Validation of a portable indirect calorimetry system for measurement of energy expenditure in sick preterm infants

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Abstract
A portable indirect calorimeter adapted from adult use was validated for use in preterm infants. Oxygen consumption (\(V_O_2\)) and carbon dioxide production (\(V_CO_2\)) were subsequently measured in 16 preterm infants breathing spontaneously in room air (canopy mode) and in nine preterm infants receiving intermittent positive pressure ventilation (ventilator mode). Validation of the system was performed using a gas injection technique with nitrogen to simulate \(V_O_2\) and carbon dioxide for \(V_CO_2\). Mean errors in validation of the canopy mode were 1-4% and 0-2% for \(V_O_2\) and \(V_CO_2\) with limits of agreement of 0-6 (+2SD) ml/min and -1-3 (-2SD) ml/min, and 0-9 (+2SD) ml/min and -2-3 (-2SD) ml/min respectively. In validation of the ventilator mode mean errors were -1-8% and -5-0% for \(V_O_2\) and \(V_CO_2\) with limits of agreement of 1-02 (+2SD) ml/min and -0-74 (-2SD) ml/min, and 0-93 (+2SD) ml/min and -1-45 (-2SD) ml/min respectively. Values of \(V_O_2\) and \(V_CO_2\) in 16 preterm infants in the canopy mode were 6-2 ml/kg/min (0-5 1SD) and 6-7 ml/kg/min (0-6 1SD) and in nine preterm infants in the ventilator mode 4-98 ml/kg/min (1-09 1SD) and 4-74 ml/kg/min (1-08 1SD) respectively. Mean energy expenditure was 45-5 kcal (191 kJ)/kg/day for infants measured in the canopy mode and 35-5 kcal (149 kJ)/kg/day for ventilated infants. This metabolic system can be adapted for use in the newborn but accuracy is reduced when it is used in those weighing less than 1000 g.

(Arch Dis Child 1992; 67:1207–11)

The measurement of the rate of oxygen consumption (\(V_O_2\)) and carbon dioxide production (\(V_CO_2\)) by indirect calorimetry may be used to calculate a number of parameters important in research into human metabolism. These include the respiratory quotient (RQ), energy expenditure, and rates of fat and carbohydrate utilisation. \(^1\) \(V_CO_2\) is also required when carbon labelled stable isotopes are infused for the measurement of substrate oxidation and estimation of whole body protein turnover. \(^2\) The first calorimeters used a closed circuit design \(^3\) and have progressed to open circuit indirect calorimeters \(^4\) and ventilated hood or canopy systems. \(^5\) Indirect calorimetry has been widely used in metabolic studies in adults and has provided much information about metabolic processes in both health \(^6\) and disease. \(^7\) Studies in children and infants are technically more difficult and this is particularly so for sick preterm infants. However a knowledge of energy expenditure and rates of substrate utilisation in this latter group is of vital importance in determining optimal environmental conditions and appropriate feeding regimens. There at present exists a paucity of calorimetry data on sick, and particularly ventilated, preterm infants.

Many factors have been described that may influence the measurement of energy expenditure in the infant including the effect of feeding, \(^8\) \(^9\) medical procedures, \(^10\) sleep state, \(^11\) maturity, \(^12\) and type of feed. \(^13\) Variability in measurements from day to day have been reported \(^14\) and investigation of infants over five days have been performed. \(^15\) These studies have concentrated on healthy growing infants and there is a need for suitable apparatus to be used with the sick preterm infant for similar work. The difficulties that arise when using indirect calorimetry in low birthweight infants are primarily due to their very small respiratory volumes. In ventilated hood or canopy systems the change in gas concentrations for infants of between 1000 and 2500 g will be in the order of 0-05 to 0-15% when a canopy ventilation rate of about 10 l/min is used. It is important that any system used to measure \(V_O_2\) and \(V_CO_2\) must be able to accurately determine such small differences. Previous investigators have reported variations from expected values in \(V_O_2, V_CO_2\) and RQ of between 2 to 5%. \(^15\) \(^16\)

