Left ventricular function in persistent pulmonary hypertension of the newborn

Computer analysis of the echocardiogram

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SUMMARY Regional and global left ventricular function was assessed in 23 neonates with persistent pulmonary hypertension using computer assisted analysis of their left ventricular echocardiograms and compared with that in 50 healthy neonates. End diastolic left ventricular dimension was normal and end systolic dimension increased while percentage left ventricular shortening and peak velocity of circumferential fibre shortening decreased indicating impaired systolic performance. The peak rate of increase in left ventricular diameter in early diastole was significantly decreased and the durations of the rapid filling and isovolumic relaxation periods were prolonged suggesting resistance to left ventricular filling due to changes in diastolic myocardial properties. This abnormal left ventricular cavity function may have been due to a combination of increased diastolic wall thickness, reduced percentage systolic wall thickening, increased relative wall thickness, and pronounced reduction in peak rates of systolic wall thickening and diastolic wall thinning.

Seven neonates with persistent pulmonary hypertension died, and of the three examined at necropsy all had left ventricular hypertrophy and two extensive subendocardial haemorrhage and infarction affecting the right and left ventricular papillary muscles. Thus left ventricular dysfunction appears to be a common feature in neonates with this disorder and may be readily detected using computer analysis of left ventricular echocardiograms. Unfortunately, no single echo measurement was useful prognostically. Left ventricular dysfunction in persistent pulmonary hypertension probably results from a combination of hypoxaemia, acidaemia, and pulmonary hypertension, and although it may contribute to the high mortality in this syndrome, a correlation between the severity of left ventricular dysfunction and clinical outcome could not be shown.

Persistent pulmonary hypertension of the newborn, or persistent fetal circulation, describes a syndrome occurring in normal birthweight neonates with normal cardiac anatomy in whom the pulmonary vascular resistance fails to fall in the immediate neonatal period. This results in reduced pulmonary blood flow, persistent pulmonary hypertension, tachypnoea, acidaemia, and cyanosis due to right to left shunting via the ductus arteriosus and the foramen ovale. In most infants with this disorder no definitive precipitating or causative factor is apparent.

The presence of left ventricular dysfunction has recently been recognised and may possibly contribute to the unacceptably high mortality rate. The purpose of this study was to assess the frequency and severity of left ventricular dysfunction in persistent pulmonary hypertension and to determine whether or not its presence affected prognosis or contributed to mortality. To achieve these aims we used computer assisted analysis of the left ventricular echocardiograms, a technique that has been used to evaluate both diastolic and systolic global and regional function in children and adults. Left ventricular function in neonates with persistent pulmonary hypertension was then compared with that in 50 healthy neonates.
Patients and methods

CONTROLS
Echocardiograms were recorded from 50 clinically healthy neonates (age range 2–64 hours, mean age 19 hours); 26 were girls and 24 boys with birth weights ranging from 2-6 to 4-5 kg as previously reported. These strict criteria were selected so that there was no doubt that all the neonates in this study had persistent pulmonary hypertension, but in so doing we included those with the most severe disease which may represent only the severe end of this disease spectrum. Table 1 shows individual patient data, respiratory assistance, blood gas pressures, blood glucose concentrations, adjuvant drug treatment, and clinical outcome.

STUDY GROUP
Echocardiograms were recorded from 23 neonates with persistent pulmonary hypertension (10 girls and 13 boys, age range 12–96 hours) whose birth weights ranged from 2-4 to 4-7 kg. Neonates were included in this study only if they fulfilled all of the following criteria: (a) full term gestation (>38 weeks); (b) normal birth weight; (c) no evidence of anatomical cardiac abnormality either clinically or by M-mode and cross sectional echocardiography (when the diagnosis was uncertain and congenital heart disease suspected, as it was in two infants, cardiac catheterisation was performed; this showed pulmonary hypertension and normal cardiac anatomy); (d) pulmonary arterial oxygen pressure (Pao2) <50 mm Hg on maximum ventilatory support with a fractional inspired oxygen content (FiO2) of 100%, an inspiratory pressure above 32 cm of water, and positive end expiration pressure (PEEP) of 4–10 cm of water; and (e) sinus rhythm with no electrocardiographic evidence of conduction disturbances.

