**INTRODUCTION**

In order to identify and treat unnatural circulatory changes in critically ill patients, it is important to know the limits of normality. Indexing of haemodynamic variables has therefore been used to establish comparable body size-independent reference values valid for the whole population.¹ In the growing child, haemodynamic reference values are constantly changing making their estimation challenging.² ⁴ Indexing is also complicated by the fact that children have a relatively larger body surface area (BSA) compared with body weight (BW) than adults, resulting in a non-linear-indexed haemodynamic variables.⁵ Empirical formulas

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**Background:** Haemodynamic studies in children are rare and most studies have included few subjects in the youngest age group. Haemodynamic variables need to be indexed to establish a reference of normality that is valid in all populations. The traditional way to index haemodynamic variables with body surface area (BSA) is complicated in young children due to its non-linear relationship with body weight (BW). We examined several haemodynamic variables in children by indexing them with BSA and BW.

**Methods:** A single-centre, observational cohort study comparing non-indexed and indexed haemodynamic variables in children undergoing heart surgery (divided into three weight groups: 1-5 kg, >5-10 kg and >10-15 kg).

**Results:** A total of 68 children were included in this study, mean age 11.1 months ± 11.1 month (range 0 to 43 months). All haemodynamic variables, cardiac output (CO), stroke volume (SV), total end-diastolic volume (TEDV), central blood volume (CBV) and active circulation volume (ACV), increased with weight without indexing (P < .05). Indexing variables with BW produced a more linear relationship for all haemodynamic variables between weight groups than BSA. The mean BSA-indexed haemodynamic values were CI$_{BSA}$ 3.5 ± 1.1 L/min/m² and SVI$_{BSA}$ 27.3 ± 8.9 ml/min/m². The mean BW-indexed haemodynamic values were CI$_{BW}$ 180 ± 50 ml/min/kg and SVI$_{BW}$ 1.34 ± 0.38 ml/kg. Blood volume variables indexed with BW were TEDV$_{BW}$ 12.0 ± 2.8 ml/kg, CBV$_{BW}$ 21.3 ± 6.6 ml/kg and ACV$_{BW}$ 70.3 ± 15.2 ml/kg.

**Conclusions:** Indexing haemodynamic variables with BW produces a more appropriate body size-independent scale in young children than BSA.

**Summary statement:** In this study, we studied indexing of haemodynamic variables and estimation of blood volumes in young children undergoing corrective heart surgery using an indicator dilution technology.
used to determine BSA in children often disagree in their calculations, adding to possible errors regarding estimations of indexed haemodynamic variables.\textsuperscript{6}

In 1963, Jegier and colleagues published their landmark paper using dye dilution method, on the relationship between cardiac output (CO) and body size, comparing indexing with BSA and BW.\textsuperscript{7} They found that although both size variables had a similar correlation with CO, there was a significant difference between age groups when BSA was used for CO indexing. However, their conclusion was that this difference in indexing with BSA and BW was not significant enough to justify a change in the already established tradition of using BSA for indexing of haemodynamic variables. These findings, almost 60 years ago, strongly suggested that indexing CO with BSA might be suboptimal due to a non-linear relationship with BW in young children.

Our hypothesis in the present study was that BW would be a more appropriate indexing alternative than BSA, to establish body size-independent reference values for haemodynamic variables in young children. The primary aim was to examine several haemodynamic variables indexed with BSA and BW. A secondary aim was to define indexed reference blood volume values and investigate the effect of intracardiac shunts on their estimations.

2 | MATERIALS AND METHODS

We performed a single-centre, retrospective observational cohort study in young children with congenital heart defects undergoing corrective heart surgery of atrial septal and/or ventricle septal defects. The inclusion criteria for this study were written informed parental consent, elective open-heart surgery and weight less than 15 kilograms. All children included in this study were enrolled from two departments,\textsuperscript{8,9} at Lund Children’s Hospital from February 2010 to March 2017. These studies were approved and registered by the Ethics Committee of Lund University, Sweden (Dnr 2009/708 and Dnr 2013/636).

