Weisz DE, Jain A, McNamara PJ, El-Khuflash A. Non-invasive cardiac output monitoring in neonates using bioimpedance: a comparison with echocardiography. Neonatology. (2012) 102:61–7. doi:10.1159/000337295

26. Blohm ME, Obrecht D, Hartwich J, Mueller GC, Kersten JF, Weil J, et al. Impedance cardiography (electrical velocimetry) and thoracostethoracic echocardiography for non-invasive cardiac output monitoring in pediatric intensive care patients: a prospective single-center observational study. Crit Care. (2014) 18:603. doi:10.1186/s13054-014-0603-0

27. Grollmus O, Gonzalez P. Non-invasive cardiac output measurement in low and very low birth weight: a method comparison. Front Pediatr. (2014) 2:16. doi:10.3389/fped.2014.00016

28. Song R, Rich W, Kim JH, Finer NN, Katheria AC. The use of electrical cardiometry for continuous cardiac output monitoring in preterm neonates: a validation study. Am J Perinatol. (2014) 31:1105–10. doi:10.1055/s-0034-1371707

29. Weisz DE, Jain A, Ting J, McNamara PJ, El-Khuflash A, Weisz DE, et al. Non-invasive cardiac output monitoring in preterm infants undergoing patent ductus arteriosus ligation: a comparison with echocardiography. Neonatology. (2014) 106:330–6. doi:10.1159/000365278

30. Torigoe T, Sato S, Nagayama Y, Sato T, Yamazaki H. Influence of patent ductus arteriosus and ventilators on electrical velocimetry for measuring cardiac output in very-low/low birth weight infants. J Perinatol. (2015) 35:485–9. doi:10.1111/jep.12345

31. Blohm ME, Hartwich J, Obrecht D, Kersten JF, Singer D, Blohm ME, et al. Effect of patent ductus arteriosus and patent foramen ovale on left ventricular stroke volume measurement by electrical velocimetry in comparison to transthoracic echocardiography in neonates. J Clin Monit Comput. (2017) 31:589–98. doi:10.1007/s10877-016-9878-9

32. Boet A, Jourdian D, Demontoux S, De Luca D, Boet A, Jourdian D, et al. Stroke volume and cardiac output evaluation by electrical cardiometry: accuracy and reference nomograms in hemodynamically stable preterm neonates. J Perinatol. (2016) 36:748–52. doi:10.1038/jp.2016.65

33. Hsu K-H, Wu T-W, Wu I-H, Lai M-Y, Hsu S-Y, Huang H-W, et al. Electrical cardiometry to monitor cardiac output in preterm infants with patent ductus arteriosus: a comparison with echocardiography. Neonatology. (2011) 112:231–7. doi:10.1159/000475774

34. Van Wyk L, Smith J, Lawson J, De Boode WP. Agreement of cardiac output measurements between non-invasive and transthoracic echocardiography in preterm infants during the transitional phase: a single-centre, prospective study. Neonatology. (2020) 117:271–8. doi:10.1055/s-0036-1506203

35. Hassan MA, Bryant MB, Hummeler HD. Comparison of cardiac output measurement by electrical velocimetry with echocardiography in extremely low birth weight neonates. Neonatology. (2021) 119:18–85. doi:10.1055/s-0036-159713

36. Van Wyk L, Smith J, Lawson J, Lombard CJ, De Boode WP. Bioreactance cardiac output trending ability in preterm infants: a single centre, longitudinal study. Neonatology. (2021) 118:600–8. doi:10.1055/s-0036-158656

37. Joosten A, Desebbe O, Suehiro K, Murphy LS, Essiet M, Alexander B, et al. Accuracy and precision of non-invasive cardiac output monitoring devices in perioperative medicine: a systematic review and meta-analysis. Br J Anaesth. (2017) 118:298–310. doi:10.1093/bja/aew461

38. Sanders M, Servaas S, Slagt G. Accuracy and precision of non-invasive cardiac output monitoring by electrical cardiometry: a systematic review and meta-analysis. J Clin Monit Comput. (2020) 34:433–60. doi:10.1007/s10877-019-00330-y

39. Mansfield RC, Kaza N, Charalambous A, Milne A, Sathiaymurthy S, Banerjee J. Cardiac output measurement in neonates and children using noninvasive electrical bioimpedance compared with standard methods: a systematic review and meta-analysis. Crit Care Med. (2022) 50:126–37. doi:10.1097/CCM.0000000000005144

40. Peyton PJ. Continuous minimally invasive peri-operative monitoring of a cardiac output by pulmonary capnometry: comparison with thermistor and transesophageal echocardiography. J Clin Monit Comput. (2012) 26:1211–32. doi:10.1007/s10877-012-9342-4

41. Wetersens M, Moller-Sorensen H, Johansen RR, Perner A. Systematic review of cardiac output measurements by echocardiography vs. thermistion: the techniques are not interchangeable. Intensive Care Med. (2016) 42:1223–33. doi:10.1007/s00134-016-4528-y