ECHOCARDIOGRAPHY
Echocardiograms were obtained with a Hoffrel ultrasonoscope using an Aerotech 5 mHz non-focused 6 mm diameter transducer providing an axial resolution of 0-4 mm with a repetition frequency of 1000 cycles/s. Recordings were made with an Irex multichannel physiological recorder at paper speeds of 75–100 mm/s with a simultaneous electrocardiographic lead chosen to demonstrate most clearly early ventricular depolarisation (Q waves). A true maximal minor axis of the left ventricular cavity was obtained by scanning from the apex of the left ventricle to the aorta and left atrium from a perpendicular recording position of the mitral valve echo. Once the mitral ring echo was identified the transducer was tilted inferiorly until strong ventricular and septal endocardial echoes were visualised (Fig. 1).

Echocardiograms were digitised by one of us without any knowledge of the clinical findings. Although

Fig. 1  Echocardiogram of a healthy neonate showing right and left ventricles. AW, anterior wall; VS, septum; PW, posterior wall; MV, mitral valve; RV, right ventricle.
### Table 1: Summary of clinical data

<table>
<thead>
<tr>
<th>Case No</th>
<th>Birth weight at echo (kg)</th>
<th>Age in days at echo (h)</th>
<th>Admission diagnosis</th>
<th>FIO(2)/Pao(2) (% &amp; kPa)</th>
<th>Paco(2) (% &amp; kPa)</th>
<th>pH</th>
<th>Blood glucose concentration (mmol/l)</th>
<th>Respiratory assistance (pressures in mm H(_2)O)</th>
<th>Tolazoline</th>
<th>Outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2-7</td>
<td>72</td>
<td>Meconium aspiration</td>
<td>1/0-5-6</td>
<td>5-8</td>
<td>7-36</td>
<td>2-5</td>
<td>Intubated (inspiration pressure 34, PEEP 6)</td>
<td>No</td>
<td>Discharged well at 12 days</td>
</tr>
<tr>
<td>2</td>
<td>4-6</td>
<td>16</td>
<td>PPHN, infant of diabetic mother</td>
<td>1/0-4-6</td>
<td>7-7</td>
<td>7-24</td>
<td>4-5</td>
<td>Intubated (inspiration pressure 36, PEEP 6)</td>
<td>No</td>
<td>Died at 24 h: necropsy showed LVH+RV+LV papillary muscle necrosis and subendocardial haemorrhage and infarction</td>
</tr>
<tr>
<td>3</td>
<td>3-2</td>
<td>36</td>
<td>PPHN, CHD</td>
<td>1/0-6-0</td>
<td>6-6</td>
<td>7-34</td>
<td>4-9</td>
<td>Intubated (inspiration pressure 30, PEEP 4)</td>
<td>Yes and dopamine</td>
<td>Discharged well at 16 days</td>
</tr>
<tr>
<td>4</td>
<td>3-5</td>
<td>48</td>
<td>CHF, ?PPHN, birth asphyxia</td>
<td>1/0-4-0</td>
<td>9-4</td>
<td>7-26</td>
<td>1-9</td>
<td>Intubated (inspiration pressure 30)</td>
<td>No</td>
<td>Died at 9 days: necropsy showed LVH, subendocardial haemorrhage and RV and LV papillary muscles</td>
</tr>
<tr>
<td>5</td>
<td>3-2</td>
<td>12</td>
<td>?Pneumonia, PPHN, ?pneumothorax</td>
<td>1/0-4-5</td>
<td>8-1</td>
<td>7-14</td>
<td>5-3</td>
<td>Intubated (inspiration pressure 40, PEEP 4)</td>
<td>Yes</td>
<td>Died at 2 days. No necropsy, became cyanosed and hypotensive led to cardiac arrest</td>
</tr>
<tr>
<td>6</td>
<td>2-9</td>
<td>36</td>
<td>PPHN</td>
<td>1/0-5-7</td>
<td>8-8</td>
<td>7-16</td>
<td>3-7</td>
<td>Intubated (inspiration pressure 40, PEEP 6)</td>
<td>Yes</td>
<td>Steady improvement, discharged at 27 days</td>
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<td>7</td>
<td>3-7</td>
<td>12</td>
<td>Aspiration, infant of diabetic mother, CHD, PPHN</td>
<td>1/0-5-2</td>
<td>7-2</td>
<td>7-25</td>
<td>5-9</td>
<td>Intubated (inspiration pressure 40, PEEP 6)</td>
<td>Yes</td>
<td>Died at 24 h: cardiac catheterisation: RV 80/7, PA 80/40, LV 80/8, Ao 80/40. R to L shunt at PDA, normal cardiac anatomy. No necropsy</td>
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<tr>
<td>8</td>
<td>3-3</td>
<td>18</td>
<td>Infant of diabetic mother, CHF, PPHN</td>
<td>1/0-5-7</td>
<td>7-8</td>
<td>7-25</td>
<td>3-6</td>
<td>Intubated (inspiration pressure 40, PEEP 10)</td>
<td>No</td>
<td>Discharged well at 17 days</td>
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<tr>
<td>9</td>
<td>2-8</td>
<td>24</td>
<td>Aspiration, ?pneumothorax, ?PPHN</td>
<td>1/0-6-6</td>
<td>7-2</td>
<td>7-27</td>
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<td>Intubated (inspiration pressure 48, PEEP 4)</td>
<td>Yes and dopamine</td>