In contrast to the measurement of \(V_O_2\) and \(V_CO_2\) in well, stable infants, there are few reports of their measurements in the ventilated, newborn infants. Lucas et al have reported measurement of \(V_CO_2\) but not \(V_O_2\), on 11 ventilated newborn infants between 26 and 32 weeks’ gestation. \(^17\) They have suggested that by measuring \(V_CO_2\) alone and taking RQ as 0-85 that errors in the calculation of energy expenditure would be no greater than 14%. In an artificial test lung recovery rates of carbon dioxide ranged from 97 to 99%. Richardson et al measured \(V_O_2\) but not \(V_CO_2\), by an oxygen replenishment technique in 14 infants with respiratory distress syndrome. \(^18\) Combined measurements of \(V_O_2\) and \(V_CO_2\) in sick infants, young children, and the newborn have been made using a metabolic monitor that employs a pneumotachometer, zirconium oxide oxygen sensor, and infrared carbon dioxide sensor. \(^19\) \(^20\) In those systems the...
assessment of $\dot{V}O_2$ is critically dependent on accurate measurement of the oxygen concentration entering the system and that the oxygen supplied is at an absolute constant concentration. Small changes in the ambient oxygen of 0.05% may lead to large errors of 30 to 100% in $\dot{V}O_2$. Measurements of $\dot{V}CO_2$ are less subject to error because of the low ambient concentrations of carbon dioxide. The lack of suitable validations in indirect calorimetry systems have been highlighted previously.\textsuperscript{21} In order to undertake metabolic studies in sick low birthweight infants we have validated a portable system (Deltatrac II Metabolic Monitor, Datex, Finland) that has been adapted for use in children by changing the flow rate through the apparatus, and tested it for its suitability for ventilated and non ventilated preterm infants. It is designed to be used as a ventilated canopy system, for those infants not requiring ventilatory support or oxygen therapy, and also in a closed circuit for ventilated infants.

**Patients and methods**

**MONITOR**

In the open canopy mode room air is drawn into a Perspex hood by a centrifugal fan at a constant flow rate of 10-3 l/min and pumped into the monitor. The Perspex hood covers the infant and a plastic skirt further covers the area of cot surrounding the infant creating a partial seal. Concentrations of carbon dioxide and oxygen are measured both in ambient room gas entering the hood, via an external sampling line, and in gas leaving the hood by an infrared carbon dioxide sensor and a paramagnetic differential oxygen sensor. With a known constant flow rate (Q), $\dot{V}O_2$ and $\dot{V}CO_2$ can be calculated from the difference between inspired and expired values; see equations (1) and (2):

\begin{align*}
(1) \quad &\dot{V}O_2 = Q \times (1-Fio_2) \times (F_{CO_2} - F_{CO_2}\times F_{O_2}) \\
(2) \quad &\dot{V}CO_2 = Q \times (F_{FEO_2} - F_{FICO_2})
\end{align*}

Where $Fio_2$ is the concentration of inspired oxygen, $FEO_2$ the concentration of expired oxygen, $FICO_2$ the concentration of inspired carbon dioxide, $FECO_2$ the concentration of expired carbon dioxide, $F_{o_2}$ is $Fio_2 - FEO_2$ and $F_{CO_2}$ is $FICO_2 - FECO_2$. The formula for $\dot{V}O_2$ includes the Haldane transformation.