2.1 | Experimental protocol

Anaesthesia was performed with a bolus of fentanyl (5 µg/kg) and thiopental (5 mg/kg). Pancuronium (0.2 mg/kg) was used for muscle relaxation and to facilitate endotracheal intubation. Isoflurane (0.5-1.0%) was used for maintaining anaesthesia for the duration of the operation. A Dräger Apollo anaesthesia machine (Drägerwerk AG & Co, Lübeck, Germany) maintained ventilation with tidal volumes of 7-8 ml/kg and FiO2 <0.40. All children received a central venous catheter (Multicath triple lumen 6 cm, 4.5 F, Vygon Ltd, Swindon, UK) into the internal jugular vein and a peripheral arterial catheter in the radial artery (Neoflon 24 G < 5 kg patient and Venflon 22 G > 5 kg patient, BD Ltd, Wokingham, UK). The COstatus monitor and the arteriovenous loop were then connected to the arterial and central venous catheters in preparation for measurements. Up to five repeated consecutive haemodynamic measurements were performed before and after the surgical correction.\textsuperscript{8,9} Transoesophageal echocardiography was performed after correction to exclude residual shunts.

2.2 | Determination of cardiac output

The determination of CO was obtained by the novel COstatus monitor device (Transonic Systems Inc, New York, USA). It uses an extracorporeal arteriovenous circuit (AV loop) connected to the arterial and central venous catheters where blood flow is regulated by a roller pump. External ultrasound sensors are attached to the venous and arterial sides of the AV loop. The ultrasound sensor on the venous side determines the amount of blood dilution after a venous injection of a normothermic saline bolus (0.5-1.0 ml/kg), and the ultrasound sensor on the arterial side determines the level of dilution after transcardiopulmonary passage of the blood. CO is determined by analysing the transcardiopulmonary blood dilution curve based on Stewart-Hamilton’s indicator dilution principle.\textsuperscript{10-12} CO and derived variables, such as stroke volume (SV), are displayed on the monitor. The technology used by COstatus has been validated and shown to be accurate, precise and safe in several paediatric studies.\textsuperscript{9,13-16}

2.3 | Determination of blood volume

The use of an AV loop with constant blood flow enables measurements of other haemodynamic variables, such as total end-diastolic volume (TEDV), central blood volume (CBV) and active circulation volume (ACV), from the dilution curve.

Total end-diastolic volume (TEDV) is defined as blood volume in the four chambers of the heart at the end of diastole. Its calculation is based on the assumption that the increase in width of the arterial dilution curve, compared to the venous dilution curve, is due to a delay in the propagation of the indicator when it travels through a volume expanding capacitor, such as the heart chambers.\textsuperscript{15} The width of the dilution curves is determined at half their height and is denoted as chords (CH). CH\textsubscript{art} and CH\textsubscript{ven} are the length of the chords of the arterial and venous curves in time units.

\begin{equation}
TEDV = (CO \times (1.62/HR + 0.77 \times CH\textsubscript{ven})).
\end{equation}

HR: Heart rate (beats per minute).
The mean BSA-indexed haemodynamic values were CI_{BSA} 3.5 ± 1.1 L/min/m² and SVI_{BSA} 27.3 ± 8.9 ml/min/m². The mean BW-indexed haemodynamic values were CI_{BW} 180 ± 50 ml/min/kg and SVI_{BW} 1.34 ± 0.38 ml/kg. CO without indexing increased with weight groups. CO indexed for BSA (CI_{BSA}) tended to increase with weight groups but was not statistically significant (P = .091). CO indexed to weight (CI_{BW}) did not change between the weight groups. A linear regression analysis of CO indexing for BSA (CI_{BSA}) and weight (CI_{BW}) showed a significant deviation of CI_{BSA} (P = .003), but CI_{BW} was stable in all the children's weights included in the cohort (Figure 1). BSA/BW ratio was exponentially increased in our study cohort compared expected BSA/BW ratio in older individuals confirming a non-linear relationship between BSA and BW in different age groups (Figure 2).

3.1 Indexing of haemodynamic variables

Blood flow (CO) and blood volume (SV, TEDV, CBV and ACV) variables indexed for BSA and BW are presented in Table 1.