<td>Discharged well at 27 days</td>
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<td>10</td>
<td>3-1</td>
<td>12</td>
<td>CHD, ?PPHN</td>
<td>1/0-6-1</td>
<td>7-2</td>
<td>7-40</td>
<td>5-6</td>
<td>Intubated (inspiration pressure 40, PEEP 6)</td>
<td>No</td>
<td>Discharged well at 11 days</td>
</tr>
<tr>
<td>11</td>
<td>3-7</td>
<td>48</td>
<td>Meconium aspiration, infant of diabetic mother</td>
<td>1/0-4-6</td>
<td>13-1</td>
<td>6-97</td>
<td>3-2</td>
<td>Intubated (inspiration pressure 40)</td>
<td>Yes and epinephrine</td>
<td>Died at 9 days: necropsy, LVH and mild LV dilatation</td>
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<tr>
<td>12</td>
<td>3-2</td>
<td>8</td>
<td>Aspiration, ?PPHN</td>
<td>1/0-4-0</td>
<td>8-6</td>
<td>7-12</td>
<td>3-9</td>
<td>Intubated (inspiration pressure 32)</td>
<td>No</td>
<td>Discharged well at 14 days</td>
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<td>4-1</td>
<td>12</td>
<td>?TGA, ?PPHN</td>
<td>1/0-4-6</td>
<td>6-6</td>
<td>7-28</td>
<td>4-8</td>
<td>Intubated (inspiration pressure 36, PEEP 8)</td>
<td>No</td>
<td>Cardiac catheterization: RV 70/7, PA 70/49, LV 65/12, Ao 63/48, R to L shunt at PDA, normal cardiac anatomy; discharged well at 22 days</td>
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<tr>
<td>14</td>
<td>4-0</td>
<td>12</td>
<td>?PPHN, ?CHF</td>
<td>1/0-5-3</td>
<td>6-5</td>
<td>7-33</td>
<td>4-0</td>
<td>Intubated (inspiration pressure 38, PEEP 8)</td>
<td>Yes and epinephrine</td>
<td>Progressive increase in cyanosis and hypotension; died from cardiac arrest at 3 days. No necropsy</td>
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<tr>
<td>15</td>
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<td>12</td>
<td>Meconium aspiration, ?PPHN</td>
<td>1/0-3-6</td>
<td>4-6</td>
<td>7-40</td>
<td>3-4</td>
<td>Intubated (inspiration pressure 40, PEEP 6)</td>
<td>Yes and dopamine</td>
<td>Discharged well at 11 days</td>
</tr>
<tr>
<td>16</td>
<td>2-9</td>
<td>30</td>
<td>?Sepsis, pneumonia, hypotension ?cause peritonitis</td>
<td>1/0-6-5</td>
<td>5-7</td>
<td>7-27</td>
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<td>Intubated (inspiration pressure 32, PEEP 6)</td>
<td>No</td>
<td>Discharged well at 30 days</td>
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<tr>
<td>17</td>
<td>4-1</td>
<td>24</td>
<td>?PPHN</td>
<td>1/0-5-4</td>
<td>4-6</td>
<td>7-39</td>
<td>4-7</td>
<td>Intubated (inspiration pressure 32)</td>
<td>No</td>
<td>Discharged well at 11 days</td>
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<tr>
<td>18</td>
<td>3-5</td>
<td>24</td>
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<td>1/0-5-7</td>
<td>6-1</td>
<td>7-29</td>
<td>4-3</td>
<td>Intubated (inspiration pressure 34, PEEP 4)</td>
<td>Yes and dopamine</td>
<td>Discharged well at 19 days</td>
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<tr>
<td>19</td>
<td>3-9</td>
<td>36</td>
<td>PPHN</td>
<td>1/0-4-8</td>
<td>6-1</td>
<td>7-40</td>
<td>3-9</td>
<td>Intubated (inspiration pressure 32)</td>
<td>No</td>
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<tr>
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<td>120</td>
<td>Aspiration</td>
<td>1/0-6-0</td>
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<td>7-32</td>
<td>4-4</td>
<td>Intubated (inspiration pressure 40, PEEP 6)</td>
<td>No</td>
<td>Discharged well at 34 days</td>
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<td>21</td>
<td>3-2</td>
<td>24</td>
<td>PPHN</td>
<td>1/0-4-9</td>
<td>6-8</td>
<td>7-33</td>
<td>3-6</td>
<td>Intubated (inspiration pressure 34)</td>
<td>Yes and dopamine</td>
<td>Died at 3 days: hypotension and increasing cyanosis led to cardiac arrest</td>
</tr>
<tr>
<td>22</td>
<td>4-7</td>
<td>24</td>
<td>Asphyxia, meconium aspiration</td>
<td>1/0-5-2</td>
<td>5-7</td>
<td>7-26</td>
<td>3-0</td>
<td>Intubated (inspiration pressure 36)</td>
<td>No</td>
<td>Discharged well at 15 days</td>
</tr>
<tr>
<td>23</td>
<td>2-7</td>
<td>48</td>
<td>?PPHN, ?CHD</td>
<td>1/0-4-9</td>
<td>4-9</td>
<td>7-38</td>
<td>2-0</td>
<td>Intubated (inspiration pressure 40, PEEP 8)</td>
<td>No</td>
<td>Discharged well at 16 days</td>
</tr>
</tbody>
</table>