In the ventilator mode gases expired from the ventilator are collected from the expiratory port and pass to a mixing chamber via wide bore tubing and a two litre low pressure reservoir (Ohmeda anaesthetic rebreathing bag) which limits pressure and flow oscillations in the system. Carbon dioxide concentration is measured in the mixing chamber and again after dilution to the known total flow rate. Thus by measurement of carbon dioxide dilution the minute expiratory flow rate can be calculated. $\dot{V}CO_2$ is calculated from the formulæ identical to that shown previously. Inspired oxygen concentration is measured from a sampling line, placed in the inspiration limb of the ventilator circuit, and the difference between this concentration and the mixed gas in the mixing chamber from the expiratory limb is measured using the differential paramagnetic oxygen sensor. By use of the Haldane transformation the RQ can be calculated and $\dot{V}O_2$ defined; see equations (3) and (4).

\begin{align*}
(3) \quad &RQ = \frac{\dot{V}CO_2/\dot{V}O_2}{1 - Fio_2} \\
(4) \quad &\dot{V}O_2 = RQ\dot{V}CO_2
\end{align*}

Figures 1 and 2 summarise the operating conditions.

All values are converted to standard temperature (0°C) and pressure (760 mm Hg) dry (STPD). The metabolic monitor uses tubing that is freely permeable to water vapour and thus equalises the water vapour pressure of the gas inside with that of the ambient air. All gas samples, including dry calibration and validation gases, pass through this tubing. Thus all measurements and calibrations are under identical conditions of humidity. Further detailed information of operating conditions and formulas used for calculations can be obtained.\textsuperscript{22}

**VALIDATION**

The constant flow of the system was measured by burning a 5 ml volume of 95% Analar grade ethanol ($CH_2CH_2OH$) in a glass test lung (supplied by Datex, Finland) attached to the system. From a known volume of ethanol the expected oxygen consumption and carbon dioxide production are calculated; see equation (5):

\begin{equation}
CH_2CH_2OH + 3O_2 \rightarrow 2CO_2 + 3H_2O
\end{equation}

As the volume occupied by a mol or g molecular weight of any gas at standard conditions is 22.414 litres (l) then it may be predicted that 1 g of ethanol consumes 3 x 22.414/46.069 = 1.46 l O_2 (STPD) and produces 2 x 22.414/46.069 = 0.973 l CO_2 (STPD)

This can be compared with the values measured by the apparatus and appropriate calibrations made. It was not possible with the system provided to reproduce rates of $\dot{V}CO_2$ and $\dot{V}O_2$ equivalent to those found in the neonate so this system was only used for calibration of the constant flow. Complete combustion of the ethanol was noted by the presence of a completely blue and even flame. Burning time for the ethanol was approximately 30 minutes.

To assess the accuracy of the measurement of $\dot{V}O_2$ and $\dot{V}CO_2$ in the canopy mode, at rates similar to those produced by preterm infants, nitrogen and carbon dioxide were infused at known rates into the canopy using certified calibration gas (Cryoservice Ltd, 25-2%±0.5% carbon dioxide and 74.8±1.5% nitrogen) and a certified precision flowmeter scaled for this gas mixture (Rotameter 1100 series, KDG Instruments). Appropriate corrections
Figure 1 Operating conditions for the ventilator mode.

Figure 2 Operating conditions for the canopy mode.

were made for operating temperature and pressure to give gas flows at standard temperature and pressure. In the ventilator mode gas was infused into the ventilator circuit via an artificial glass lung of compliance similar to that of an infant of 1500 g with respiratory distress syndrome. The ventilator was connected to gas supplies and humidifier in the normal way. With known infusion rates the injection of nitrogen and carbon dioxide will simulate VO₂ and VCO₂; see equations (6) and (7).

\[
\begin{align*}
(6) \quad & \dot{V}O_2 = (F_{iO_2} / (1 - F_{iO_2})) \times \dot{W}N_2 \\
(7) \quad & \dot{V}C0_2 = \dot{W}CO_2 (\dot{W}CO_2 = CO_2 \text{ injection})
\end{align*}
\]

Where \( \dot{W}N_2 \) is the nitrogen injection (ml/min) and \( \dot{W}CO_2 \) is the carbon dioxide injection (ml/min).

For the canopy system \( \dot{V}O_2 \) and \( \dot{V}C0_2 \) rates were simulated over a range of 5 to 24 ml/min and 4 to 30 ml/min respectively and for the ventilated system 4 to 50 ml/min and 3 to 34 ml/min respectively.

