SHORT REPORT

Gastrointestinal manifestations of postnatal cytomegalovirus infection in infants admitted to a neonatal intensive care unit over a five year period

J L Y Cheong, F M Cowan, N Modi


Sixteen cases of postnatal cytomegalovirus (CMV) infection were identified in a neonatal intensive care unit population over a five year period. Eleven of these infants had gastrointestinal signs at the time of presentation. These ranged from minor and transient (abdominal distension and enteral feed intolerance) to severe and life threatening (protein losing enteropathy, diarrhoea, and hypernatraemic dehydration). An initial diagnosis of necrotising enterocolitis was common, but no infant showed intestinal or hepatic portal haemorrhage. The gestational age of the infants was 24–38 weeks. All had received fresh maternal breast milk. It is suggested that CMV enteritis is added to the spectrum of clinical manifestations of postnatal CMV infection. Signs suggestive of necrotising enterocolitis with atypical features should prompt investigations for CMV infection.

Postnatal cytomegalovirus (CMV) infection has not attracted the attention received by congenital infection as it is held to result in low morbidity. The recognition of severe gastrointestinal symptoms in a small number of preterm infants with postnatal CMV infection prompted a search of the literature and a review of infants admitted to our neonatal unit over a five year period who were known to have postnatally acquired CMV infection.

METHODS

Infants were identified as a result of CMV screening in the presence of symptoms such as prolonged jaundice, unexplained enteritis, or signs of systemic sepsis in the absence of positive blood cultures for bacterial sepsis. The standard screen test was the urine CMV DEAFF (detection of early antigen fluorescent foci) test. This is a rapid culture test whereby early CMV antigens are detected by immunofluorescence after the inoculum on a cell sheet is spun for two to four days. The diagnosis of CMV infection was based on the fact that a previously negative urine CMV DEAFF test became positive coincidentally with the onset of clinical symptoms. Urine CMV DEAFF screening of infants at admission was not performed routinely although many of the infants had early screening. However, all of our cases had negative urine CMV DEAFF tests before the positive tests. In cases where the symptomatology was severe, additional tests such as CMV blood IgM and antigen detection were carried out. As the infants were sick preterm babies, it was not feasible to obtain gastrointestinal biopsy samples to confirm the diagnosis of CMV gastrointestinal disease.

Our unit feeding policy is to start enteral feeds early once the infant is clinically stable. The preference is for the mother’s own breast milk. If not available, then with parental consent, banked breast milk is given. The mother’s own milk is given fresh within 48 hours of expression. This milk would have been stored at 4°C. If fresh mother’s milk is unavailable or insufficient, frozen maternal breast milk or banked donor milk is used. Our unit has its own milk bank on site. Milk is obtained from donors who are screened for a panel of viruses (hepatitis B, hepatitis C, HIV-1 and HIV-2, human T cell leukaemia/lymphoma virus type 1 or II, and syphilis.). The milk is pasteurised (heated to 62.5°C for 30 minutes) and then frozen at −20°C.

RESULTS

Sixteen infants were identified out of 2830 admissions to the Queen Charlotte’s and Chelsea and Hammersmith Hospital neonatal unit over a five year period. They ranged from a median age of 25 weeks (24–29) and median birth weight of 801 g (705–978) (values in parentheses are interquartile ranges). All infants had received fresh maternal expressed breast milk before the onset of symptoms, and, at the time of diagnosis, were receiving a combination of fresh maternal expressed breast milk and banked breast milk. Maternal CMV serostatus was not available for all infants.

Table 1 shows basic data and clinical characteristics of the cases. The infant in case 2 received ganciclovir from day 32 until 44. There was no appreciable improvement of his clinical status that could be attributed to the antiviral treatment. His gastrointestinal symptoms subsequently resolved and he was eventually discharged home. He now has developmental delay.

Table 2 shows the identified cases as a percentage of admissions in gestational age groups. Two of the infants died (cases 5 and 11).

DISCUSSION

During the past decade, developments in the diagnosis and management of CMV infection and disease have led to refinements in the definitions of “infection” and “disease” in an attempt to develop consistent reporting in clinical trials. “CMV infection” is defined as the isolation of the CMV virus or detection of viral proteins or nucleic acid in any body fluid or tissue specimen. “CMV disease”, or more specifically in the case of “CMV gastrointestinal disease”, the definition encompasses clinical symptoms from the upper and lower gastrointestinal tract, findings of macroscopic mucosal lesions on endoscopy, and demonstration of CMV infection in a gastrointestinal tract biopsy specimen. These definitions were developed primarily for the transplant population and may not be applicable to a newborn population. In particular, it is not a feasible option to perform endoscopy and obtain biopsy specimens on premature newborns without an open

Abbreviations: CMV, cytomegalovirus; DEAFF, detection of early antigen fluorescent foci; NEC, necrotising enterocolitis; PCR, polymerase chain reaction
liver, and subsequently died. In four infants, the initial diagnosis was that of necrotising enterocolitis (NEC) because of sepsis in the absence of positive bacterial cultures. Seven of the 16 babies had severe diarrhoea during the time of CMV infection. Of these, one had a severe protein losing enteropathy and one developed severe acute hypernatraemic dehydration and subsequently died. In four infants, the initial diagnosis was that of necrotising enterocolitis (NEC) because of the presence of abdominal distension, tenderness, and fresh blood staining of the stools. However, no infant had radiological evidence of intramural or hepatic portal pneumatosis, and in none of these cases was NEC subsequently confirmed.