CHD, congenital heart defect; CHF, congestive heart failure; LV, left ventricular; PEEP, positive end expiratory pressure; PPHN, persistent pulmonary hypertension of the newborn; RV, right ventricular; TGA, transposition of the great arteries; FIO\(2\), fractional inspired oxygen; Paco\(2\), arterial carbon dioxide pressure; PaO\(2\), arterial oxygen pressure.

Conversion: ST to traditional units—glucose: 1 mmol/l = 18-01 mg/100 ml; arterial pressure: 1 kPa = 7-5 mm Hg.
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Throughout the cardiac cycle were digitised as described using a Digisonics digitiser and microprocessor. To minimise errors the depth scale was expanded so that each cm exceeded 15 mm. Echoes were calibrated with points defining a time interval of 1 s, a depth of 3 cm, and two successive Q waves on the EKG, enclosing the cardiac cycle to be analysed. Data points were generated at intervals of 10 ms so that strings of XY coordinates were obtained for the abovementioned surface boundaries. From these data plots were obtained from an online incremental plotter of continuous left ventricular cavity dimension and posterior left ventricular wall thickness and their respective rates of change expressed in cm/s or normalised by dividing by instantaneous cavity dimension or posterior wall thickness (Fig. 2). From these plots the following measurements were made: (a) heart rate (beats/minute); (b) dimensions—(i) end systolic and end diastolic left ventricular cavity dimension (mm), (ii) end systolic and end diastolic posterior wall thickness (mm), and (iii) end diastolic relative wall thickness (ratio of posterior wall thickness to left ventricular cavity radius at end diastole (h/R)); (c) left ventricular systolic function—(i) percentage left ventricular cavity shortening, (ii) peak velocity of circumferential fibre shortening (peak VCF/s), (iii) percentage systolic thickening of the posterior left ventricular wall, and (iv) normalised peak rate of posterior left ventricular wall thickening (per second); and (d) left ventricular diastolic function—(i) peak rate of increase in left ventricular dimension (cm/s), (ii) normalised peak rate of left ventricular wall thinning (s⁻¹), (iii) duration of rapid filling (ms) defined as the time interval from minimum left ventricular cavity dimension to that at which the peak rate of increase in left ventricular dimension fell to 20% of its peak value,⁹ and (iv) duration of the isovolumic relaxation period (ms) defined as the time interval from minimum left ventricular dimension to the onset of the initial separation of the mitral valve leaflets.