**Patients**

Determinations of \( \dot{V}O_2 \) and \( \dot{V}CO_2 \) were made in 16 infants using the canopy mode over a period of approximately one hour. Gestational age varied between 28 to 33 completed weeks (median 31 weeks) study weight 1190 g to 2460 g (median 1822 g), and postnatal age of 7 to 19 days (median 11 days). All infants were clinically stable and in air at the time of studies and at least one hour had elapsed after their last feed.

Nine ventilated infants, gestation 27 to 36 weeks (median 28 weeks), with weights between 880 g to 2500 g (median 1300 g) were studied in the closed loop ventilator mode. All infants were being ventilated with Draeger Babylog ventilators, size 3-0 mm endotracheal tubes, and receiving intravenous glucose or parenteral nutrition with the exception of one infant receiving all nutrition by the oral route. Ages ranged from 3 to 28 days old (median 4 days). The concentration of inspired oxygen in these infants was between 21% and 60%. An assessment of the leak from the endotracheal tube was made by measurement of carbon dioxide concentrations in a plastic mask covering the face. This included a slit which allowed the endotracheal tube and nasogastric tube to pass through the mask. A partial seal was created and air was drawn from the mask at a rate of 100 ml/min as measured by an 1100 series Rotameter flowmeter. Carbon dioxide concentrations were measured by infrared analysis (Beckman LB-2 Analyser, Beckman Instruments).

All infant temperatures were within the thermoneutral range for that infant at the time of the studies. Urine was collected during the period of the study for estimation of urinary nitrogen excretion (Un) for calculation of energy expenditure values. Energy expenditure (EE) for the infants was calculated from the classical Weir equation\(^{22}\); see equation (8):

\[
(8) \quad EE(\text{kcal/day}) = 5.67 \dot{V}O_2 (\text{ml/min}) + 1.59 \dot{V}CO_2 (\text{ml/min}) - 2.17 \text{Un} (\text{g/day})
\]

All studies were approved by the district ethical committee and informed consent was obtained from the parents prior to the study.

**Results**

**Validation**

The constant flow of gas through the analyser was calculated to be 10-3 l/min after three repeated burnings of ethanol showed less than 1% difference for the calculated value between each burning cycle. Mean error was calculated as [(measured value-true value)/true value]×100 where the true value is the test gas infusion. In addition the test gas infusion method, as the gold standard for reproducing \( \dot{V}CO_2 \) and \( \dot{V}O_2 \), was compared with values obtained from the metabolic monitor using the statistical method as proposed by Bland and Altman for limits of agreement.\(^{24}\)

This is able to give a 95% confidence level for individual measurements when compared with an absolute or standard and accurate method (that is test gas infusion).

Mean errors in validation of the canopy mode were 1.4% and 0.2% for \( \dot{V}O_2 \) and \( \dot{V}CO_2 \), with limits of agreement of 0.9 ml/min.
(+2SD) and −2.3 mls/min (−2SD), and 0.6 mI/min (+2SD) and −1.3 mls/min (−2SD) respectively. In validation of the ventilator mode mean errors were −1.8% and −5.0% for VO₂ and VCO₂ with limits of agreement of 1.02 ml/min (+2SD) and −0.74 ml/min (−2SD), and 0.93 ml/min (+2SD) and −1.45 ml/min (−2SD) respectively.

Mean values for infants measured in the canopy mode for VO₂ and VCO₂ were 6.2 ml/kg/min (0.5 1SD) and 6.7 ml/kg/min (0.6 1SD), RQ was 1.07 and mean energy expenditure was 45.5 kcal (191 kJ)/kg/day.

Mean values for infants measured in the ventilator mode for VO₂ and VCO₂ were 4.98 ml/kg/min (1.09 1SD) and 4.74 ml/kg/min (1.08 1SD), RQ was 0.95 and mean energy expenditure was 35.5 kcal (149 kJ)/kg/day. We were not able to detect a leak of greater than 0.1 ml/min carbon dioxide from the endotracheal tube in the three infants in whom measurements of leakage were made.