The mean BSA-indexed haemodynamic values were CI_{BSA} 3.5 ± 1.1 L/min/m² and SVI_{BSA} 27.3 ± 8.9 ml/min/m². The mean BW-indexed haemodynamic values were CI_{BW} 180 ± 50 ml/min/kg and SVI_{BW} 1.34 ± 0.38 ml/kg. CO without indexing increased with weight groups. CO indexed for BSA (CI_{BSA}) tended to increase with weight groups but was not statistically significant (P = .091). CO indexed to weight (CI_{BW}) did not change between the weight groups. A linear regression analysis of CO indexing for BSA (CI_{BSA}) and weight (CI_{BW}) showed a significant deviation of CI_{BSA} (P = .003), but CI_{BW} was stable in all the children's weights included in the cohort (Figure 1). BSA/BW ratio was exponentially increased in our study cohort compared expected BSA/BW ratio in older individuals confirming a non-linear relationship between BSA and BW in different age groups (Figure 2).

SV increased with weight groups and after indexing for BSA (SVI_{BSA}), but not when it was indexed for weight (SVI_{BW}).

TEDV increased with weight groups and after indexing for BSA (TEDVI_{BSA}), but it did not when indexed for weight (TEDVI_{BW}).

CBV increased with weight groups and after indexing for BSA (CBVI_{BSA}), but not when it was indexed for BW (CBVI_{BW}).

ACV increased significantly with weight. After indexing for BSA (ACVI_{BSA}), it tended to increase, but this was not statistically significant. ACV indexing with BW (ACVI_{BW}) did not change between weight groups.
TABLE 1  Haemodynamic variables and blood volume by separate weight groups (N = 68)

| Haemodynamic variables | Weight groups (N = number of subjects) | Statistical significance |
|------------------------|-----------------------------------------|-------------------------|
|                        | 1−5 kg (N = 22)                         | >5−10 kg (N = 32) |
| Mean (CO) ±SD (l/min)  | 0.77 ± 0.28±****                       | 1.24 ± 0.53±***       |
| Mean (CO/BSA) ±SD (l/min/m²) | 3.18 ± 1.04                            | 3.48 ± 1.07         |
| Mean (CO/BW) ±SD (ml/min/kg) | 190 ± 50                                | 180 ± 50            |
| Mean (SV) ±SD (ml)    | 5.7 ± 2.0±****                         | 9.8 ± 4.7±***       |
| Mean (SV/BSA) ±SD (ml/m²) | 23.0 ± 6.6±***                         | 27.4 ± 8.9**        |
| Mean (SV/BW) ±SD (ml/kg) | 1.4 ± 0.4                               | 1.4 ± 0.4           |
| Mean (TEDV) ±SD (ml)  | 52.4 ± 17.6±****                       | 76.7 ± 24.9±****   |
| Mean (TEDV/BSA) ±SD (ml/m²) | 214.0 ± 61.5                            | 221.2 ± 45.5±***   |
| Mean (TEDV/BW) ±SD (ml/kg) | 12.7 ± 3.6                              | 11.4 ± 1.9         |
| Mean (CBV) ±SD (ml)   | 82.8 ± 42.2±****                       | 132.7 ± 67.0±****  |
| Mean (CBV/BSA) ±SD (ml/m²) | 355.8 ± 150.0±***                      | 372.1 ± 134.0±***  |
| Mean (CBV/BW) ±SD (ml/kg) | 19.8 ± 8.5                              | 19.3 ± 5.5         |
| Mean (ACV) ±SD (ml)   | 312.0 ± 93.6±****                      | 475.2 ± 138.1±***  |
| Mean (ACV/BSA) ±SD (ml/m²) | 1269.9 ± 311.3                         | 1366.1 ± 282.0     |
| Mean (ACV/BW) ±SD (ml/kg) | 75.5 ± 19.5                            | 69.8 ± 13.3        |

Abbreviations:: CO, cardiac output; SD, standard deviation; BSA, body surface area; BW, body weight; SV, stroke volume; TEDV, total end-diastolic volume; CBV, central blood volume; ACV, active circulation volume.

| Effect of intracardiac shunt on precision |
|------------------------------------------|

The variability of the measurements decreased and precision improved for CIBW, SVIBW, CBVIBW and ACVIBW after the correction of the intracardiac shunt (Table 3). Repeated haemodynamic measurements were available from 44 children before and after correction.

4 | DISCUSSION

In young children, BW was a more appropriate indexing alternative than BSA to eliminate the influence of body size on haemodynamic variables. This indexing enables the comparison of values between children of different sizes and makes it possible to define normal haemodynamic reference values.