Reproducibility and statistical method
Although the theoretical axial resolution of the 5 mHz transducer driven by the Hoffrell ultrasonoscope was 0.4 mm we recorded wall thickness and left ventricular dimensions from the online printer to the nearest millimetre.

The significance of differences between the controls and the study group was assessed using the unpaired Student's t test. Intraobserver beat to beat variability in left ventricular dimension and its rate of change was 0–7% (mean 5%). The interobserver error in the same measurements among four individuals varied from 0 to 12% (mean 7%), provided that there were optimal expansion of the depth scale and high paper speeds.
Results

HEART RATE
Heart rates in the controls varied from 110 to 146 beats/minute (mean 124), which were similar to those in the study group in whom heart rate varied from 108 to 150 beats/minute (mean 127), enabling a comparison to be made of cavity and regional ventricular dynamics.

DIMENSIONS
End systolic and end diastolic left ventricular cavity size, wall thicknesses, and relative wall thickness at end diastole for the study group and controls are shown in Table 2. End diastolic and end systolic left ventricular dimensions in the two groups varied over the same wide range; however, while the mean end diastolic dimension was normal in the study group mean end systolic dimension was increased (p<0-03) (Table 2) (Fig.3). Left ventricular wall thicknesses at end systole and end diastole were recorded to the nearest millimetre. Although differences in wall thicknesses between the study group and the controls were small, all records were digitised by only one of us who had no knowledge of the clinical details, thus obviating interobserver error but incurring a possible mean beat to beat variability of 5% for each group. Nevertheless, mean values for left ventricular wall thicknesses at end diastole were significantly increased in the study group compared with the controls (p<0.001) (Fig. 3). In addition, diastolic relative wall thickness or h/R ratio was also significantly increased in the study group (p<0.001) (Fig. 3) (Table 2).

DIASTOLIC VENTRICULAR FUNCTION (FIG. 3)
There was little overlap of the peak rates of increase in left ventricular cavity diameter in diastole and the peak rates of left ventricular wall thinning between the study group and the controls, and the mean values were significantly decreased in the former (p<0.001). In addition the duration of the rapid filling period was increased (p<0.001) (Table 2). These abnormal left ventricular filling characteristics are shown on the plot of continuous left ventricular dimensions and its first derivative (Fig. 2); left ventricular filling in the study group was not only slowed but also prolonged, with shortening or abolition of the normal period of diastasis. The isovolumic relaxation period, which we were able to measure in only 14 patients as the time interval from minimum left ventricular dimension to mitral valve opening, was also significantly longer in the study group than in the controls (Table 2) suggesting reduced left ventricular diastolic compliance.