Discussion

There is a need for further understanding of energy requirements and substrate utilisation of different fuels in the preterm infant and newborn. Indirect calorimetry provides a means for studying this area. A system that is compact and mobile, providing the facility to measure both VO₂ and VCO₂ simultaneously is of great potential value in making indirect calorimetry measurements. However, rigorous validation of systems for the measurement of VO₂ and VCO₂ are needed as errors may be significant.

The system we have used is acceptable to the infants and their parents, is non-obtrusive in the canopy mode, and may be used in the laboratory or clinical setting. Used in the ventilator mode it may be integrated into the clinical environment without any disruption to normal ventilator function.

Validation of a system by gas injection allows measurement of VO₂ and VCO₂ at levels that are found in the preterm and newborn infant in conditions identical to those of normal clinical use. We did not have facility for measurement of the gas infusion separately by mass spectrometric methods.

Mean errors were small for both the canopy mode and ventilation mode. These errors would result in a maximum error of RQ measurement in the canopy mode of 1% and ventilator mode of 3%-5%. These errors are well within the previous published limits and make only a small difference to RQ estimation.

The limits of agreement for the canopy mode and the ventilator mode are of a similar order. The apparent errors when using the mean error for validation are small. By using the method of Bland and Altman it is possible to judge better the performance of the measuring apparatus over the whole range of values likely to be obtained in studies on newborn infants. This is preferable to using mean errors or correlation coefficients which may hide large errors in individual measurements. Our validation results show that in the small infant, particularly if less than 1000 g, there exists the possibility of significant errors in measurement. For such infants in the ventilator mode the 95% confidence limits for VO₂ and VCO₂ may be as high as 20% and 29% respectively if VO₂ and VCO₂ levels are each 5 ml/min (maximum error 1.02 ml/min and 1.45 ml/min respectively). The largest calculated error in RQ measurement for true values of 1 and 0.7 would be −0.4 and −0.3, which are unacceptable for most purposes. At higher rates of VO₂ and VCO₂ these variations become less significant. We would suggest that limits of agreement be published when validating any new system. As this approach to presenting results has not previously been extensively used then there exists no previous data for comparison with other systems.

Measurements are difficult to make in situations of raised oxygen concentration in the ventilator mode where fluctuations may occur in inspired oxygen levels. This problem is reduced by sampling the inspired oxygen concentration after the humidifier which acts as a mixing chamber and reduces fluctuations. In addition it is important to note that the system as described for our methods used the Draeger Babylog ventilator and capture of expiratory gas from the expiratory outlet of the ventilator. In the other ventilator commonly used on our unit and other neonatal units (Sechrist infant ventilator IV-100B, EME) there is mixing, at the expiratory outlet, of gas from the infant with gas from a purging system that is a different concentration from that received by the infant. This results in considerable errors due to dilution of the expired gas. We have tried to eliminate this problem by use of an oxygen-air mixer between the external oxygen and air supplies and the ventilator blender, so that the purging gas and inspired gas are at the same concentration. However large systematic variations in VO₂ and VCO₂ and undermeasurement of these values still occur in infants measured this way. High levels of RQ errors may be seen if this ventilator exceeding 10.3/min and loss of expired gas from the collecting system would explain this error. Before using this system with any other ventilators other than the Draeger Babylog it will be necessary for investigators to validate their measurements.

Values for VO₂ and VCO₂ in the infants in our study are similar to those previously published for infants measured in open hood systems and for those on ventilators. Energy expenditure values are also comparable with other published figures.

The mean RQ for those infants in the canopy mode was 1.07. Other studies have reported RQ values >1 in the neonate and this suggests that lipogenesis from glucose is occurring. Our values were obtained over a period of 60 minutes and postprandial by at least one hour. Had we continued studies longer then net lipid synthesis from glucose may not be apparent. This equipment would be appropriate for such studies. Because the canopy mode operates in conditions of ambient air it is not recommended for use in infants receiv-
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