In general, regarding all species, physiological metabolic variables, such as oxygen consumption and alveolar ventilation, seem to follow allometric principles in an exponential manner. Same exponential relationship has been believed to apply to CO as a physiological flow variable. At the same time, it has been noticed that volume variables tend to follow a more linear relationship with BW. Several efforts have been made to achieve haemodynamic reference values in young children in the past, but the cohorts are often small and based on the accepted routine to index the values according to BSA.19-21 These studies have shown that the use of BSA as an index, or normalization, is still unsupported by data in infants and that the exponential relationship between BSA and some haemodynamic variables in smaller children needs to be explained.22-23 This is supported by the exponential increase in the BSA/weight ratios that occurs in smaller children (Figure 2).

We found an extraordinarily good agreement in the indexing of CO to kilogram BW (CIBW) in our weight groups of young children (Table 1). However, it is obvious that there is an upper cut-off weight where the indexing to BW overestimates CO (it is unrealistic to believe a normal individual weighing 100 kg to have a CO of 18 l/min). It has been shown that the Du Bois equation, which was used in this study, deviates significantly from the BSA versus BW curve in weights below 15 kg.5 This may indicate that there is an optimal weight level and that CO is better indexed by weight below this level. This must be defined, but according to our results, it is probably near our study maximal weight of around 15 kg.
Intracardiac blood volumes, such as SV and TEDV, are routinely indexed for BSA, analogous to CO. In our analysis, both SV and TEDV increased after indexing for BSA, but, when indexing with BW (SVI_{BW} and TEDVI_{BW}), they were remarkably stable across all weight groups. The same non-linear relationship between intracardiac blood volume and BSA was confirmed in a magnetic resonance
study by Valsangiacomo Buechel et al, where normal ventricular volumes in children were quantified. Their results indicated that intracardiac blood volumes were better correlated with BW than with BSA. TEDV was analysed by CO status and has been found to predict values within the expected physiological range. Our data indicate a slightly higher TEDVI BW in our cohort of children, which can be expected due to a volume overloaded heart caused by the intracardiac shunting of blood. Our findings of appropriate TEDV indexing by BW in children strengthen the possibility of establishing reference values and potential clinical use, which also applies to SVI BW. The circulatory blood volumes (TEDV, CBV and ACV) are potentially important and may be used to detect hypo- or hypervolaemia when normal reference values have been established.

The precision of the measurements improved in several of the haemodynamic variables (CI BW, SVI BW and CBVI BW) after the correction of the intracardiac shunt (Table 3). This may have been caused by a more pronounced ventilator-dependent circulatory variability with an existing shunt. Alternatively, this may also have been due to the difficulty of the program algorithm in exactly defining the right limits of the dilution curve, which was delayed in the presence of a shunt.

Our results raise a concern regarding the traditional way haemodynamic variables are indexed. All CO monitors on the market index CO with BSA for estimation of CI which may produce artificially lower reference values in children compared with adults. This means that derived variables, such as pulmonary vascular resistance index (PVRi BSA) and systemic vascular resistance index (SVRI BSA), might be overestimated in children. Inaccurate haemodynamic estimations may lead to decisions to treat children with unnecessary vasoactive medication and/or fluids. Anaesthesiologists should be aware of this problem when performing haemodynamic studies and measurements in young children.

Limitations of the study were that all our patients had congenital heart disease and all measurements were taken during anaesthesia, before and immediately after corrective cardiac surgery. Their haemodynamic values do not represent normal reference values in healthy children when they are awake. However, most haemodynamic research, to date, is based on children who have some underlying illnesses, as it would not be ethical to inflict invasive measurements on healthy children. There are similar ethical limitations and clinical challenges regarding the estimation of blood volume in children. It is highly doubtful that a comparative blood volume study using dyes, carbon monoxide or radioactive markers (eg iodinated albumin) as was used in the past, would be allowed in healthy children today. Several animal studies aimed to validate the ability of the technique to accurately detect different blood volumes are available. In this study, the focus was on the youngest and smallest children, who are often

| Variable | Mean ± SD |
|----------|-----------|
| CI BW    | 180 ± 50 ml/min/kg |
| SVI BW   | 1.34 ± 0.38 ml/kg |
| CBVI BW  | 21.3 ± 6.6 ml/kg |
| ACVI BW  | 70.3 ± 15.2 ml/kg |
| TEDVI BW | 12.0 ± 2.8 ml/kg |

Note: All parameters indexed to kilogram body weight.
Abbreviations: ACVI BW, active circulation volume index; CBVI BW, central blood volume index; TEDVI BW, total end-diastolic volume index; SD, standard deviation, CI BW, cardiac index; SVI BW, stroke volume index; 95% CI, 95% confidence interval.
under-represented in larger haemodynamic studies that therefore give an inaccurate overall picture.