**Table 2**  Left ventricular cavity and wall dynamics in study group and controls. Results are means ± SD

<table>
<thead>
<tr>
<th></th>
<th>Study group (n = 23)</th>
<th>Total study group</th>
<th>Controls (n = 50)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Survivors (n = 16)</td>
<td>Non-survivors (n = 7)</td>
<td></td>
</tr>
<tr>
<td>End diastolic LV dimension (mm)</td>
<td>17.5±3.4 15.6±5.0</td>
<td>17.3±5.6</td>
<td>17.3±2.6</td>
</tr>
<tr>
<td>End systolic LV dimension (mm)</td>
<td>11.8±2.7 10.0±3.7</td>
<td>11.7±2.9***</td>
<td>9.6±2.2</td>
</tr>
<tr>
<td>% LV shortening fraction</td>
<td>33±9 36±5</td>
<td>33±8**</td>
<td>44±6</td>
</tr>
<tr>
<td>Peak VCF/s</td>
<td>3.0±0.9 3.7±0.8</td>
<td>3.0±0.9*****</td>
<td>3.8±1.0</td>
</tr>
<tr>
<td>Peak LV filling rate dD/dt diastole (cm/s)</td>
<td>3.9±1.3 3.8±0.8</td>
<td>4.0±1.1*</td>
<td>7.7±2.9</td>
</tr>
<tr>
<td>Duration of rapid filling period (ms)</td>
<td>171±30 184±37</td>
<td>177±33</td>
<td>138±28</td>
</tr>
<tr>
<td>Isovolumic relaxation period</td>
<td>40±15***</td>
<td>40±15***</td>
<td>15±10</td>
</tr>
<tr>
<td>Wall thickness (mm)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Minimum diastolic</td>
<td>3.1±0.8 4.0±1.1</td>
<td>3.5±0.9*</td>
<td>2.5±0.6</td>
</tr>
<tr>
<td>Maximum systolic</td>
<td>5.7±1.1 6.4±1.3</td>
<td>5.9±1.1</td>
<td>5.4±0.8</td>
</tr>
<tr>
<td>Systolic wall thickening (%)</td>
<td>86±28 70±23</td>
<td>78±28*</td>
<td>130±50</td>
</tr>
<tr>
<td>Relative wall thickness at end diastole</td>
<td>0.37±0.09 0.45±0.06</td>
<td>0.4±0.08**</td>
<td>0.29±0.05</td>
</tr>
<tr>
<td>Peak rate systolic wall thickening</td>
<td>4.1±1.0 3.8±1.3</td>
<td>3.9±1.1*</td>
<td>5.4±1.3</td>
</tr>
<tr>
<td>Peak rate diastolic wall thinning</td>
<td>4.5±1.4 4.6±1.3</td>
<td>4.6±1.4*</td>
<td>8.0±2.2</td>
</tr>
</tbody>
</table>

Significant difference (study group vs controls): *p<0.001, **p<0.005, ***p<0.01, ****p<0.02, *****p<0.08. LV, left ventricular; VCF, velocity of circumferential fibre shortening.

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Fig. 3 Plot of (a) left ventricular cavity dimension in healthy neonates (control group) and those with PPHN (study group) at end diastole and end systole ($x = \text{non-survivors}$); (b) posterior left ventricular wall thickness; (c) diastolic relative wall thickness and percentage systolic wall thickening, (d) peak velocity of circumferential fibre shortening and left ventricular fractional shortening, (e) peak rate of change in left ventricular wall thickness in diastole, and (f) peak left ventricular filling rate ($dD/dt$ diastole) and duration of the rapid diastolic filling period.
The study group was subdivided into those who survived and those who died. Wall and cavity dimensions and dynamics were compared in an attempt to identify any measure of left ventricular function which might be useful prognostically. There was no difference in left ventricular cavity size or wall thickness, and those who died could not be differentiated by measurements of cavity or regional systolic or diastolic function (Fig. 3). In both groups, as in the study group as a whole, there was greater impairment of diastolic than systolic cavity and regional left ventricular function.

**CLINICAL OUTCOME**

Of the 23 neonates in the study group, seven (30%) died from one to nine days (mean 3-9 days) after birth (Table 1). Three of the seven neonates who died were examined at necropsy. All three had concentric left ventricular hypertrophy and one also mild left ventricular cavity dilatation. Two of the three had severe subendocardial haemorrhage and papillary muscle necrosis in both the right and left ventricles with extensive subendocardial infarction of both ventricles.

