Screening for cystic fibrosis by examination of meconium

R. PROSSER, H. OWEN, F. BULL, B. PARRY, J. SMERKINICH, H. A. GOODWIN, and J. DATHAN

From the Department of Paediatrics and Pathology, The Royal Gwent Hospital, Nevill Hall, Monmouthshire, and City General Hospital, Stoke-on-Trent

Prosser, R., Owen, H., Bull, F., Parry, B., Smerkinich, J., Goodwin, H. A., and Dathan, J. (1974). Archives of Disease in Childhood, 49, 597. Screening for cystic fibrosis by examination of meconium. The value of detecting albumin in meconium as a screening procedure for cystic fibrosis (CF) has been assessed on 34,228 samples in South Wales and North Staffordshire over a 4-year period; simultaneously, four methods of detecting albumin were evaluated. 12 cases of CF were detected, detection rate being 60%. The incidence of the disease in the population screened was 1 in 1850, confirmed by clinical and other test procedures. Cases of CF without impairment of pancreatic function are likely to be missed by screening methods which depend on the presence of albumin in meconium.

Green, Clarke, and Shwachman (1958) showed that in comparison with the normal, the meconium in meconium ileus contains a large amount of protein, predominantly albumin. This excess albumin could also be detected in the apparently normal-looking meconium of babies who are born affected by CF without meconium ileus (Green and Shwachman, 1968; Wiser and Beier, 1964). The albumin is probably derived from liquor amnii swallowed by the fetus and not digested because of impairment of the exocrine function of the pancreas (Pritchard 1965, 1966). Several authors have suggested that the presence of albumin in meconium had potential in screening for CF (Schutt and Isles, 1968; Green and Shwachman, 1968; Wiser and Beier, 1964).

A screening programme to record the presence of albumin in meconium of all newborns was initiated on 1 January 1970 in the Royal Gwent Hospital, and was extended to include all births in Monmouthshire, and later in East Glamorgan and North Staffordshire. The aim of the survey was to assess the value of detecting albumin in meconium as a screening test and to determine the most suitable test, method of recording results, and cost.

Methods

Part of the first or second specimen of meconium was collected into a sterile disposable container and sent to the laboratory suitably labelled. Conventional hospital transport was used, or for district deliveries and distant hospitals specimens were refrigerated at 4 °C and sent in weekly batches through the post to the laboratory. The tests were performed in either the Paediatric or Pathology Departments of the Royal Gwent Hospital, Newport; the City General Hospital, Stoke-on-Trent; Nevill Hall Hospital, Abergavenny; East Glamorgan Hospital, Pontypridd; and Cardiff Maternity Hospital, Cardiff.

Four techniques were assessed, and frequently more than one type of test was used on a particular specimen of meconium (Table I). At the outset the trichloracetic acid (TCA) ring test (Green and Shwachman, 1968) was used. However, because of difficulty in recognizing the end point, it was soon abandoned in favour of the sulphosalicylic acid test (SSA), the predominant technique in the survey. All the specimens which were tested by Albustix and BM meconium test also had an SSA test done, and all positive cases of immunodiffusion also had an SSA test done.

SSA test. Approximately 1 g meconium was diluted 1 in 8 with saline and shaken mechanically for 30 minutes at 100 oscillations/min; 5 ml of homogenate was poured into plastic tubes and centrifuged at 3000 r.p.m. for 15 minutes. Into duplicate 10 x 75 mm plastic tubes 1.5 ml saline was dispensed, and to each tube 0.5 ml of the meconium supernatant was added, giving an initial testing dilution of 1 in 32. The dilution was thoroughly mixed with a vortex mixer and 0.2 ml of 25%, sulphosalicylic acid and added to one of each of the duplicate...
TABLE I
Number of specimens of meconium tested by each method

<table>
<thead>
<tr>
<th>Method</th>
<th>Trichloracetic acid (TCA)</th>
<th>Sulphosalicylic acid (SSA)</th>
<th>Immunodiffusion</th>
<th>Albustix</th>
<th>Boehringer (BM)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Trichloracetic acid (TCA)</td>
<td>330</td>
<td>29,984</td>
<td>4535</td>
<td>20,087</td>
<td>2106</td>
</tr>
</tbody>
</table>

Results

There have been 40,801 births included in the survey area and 34,228 specimens of meconium have been tested, i.e. 84% of all possible births. 22 cases of CF have been diagnosed among the 40,801 births to date, an incidence of about 1 in 1850. 12 of these were detected by meconium screening. 8 were negative when the meconium was tested for albumin, and the remaining 2 were among the group from which no meconium was collected (Table II).