CMV enteritis with ischaemic necrosis is common and well recognised in patients with AIDS and in immunosuppressed transplant patients. The spectrum of gastrointestinal involvement in this patient group includes oesophageal, gastric, small intestinal, and colonic disease as well as cholangitis and pancreatitis. Clinical manifestations include abdominal pain, watery diarrhoea, gastrointestinal bleeding, vomiting, and protein losing enteropathy. Small bowel perforation may occur. Dolgin et al12 described a case of CMV enteritis in a child with AIDS resulting in massive haemorrhage and fatal small bowel obstruction. At laparotomy, large yellow plaques with central ulceration were found along the entire small bowel. Smith et al13 reported on two infants with extensive leio-myolysis due to CMV enterocolitis in association with HIV infection.

We identified only a single published report of CMV enteritis in the newborn. This described a 2.2 kg neonate who presented with a diagnosis of NEC. These authors refer to four additional reports of neonatal CMV enteritis in the non-English language literature.

The preterm neonate is an immunocompromised host, and therefore the common observation of clinically significant

### Table 1: Details of the 16 cases of cytomegalovirus (CMV) infection identified over a five year period

<table>
<thead>
<tr>
<th>Case</th>
<th>Sex</th>
<th>GA (weeks)</th>
<th>BW (grams)</th>
<th>Neg test</th>
<th>CMV test</th>
<th>Other positive tests</th>
<th>Symptom onset</th>
<th>Symptoms</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>F</td>
<td>29</td>
<td>890</td>
<td>3</td>
<td>65</td>
<td>–</td>
<td>64</td>
<td>Diarrhoea, NEC</td>
</tr>
<tr>
<td>2</td>
<td>M</td>
<td>24</td>
<td>550</td>
<td>10</td>
<td>27</td>
<td>ETA (27)</td>
<td>25</td>
<td>Diarrhoea, abdominal distension, protein losing enteropathy, pleural effusions, treated with ganciclovir</td>
</tr>
<tr>
<td>3</td>
<td>F</td>
<td>25</td>
<td>730</td>
<td>2</td>
<td>37</td>
<td>Blood IgM (37)</td>
<td>37</td>
<td>No abdominal symptoms, lethargy, thrombocytopenia</td>
</tr>
<tr>
<td>4</td>
<td>M</td>
<td>27</td>
<td>1010</td>
<td>3</td>
<td>61</td>
<td>–</td>
<td>56</td>
<td>No abdominal symptoms</td>
</tr>
<tr>
<td>5</td>
<td>M</td>
<td>25</td>
<td>908</td>
<td>1</td>
<td>35</td>
<td>–</td>
<td>30</td>
<td>No abdominal symptoms</td>
</tr>
<tr>
<td>6</td>
<td>M</td>
<td>29</td>
<td>1485</td>
<td>2</td>
<td>61</td>
<td>–</td>
<td>60</td>
<td>No abdominal symptoms</td>
</tr>
<tr>
<td>7</td>
<td>M</td>
<td>29</td>
<td>1130</td>
<td>2</td>
<td>48</td>
<td>–</td>
<td>45</td>
<td>No abdominal symptoms</td>
</tr>
<tr>
<td>8</td>
<td>M</td>
<td>32</td>
<td>978</td>
<td>3</td>
<td>80</td>
<td>–</td>
<td>70</td>
<td>Diarrhoea</td>
</tr>
<tr>
<td>9</td>
<td>F</td>
<td>25</td>
<td>642</td>
<td>2</td>
<td>56</td>
<td>ETA (80)</td>
<td>40</td>
<td>NEC, died</td>
</tr>
<tr>
<td>10</td>
<td>F</td>
<td>24</td>
<td>715</td>
<td>55</td>
<td>66</td>
<td>ETA (65)</td>
<td>60</td>
<td>Diarrhoea</td>
</tr>
<tr>
<td>11</td>
<td>F</td>
<td>24</td>
<td>801</td>
<td>1</td>
<td>67</td>
<td>–</td>
<td>60</td>
<td>Severe GOR requiring fundoplication, abdominal LFT</td>
</tr>
<tr>
<td>12</td>
<td>M</td>
<td>24</td>
<td>775</td>
<td>1</td>
<td>105</td>
<td>–</td>
<td>103</td>
<td>No abdominal symptoms, lethargy</td>
</tr>
<tr>
<td>13</td>
<td>M</td>
<td>25</td>
<td>908</td>
<td>1</td>
<td>45</td>
<td>–</td>
<td>42</td>
<td>No abdominal symptoms, lethargy</td>
</tr>
<tr>
<td>14</td>
<td>M</td>
<td>24</td>
<td>705</td>
<td>5</td>
<td>66</td>
<td>–</td>
<td>66</td>
<td>No abdominal symptoms, lethargy</td>
</tr>
<tr>
<td>15</td>
<td>F</td>
<td>24</td>
<td>801</td>
<td>1</td>
<td>67</td>
<td>–</td>
<td>60</td>
<td>Severe GOR requiring fundoplication, abdominal LFT</td>
</tr>
<tr>
<td>16</td>
<td>F</td>
<td>24</td>
<td>801</td>
<td>1</td>
<td>67</td>
<td>–</td>
<td>60</td>
<td>Severe GOR requiring fundoplication, abdominal LFT</td>
</tr>
</tbody>
</table>