**Discussion**

Persistent pulmonary hypertension of the newborn has been regarded primarily as a pulmonary vascular or right heart disorder. Computer analysis of the left ventricular echocardiograms in the 23 neonates in this study showed significant abnormalities of both systolic and diastolic left ventricular function. About one quarter of the patients died despite intubation and maximum ventilatory support with increased inspiratory pressure, correction of acidaemia, and treatment with vasodilator drugs. This inordinately high mortality may have been due in part to our selecting only infants with the most severe pulmonary hypertension for inclusion in the study.

Although M-mode echocardiography permits static measurements of ventricular chamber size, a possible shortcoming in assessing dynamic left ventricular function is that only a basal slice of the ventricle is reproducibly visualised. Nevertheless, in the absence of segmental abnormalities, which we presumed to be the case in these neonates with generalised hypoxaemia, echocardiographic recordings from the base were regarded as representative of the left ventricle as a whole.

Although end diastolic left ventricular cavity size was normal, percentage fractional shortening and peak velocity of circumferential shortening varied over a wide range. While these systolic variables could not reliably identify individual neonates with persistent pulmonary hypertension, the mean values for the group as a whole were significantly lower than for the controls, indicating impaired contractile function overall. More impressive even than the abnormalities of systolic cavity function was the impairment of diastolic cavity dynamics manifested as abnormal left ventricular filling characteristics. The rate of left ventricular filling was universally decreased with virtually no overlap with the controls. The duration of the rapid diastolic filling period was also prolonged in most infants in the study group with consequent shortening or abolition of the normal period of diastasis (Fig. 2). Since the mitral valve leaflets and their motion pattern were normal and did not therefore obstruct blood flow at the left ventricular inflow tract the reduction in left ventricular filling was most likely to have been caused by decreased myocardial compliance, resulting at least in part from the concentric left ventricular hypertrophy. Further evidence for decreased left ventricular compliance in persistent pulmonary hypertension was the prolongation of isovolumic relaxation.4-13

Left ventricular diastolic wall thickness varied, but the mean value for the study group as a whole was significantly increased compared with that for the controls, and in addition, the presence of concentric left ventricular hypertrophy was confirmed in the three neonates who were examined at necropsy. Relative wall thickness, which is virtually constant in the normal human left ventricle from birth to old age,11 14 was significantly increased and may have resulted in part from the increased systemic vascular resistance recently reported in this disorder.15

Regional myocardial function, the integral of which determines left ventricular cavity function, was also severely abnormal. Systolic and diastolic wall dynamics, which directly reflect myocardial contraction and relaxation respectively, were both significantly reduced. This impairment in cardiac muscle function in the study group could not be accounted for by any differences in heart rate, as these were the same as in the controls. It may, however, have resulted in part from the increased wall thickness and changed cavity architecture (relative wall thickness). The greater impairment of diastolic than systolic regional function was unexplained, but it was consistent with the similarly greater impairment of diastolic cavity function, since cavity function is simply an expression of endocardial movement, the determinants of which are muscle thickening and thinning.

To determine whether this detailed computer analysis of left ventricular function could identify some measurement of regional or cavity dynamics which might prove useful in prognosis, we compared survivors with non-survivors. Systolic and diastolic cavity and regional function were, however, equally abnormal in both groups (Table 2), and no single or combination of echocardiographic measurements
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proved useful in differentiating between the two groups.

The precise aetiology of the left ventricular dysfunction in persistent pulmonary hypertension of the newborn is not resolved, but there are four important contributing factors: pulmonary hypertension by itself, alteration in left ventricular geometry by the pressure overloaded right ventricle, hypoxaemia resulting in generalised myocardial ischaemia, and metabolic acidaemia.

The effects of pulmonary hypertension by itself on left ventricular function have been evaluated in adults with pulmonary artery pressures at subsystemic values and have shown reduced rates of diastolic filling and decreased systolic performance. Furthermore, in experimental animals with moderate right ventricular hypertension myocardial blood flow to the left as well as the right ventricle is significantly reduced, and it has been suggested that this may account for the reduced left ventricular compliance and secondary resistance to left ventricular filling. This mechanism may be concerned in the pathogenesis of left ventricular dysfunction in persistent pulmonary hypertension, in which the right ventricle often develops systemic or greater than systemic pressures.