SSA test (Table III). A titre of 1/128 or higher appeared to be significant for recognizing CF and was considered as a positive result. The 8 cases of CF missed by the test gave negative results at 1 in 8 dilution. 2 of these infants were sibs of known cases and were therefore anticipated as possible cases of CF. Their meconium was negative to SSA and Albustix and in one instance to immunodiffusion test.

Immunodiffusion test (Table IV). To date no case of CF has been detected in the survey by this

TABLE II
Overall results of meconium testing

<table>
<thead>
<tr>
<th></th>
<th>Known cases of CF</th>
<th>Meconium not tested</th>
<th>Detected by examination of meconium</th>
<th>Not detected by examination of meconium</th>
<th>% detected by test</th>
</tr>
</thead>
<tbody>
<tr>
<td>South Wales</td>
<td>16</td>
<td>2</td>
<td>9*</td>
<td>5</td>
<td>64</td>
</tr>
<tr>
<td>North Staffordshire</td>
<td>6</td>
<td>0</td>
<td>3</td>
<td>3</td>
<td>50</td>
</tr>
<tr>
<td>Total</td>
<td>22</td>
<td>2</td>
<td>12</td>
<td>8</td>
<td>60</td>
</tr>
</tbody>
</table>

*1 case of meconium ileus.

TABLE III
Results of SSA test

<table>
<thead>
<tr>
<th>No. of births</th>
<th>No. not tested</th>
<th>Infants tested—serial dilutions</th>
<th>% of false positives</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>SSA only</td>
<td>SSA + electrophoresis*</td>
</tr>
<tr>
<td>35,688</td>
<td>5702 (2)</td>
<td>26620 (7)</td>
<td>508</td>
</tr>
</tbody>
</table>

Numbers in parenthesis denote number of infants with CF.

*Meconium with a titre of ≥1/128 was subjected to electrophoresis.
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TABLE IV
Results of immunodiffusion test

<table>
<thead>
<tr>
<th>No. of births</th>
<th>No. not tested</th>
<th>Negative (&lt;20 mg/100 ml)</th>
<th>Positive (&gt;20 mg/100 ml)</th>
<th>% false positives</th>
</tr>
</thead>
<tbody>
<tr>
<td>5111</td>
<td>576*</td>
<td>4525*</td>
<td>10</td>
<td>0.2</td>
</tr>
</tbody>
</table>

*One infant with CF.

TABLE V
Results of Albustix test

<table>
<thead>
<tr>
<th>No. of births</th>
<th>No. not tested</th>
<th>Infants tested</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Negative 1/64</td>
</tr>
<tr>
<td>20788</td>
<td>2250</td>
<td>18304 (3)</td>
</tr>
</tbody>
</table>

Numbers in parentheses denote number of infants with CF.

test; 1 case gave a negative result, i.e. <20 mg/100 ml albumin in a 1/5 meconium extract. It was also negative to SSA. 3 of the positives by this immunodiffusion technique were also positive for occult blood.

**Albustix** (Table V). The results were effectively the same as with the SSA test. Because of the green colour of the supernatant fluid (biliverdin), the interpretation of positive results was difficult. 5 cases of CF were detected using this technique, and 4 gave negative results. The 5 positive cases were also detected by the SSA test and the 4 negative cases were missed by the SSA test.

**BM test** (Table VI). 1 case of CF was detected and confirmed by this test. The number of false positives was small.

**Electrophoresis** (Table VII). Protein electrophoresis is now performed routinely at the Royal Gwent Hospital on meconiums with a negative occult blood but with a positive SSA titre of 1/128.

TABLE VI
Results of BM meconium test

<table>
<thead>
<tr>
<th>No. tested</th>
<th>No. negative</th>
<th>No. positive</th>
<th>% false positives</th>
</tr>
</thead>
<tbody>
<tr>
<td>2229</td>
<td>2216</td>
<td>13*</td>
<td>0.6</td>
</tr>
</tbody>
</table>

*One infant with CF.

This readily distinguishes between false and true positives; the latter are recognized by a dense albumin band. Only 8 of the 141 with false positive titres at 1/128 showed albumin; in 6 of these the albumin was present in a small amount only. The 2 exceptions with a dense albumin band included a small premature infant weighing 1000 g who has subsequently thrived, and 1 normal infant who also gave repeatedly abnormal sweat chlorides in the first 2 weeks of life, with a gradual return to normal levels.