In our opinion, the number of patients presented in this study was sufficient to give an overall view of haemodynamic values in these weight groups after heart surgery. All measurements were performed by the same researchers in a very homogenous group of patients. In general, minimal inotropic and vasopressor support was needed and there was no difference between weight groups. As all measurements were part of earlier comparison studies, good agreement and precision were confirmed, with a percentage error below 30%.

**TABLE 3** Variability of variables before and after correction of the intracardiac shunt (N = 44)

| Variable      | Before surgical correction | After surgical correction | P value |
|---------------|----------------------------|---------------------------|---------|
| CIBW          | 8.1 (3.5)                  | <.001                     |
| SVI BW        | 8.2 (3.8)                  | <.001                     |
| ACVI BW       | 8.2 (6.6)                  | .14                       |
| CBVI BW       | 5.6 (3.4)                  | .004                      |
| TEDVI BW      | 3.4                        |                           |

**Abbreviation:** CE, coefficient of error (CE = CV/√n) (CV, coefficient of variation, n, number repeated measurements). CIBW, cardiac index; SVI BW, stroke volume index; ACVI BW, active circulation volume index; CBVI BW, central blood volume index; TEDVI BW, total end-diastolic volume index.

TEDVI is not estimated in the presence of an intracardiac shunt.

Note: All variables indexed to kilogram body weight.

**FIGURE 3** Effects of intracardiac shunt on haemodynamic variables (N = 68). There were no significant changes in CIBW (a), SVI BW (b), CBVI BW (c) and ACVI BW (d) before and after surgical closure of an intracardiac shunt (values are means ± 95% confidence interval).

**CONCLUSION**

Indexing haemodynamic variables with BW produces a more appropriate body size-independent scale in young children than BSA. Surgical correction of an intracardiac shunt did not affect the haemodynamic variables significantly but improved the precision of the measurement values.
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CONFLICT OF INTEREST
TSS declares no competing interest. LL declares no competing interest.