42. de Sitter A, Verdaasdonk RM, Faes TJ. Do mathematic model studies settle the controversy on the origin of cardiac synchronous trans-thoracic electrical impedance variations? A systematic review. Physiol Meas. (2016) 37:R86–108. doi:10.1088/0967-3334/37/9/R88

43. Anand G, Yu Y, Lowe A, Kalra A. Bioimpedance analysis as a tool for hemodynamic monitoring: overview, methods and challenges. Physiol Meas. (2021) 42:03TR01. doi:10.1088/1361-6579/abe80e

44. Medicare Coverage Database. Decision Memo for Electrical Biopotential for Cardiac Output Monitoring (CAG-00001R). (2020). Available online at: https://www.cms.gov/medicare-coverage-database/details/nca-decision-memo.aspx?NCAID=23#fomdb=true (accessed April 5, 2021).

45. National Institute for Health Research. Thoracic Electrical Biopotentials. Cent Rev Dissec. (2001). Available online at: https://www.crd.york.ac.uk/CRDWeb/ShowRecord.asp?AccessionNumber=3200000165&UserID=0 (accessed April 5, 2021).

46. Cornette J, Laker S, Jeffery B, Lombaard H, Alberts A, Rizopoulos D, et al. Validation of maternal cardiac output assessed by transthoracic
Van Wyk et al. Systematic Review Electrical Biosensing Technology

Echocardiography against pulmonary artery catheterization in severely ill pregnant women: prospective comparative study and systematic review. *Ultrasound Obstet Gynecol.* (2017) 49:25–31. doi: 10.1002/uog.16015

47. McCarthy KN, Pavel A, Garvey AA, Hawke A-LL, Levins C, Livingstone V, et al. Feasibility of non-invasive cardiac output monitoring at birth using electrical bioimpedance in term infants. *Arch Dis Child Fetal Neonatal Ed.* (2020) 106:431–4. doi: 10.1136/archdischild-2019-318244

48. Freidl T, Baik N, Pichler G, Schwabeger B, Zingerle B, Avian A, et al. Haemodynamic transition after birth: a new tool for non-invasive cardiac output monitoring. *Neonatology.* (2016) 111:55–60. doi: 10.1159/000446468

49. Coskun S, Yüksel H, Bilgi Y, Lacin S, Tansug N, Onag A. Non-invasive evaluation of the adaptations of cardiac function in the neonatal period: a comparison of healthy infants delivered by vaginal route and caesarean section. *Acta Med Okayama.* (2001) 55:213–8. doi: 10.18926/AMO/31993

50. Rodríguez-Sánchez de la Blanca A, Sánchez Luna M, González-Pacheco N, Arriaga Redondo M, Navarro Patiño N. Electrical velocimetry for non-invasive monitoring of the closure of the ductus arteriosus in preterm infants. *Eur J Pediatr.* (2018) 177:229–35. doi: 10.1007/s00431-017-3063-0

51. Hsu KH, Wu TW, Wu IH, Lai MY, Hsu SY, Huang HW, et al. Baseline cardiac output and its alterations during ibuprofen treatment for patent ductus arteriosus in preterm infants. *BMC Pediatr.* (2019) 19:179. doi: 10.1186/s12887-019-1560-1

52. Norozi K, Beck C, Osthau WAA, Wille I, Wessel A, Bertram H. Electrical velocimetry for measuring cardiac output in children with congenital heart disease. *Br J Anaesth.* (2008) 100:88–94. doi: 10.1093/bja/aem320

53. Gatelli IF, Vitelli O, Chiesa G, De Rienzo F, Martinielli S. Noninvasive cardiac output monitoring in newborn with hypoplastic left heart syndrome. *Am J Perinatol.* (2020) 37:S54–5. doi: 10.1055/s-0040-1713603

54. Gatelli IF, Vitelli O, Fossati M, De Rienzo F, Chiesa G, Martinielli S. Neonatal septic shock and hemodynamic monitoring in preterm neonates in a nicu: added value of electrical cardiac output in real-time tailoring of management and therapeutic strategies. *Am J Perinatol.* (2021), doi: 10.1055/s-0041-1726123

55. Miletin J, Semberova J, Martin AMAM, Janota J, Stranak Z. Low cardiac output measured by bioimpedance and adverse outcome in preterm infants with birth weight less than 1250 g. *Early Hum Dev.* (2020) 149:105153. doi: 10.1016/j.earlhumdev.2020.105153

56. Elrefaee A, Eldin ZA, Mostafa FA, Elmazzahy, Bekheet S. Electrical cardiometry and functional echocardiography predicting and hemodynamic monitoring of neonatal sepsis. * Turk J Physiol Rehabil.* (2021) 32:12 040–9.

57. Nawarathna LS, Choudhary PK. A heteroscedastic measurement error model for method comparison data with replicate measurements. *Stat Med.* (2015) 34:1242–58. doi: 10.1002/sim.6424

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The remaining authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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