A further possible cause of left ventricular dysfunction in persistent pulmonary hypertension is that the pressure overloaded right ventricle changes the geometry of the left ventricle by displacing it more posteriorly or by altering the direction of septal motion, thus changing the region of the left ventricle intersected by the M-mode echo beam. Although this is a theoretical consideration, septal motion was normal in all infants with persistent pulmonary hypertension. Moreover, previous studies in patients with pronounced right ventricular volume overload with or without septal abnormalities and in right ventricular pressure overload with normal septal motion failed to show any evidence of regional or global left ventricular dysfunction. It seems unlikely therefore that abnormal geometry was a major cause of the left ventricular dysfunction in our study group.

The role of hypoxaemia in causing myocardial ischaemia and subsequent left ventricular dysfunction is not clearly established. In the normal heart, myocardial oxygen extraction is near maximal and anaerobic reserves are minimal. In persistent pulmonary hypertension of the newborn in which PaO₂ is <6-6 kPa (<50 mm Hg), the baseline delivery is reduced which, coupled with the possible reduction in myocardial perfusion due to pulmonary hypertension and the increased muscle mass from left ventricular hypertrophy, may exacerbate the imbalance between myocardial oxygen supply and demand resulting in global myocardial ischaemia.

More direct evidence for severe hypoxaemia resulting in ischaemia is provided by the subendocardial infarction which has been found at necropsy in neonates with severe cyanosis from congenital heart disease and normal coronary anatomy. It is of particular interest that these subendocardial infarctions are present in the left as well as the right ventricle. It was not until very recently, however, that myocardial infarction and papillary muscle necrosis in the right and left ventricles were recognised in neonates with severe hypoxaemia due to birth asphyxia, aspiration, intracranial haemorrhage, and hyaline membrane disease in the absence of any congenital heart disease.

Myocardial ischaemia has also been shown by thallium imaging in neonates with hypoxaemia due to the respiratory distress syndrome. Such severe hypoxaemia is commonplace in persistent pulmonary hypertension of the newborn. Of our three neonates who were examined at necropsy, two had severe subendocardial haemorrhage and infarction in the right and left ventricles indicating myocardial ischaemia as a probable aetiological factor, which in the most severe circumstances results in infarction and when of lesser severity results in major left ventricular dysfunction.

The fourth major contributing factor to left ventricular dysfunction is metabolic acidaemia, which even in the absence of hypoxaemia may decrease myocardial contractile function. When acidaemia is accompanied by severe hypoxaemia, as it is in persistent pulmonary hypertension of the newborn, the deleterious effect is greater than with either hypoxaemia or acidaemia alone, because acidaemia potentiates the vasoconstrictor effect of hypoxaemia. Thus a vicious cycle ensues in which further pulmonary vasoconstriction and rise of pulmonary artery pressure causes increased right to left shunting, greater oxygen desaturation, and exacerbation of pulmonary arterial vasospasm. Although most of our patients were in relative acid base balance at the time of their echocardiograms, the effects of earlier acidaemia and hypoxaemia on left ventricular function may still have been present.

We conclude that there is severe left ventricular dysfunction in persistent pulmonary hypertension of the newborn, with consistent abnormalities of diastolic properties which can be readily recognised by computer analysis of the echocardiogram. Unfortunately, however, no single or combination of echo measurements of left ventricular function were useful in determining prognosis in individual patients. The cause of the left ventricular dysfunction was most likely due to a combination of hypoxaemia, acidaemia, pulmonary hypertension, and possibly changes in left ventricular geometry, but we were unable to assess the respective contributions of each.
Caution must be taken in interpreting the incidence of left ventricular dysfunction in persistent pulmonary hypertension of the newborn from this study, since this disorder has a wide spectrum of severity and the criteria we used to select our patients tended to include only those with severe disease. Finally, the echocardiographic and necropsy findings of left ventricular disease in persistent pulmonary hypertension of the newborn, although not correlating with clinical outcome, may still play a part in the high mortality rate in newborns with this disorder.

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Left ventricular function in persistent pulmonary hypertension of the newborn. Computer analysis of the echocardiogram.

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