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REFERENCES
1. Egan JR, Festa M, Cole AD, et al. Clinical assessment of cardiac performance in infants and children following cardiac surgery. Intensive Care Med. 2005;31:568-573.
2. Tibby SM, Hatherill M, Marsh MJ, et al. Clinicians’ abilities to estimate cardiac index in ventilated children and infants. Arch Dis Child. 1997;77:516-518.
3. Trieu CT, Williams TM, Cannesson M, et al. Babies and children at last: Pediatric cardiac output monitoring in the twenty-first century. Anesthesiology. 2019;130:671-673.
4. Cattermole GN, Leung PYM, Ho GYL, et al. The normal ranges of cardiovascular parameters measured using the ultrasonic cardiac output monitor. Physiol Rep. 2017;5:1-9.
5. Livingston EH, Lee S. Body surface area prediction in normal-weight and obese patients. Am J Physiol Endocrinol Metab. 2001;281:586-591.
6. Sigurdsson TS, Lindberg L. Six commonly used empirical body surface area formulas disagreed in young children undergoing corrective heart surgery. Acta Paediatr. 2020;109:1838-1846.
7. Jegier W, Sekeli P, Auld PAM, et al. The relation between cardiac output and body size. Br Heart J. 1963;25:425-430.
8. Lindberg L, Johansson S, Perez de Sa V. Validation of an ultrasound dilution technology to measure cardiac output and identify shunts in small children. Pediatr Crit Care Med. 2014;15:139-147.
9. Sigurdsson TS, Aronsson S, Lindberg L. Extracorporeal arteriovenous ultrasound measurement of cardiac output in small children. Anesthesiology. 2019;130:712-718.
10. Stewart G. Researches on the circulation time and on the influences which affect it. J Physiol. 1897;22:159-183.
11. Hamilton WF, Moore JW, Kinsman JM, Spurling RG. Studies on the circulation. Am J Physiology. 1932;99:534-151.
12. Reuter DA, Huang C, Edrich T, et al. Cardiac output monitoring using indicator-dilution techniques: Basics, limits, and perspectives. Anest Analg. 2010;110:799-811.
13. Boehme M, Baustert M, Paetzol V, et al. Determination of cardiac output by ultrasound dilution technique in infants and children: a validation study against direct Fick principle. Br J Anaesth. 2014;112:469-476.
14. Crittendon I, Dreyer WJ, Decker JA, Kim JJ. Ultrasonic dilution: An accurate means of determining cardiac output in children. Pediatr Crit Care Med. 2012;13:42-46.
15. Krivitski NM, Kislukhin VV, Thuramalla NV. Theory and in vitro validation of a new extracorporeal arteriovenous loop approach for hemodynamic assessment in pediatric and neonatal intensive care unit patients. Pediatr Crit Care Med. 2008;9:423-428.
16. Saxena R, Krivitski N, Peacock K, et al. Accuracy of the transpulmonary ultrasound dilution method for detection of small anatomic shunts. J Clin Monit Comput. 2015;29:407-414.
17. Dobson A, Kislukhin VV. Heart blood volume by dilution in patients on hemodialysis. ASAIO J. 2004;50:278-284.
18. Cecconi M, Rhodes A, Poloniecki J, et al. Bench-to-bedside review: the importance of the precision of the reference technique in method comparison studies with specific reference to the measurement of cardiac output. Crit Care. 2009;13:201.
19. Grindheim G, Eidet J, Bentsen G. Transpulmonary thermodilution (PiCCO) measurements in children without cardiopulmonary dysfunction: large interindividual variation and conflicting reference values. Paediatr Anaesth. 2016;26:418-424.
20. van der Ven JPG, Sadighy Z, Valsangiocomo Buechel ER, et al. Multicentre reference values for cardiac magnetic resonance imaging derived size and function for children aged 0–18 years. Eur Heart J Cardiovasc Imaging. 2020;21:102-113.
21. Poutanen T, Jokinen E, Sairanen H, et al. Left atrial and left ventricular function in healthy children and young adults assessed by three dimensional echocardiography. Br Heart J. 2003;89:544-549.
22. Buechel EV, Kaiser T, Jackson C, et al. Normal right- and left ventricular volumes and myocardial mass in children measured by steady state free precession cardiovascular magnetic resonance. J Cardiovasc Magn Reson. 2009;11:19.
23. Sarikouch S, Peters B, Gutberlet M, et al. Sex-specific pediatric percentiles for ventricular size and mass as reference values for cardiac MRI: assessment by steady-state free-precession and phase-contrast MRI flow. Circ Cardiovasc Imaging. 2010;3:65-76.
24. Galstyan G, Bychinin M, Alexanyan M, Gorodetsky V. Comparison of cardiac output and blood volumes in intrathoracic compartments measured by ultrasound dilution and transpulmonary thermodilution methods. Intensive Care Med. 2010;36:2140-2144.
25. Sigurdsson TS, Lindberg L. Estimation of intracardiac shunts in young children with a novel indicator dilution technology. Sci Rep. 2020;10:1337.
26. Lemson J, Merkus P, van der Hoeven JG. Extravascular lung water index and global end-diastolic volume index should be corrected in children. J Crit Care. 2011;26:432-437.
27. Lopez-Herce J, Bustinza A, Sancho L, et al. Cardiac output and blood volume parameters using femoral arterial thermodilution. Pediatr Int. 2009;51:59-65.
28. Schiffmann H, Erdelenbruch B, Singer D, et al. Assessment of cardiac output, intravascular volume status, and extravascular lung water by transpulmonary indicator dilution in critically ill neonates and infants. J Cardiovasc Anesth. 2002;16:592-597.
29. Vrancken SL, van Heijst AF, Hopman JC, et al. Hemodynamic volumetry using transpulmonary ultrasound dilution (TPUD) technology in a neonatal animal model. J Clin Monit Comput. 2015;29:643-652.
30. de Simone G, Devereux RB, Daniels SR, et al. Stroke volume and cardiac output in normotensive children and adults. Assessment of relations with body size and impact of overweight. Circulation. 1997;95:1837-1843.

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