**Discussion**

Several tests have been used to assess the value of examining meconium for albumin as a screening test.
for CF. No one test appears to be more efficient than the other at detecting albumin. The end point is more difficult to recognize in the SSA test and with Albustix. Both these tests, together with the BM meconium test, because they are not albumin specific, gave slightly more false positives than the immunodiffusion test.

The number of false positives with the SSA test (0.7%), Albustix test (0.3%), and BM meconium test (0.6%) can be reduced to a negligible number (0.02%) if the positive reacting meconiums are subjected to electrophoresis, when false positives are excluded by the absence of an albumin band. The source of protein in false positives is not known, but the presence of blood is easily excluded by the occult blood test.

An SSA or Albustix titre of 1/128 or higher is considered positive and suggestive of CF. Albustix can be used more directly as a ward test (Cain, Deall, and Noble, 1972), and Hobbs (1969) described a method for using Albustix which is both direct and, by providing its own control, simplifies the recognition of positives. The BM meconium test can be performed even by a relatively unskilled operator, the blue end point being easy to recognize. The time of the reading is not critical and the strip can be retained as a permanent record in the infant's notes. The immunodiffusion test gives a more definite end point and also the opportunity to detect the exact amount of protein present. It is sensitive, albumin specific, and there are fewer false positives (0.2%). All these tests were assessed at various times on known positives and were found to give more or less identical results.

False negatives, i.e. patients subsequently shown to have CF, have occurred with three of the tests and are likely in all four. No case of CF has been detected by one and missed by the other. Of the 6 cases not identified by this test, 3 were found to have normal excretion (<2 g/day). Fat excretion of the remaining 3 cases is not known, but one had a stool trypsin of 1 in 2000. This suggests that pancreatic function in these children is not severely disturbed, and would account for the negative meconium tests.

The detection rate in this survey, 60%, is low compared to the 90% success described by Green and Shwachman (1968). However, their survey included only 196 infants over a 9-year period, all sibs of known cases of CF.

Approximately 10 to 15% of infants with CF have normal pancreatic function, and the remainder can be expected to give a positive meconium albumin test. Possibly the relatively low detection rate in this survey was due to the chance occurrence of an abnormally large number of infants with adequate pancreatic function. Green and Shwachman (1968) suggested that children with false positive results may represent heterozygotes, but this was not substantiated by us.

The protein appears stable, and transport of nonrefrigerated specimens through the post does not affect positive results. 6 specimens of meconium, which included known positives and false positives, were retested after 18 months and were found to give identical results with the SSA and Albustix tests and by electrophoresis.

Cost is important (Table VIII) and in this respect Albustix has a clear advantage if used as a ward test, but it is the least satisfactory of the four tests. When performed in the laboratory the cost is similar in the SSA and immunodiffusion tests and with Albustix, and the overall cost is only slightly increased if false positives (1 in 128 or greater) are subjected to electrophoresis. The BM meconium test is significantly more expensive, either as a laboratory or a ward test.

We conclude that the examination of meconium protein for CF is not an ideal screening test because of the large number of false negative results. However, because of its relative simplicity, low cost, absence of direct involvement of the parent and patient, and high yield of positive cases, its use on a wider scale is justified. Any future screening programme should also be used to assess the value of early prophylactic antibiotic therapy. On balance, there is little to choose between the tests, but the

<table>
<thead>
<tr>
<th>SSA</th>
<th>SSA + electrophoresis</th>
<th>Immunodiffusion</th>
<th>Albustix</th>
<th>BM</th>
</tr>
</thead>
<tbody>
<tr>
<td>8.35p</td>
<td>8.44p</td>
<td>8.85p</td>
<td>7.96p (0.4p)*</td>
<td>12p (10p)*</td>
</tr>
</tbody>
</table>

*Cost if used as a ward test.

Note: Costs have been calculated on the basis that the test will be done in the laboratory by a technician and the results properly recorded. Albustix (Schutt and Isles, 1968; Cain et al., 1972) and BM test are suitable for ward use and the cost could be further reduced if the material is tested directly from the baby's napkin.
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immunodiffusion, or the SSA test plus electrophoresis, are the most satisfactory laboratory techniques, while the BM meconium test, though expensive, is preferred as a ward test because of its more easily recognized end point. We advocate that all specimens should be tested in a laboratory where quality control can be assured and results properly recorded and notified by an appropriate secretariat.

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References


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