F, Female; M, Male; GA, gestational age in completed weeks; BW, birth weight in grams; Neg test, last negative urine CMV DEAFF test; CMV test, CMV urine DEAFF test (day of first positive test); DEAFF, detection of early antigen fluorescent; Other positive tests, other CMV tests (day of first positive test); ETA, endotracheal aspirate; Symptom onset, day of symptom onset; LFT, liver function tests; NEC, necrotising enterocolitis; GOR, Gastro-oesophageal reflux.

### Table 2: Identified cases as a percentage of admissions in gestational age groups

<table>
<thead>
<tr>
<th>Gestational age (weeks)</th>
<th>No cases</th>
<th>Percentage of total admissions for gestational age group</th>
</tr>
</thead>
<tbody>
<tr>
<td>23–27</td>
<td>10</td>
<td>3</td>
</tr>
<tr>
<td>27–31</td>
<td>4</td>
<td>1</td>
</tr>
<tr>
<td>31–35</td>
<td>1</td>
<td>0.2</td>
</tr>
<tr>
<td>&gt;35</td>
<td>1</td>
<td>0.1</td>
</tr>
</tbody>
</table>

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gastrointestinal signs during postnatal CMV infection, probably acquired enterally in fresh maternal breast milk, is not surprising. This does not, however, appear to be a recognised manifestation of postnatal CMV infection. Diagnosis is likely to be difficult, and recognition requires a high index of suspicion. Kusne et al\textsuperscript{11} showed that the specificity and sensitivity of a polymerase chain reaction (PCR) technique was better than diagnosis of CMV enteritis based on histopathology, shell viral assay, and tube culture of intestinal biopsy specimens. In any case, intestinal biopsy, other than at laparotomy, is unlikely to be widely applicable in the preterm population. However, it might be of interest to evaluate intestinal biopsy specimens taken at surgery in infants with NEC to see if some have CMV enteritis. There has been a report of the value of negative CMV PCR from stool specimens taken from immunocompromised adults in ruling out intestinal CMV disease.\textsuperscript{15} The authors did caution that this method required further evaluation with testing of larger numbers of patients with AIDS. Stool testing was unfortunately not performed in our patients, and, so far, is not advocated routinely as a confirmatory test of CMV in the neonatal population.\textsuperscript{14}

The examination of maternal CMV serological status to assess the risk of CMV transmission to the infant is controversial. Current practice is to recommend and promote the use of mother's own fresh milk. Although studies have shown that freezing breast milk at \( -20^\circ\)C reduces CMV viral titres\textsuperscript{27} and infectivity,\textsuperscript{28} there is the disadvantage of delay in introduction of enteral feeds. A recent report describes a low prevalence of postnatal CMV acquisition in a neonatal unit where it was policy to freeze breast milk before administration to the infant.\textsuperscript{29} The minimum duration of freezing was not included in the report. There is a balance between the risk of CMV acquisition from fresh breast milk and the benefits of early introduction of enteral feeds. Early initiation of enteral feeds has been shown to reduce the total duration of parenteral nutrition, promote gastrointestinal development, and improve overall tolerance of feeds.\textsuperscript{29–32} Clarification of best practice with regard to maternal CMV testing and handling of expressed breast milk will require prospective evaluation.

All our infants except one were premature, with a skewed range. The exception was a 38 week female infant with normal intrauterine growth. In the preterm population, however, it might be of interest to identify infants with NEC to see if some have CMV enteritis. There is a balance between the risk of CMV acquisition from fresh breast milk and the benefits of early introduction of enteral feeds. Early initiation of enteral feeds has been shown to reduce the total duration of parenteral nutrition, promote gastrointestinal development, and improve overall tolerance of feeds.\textsuperscript{29–32} Clarification of best practice with regard to maternal CMV testing and handling of expressed breast milk will require prospective evaluation.

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