The impact of meconium ileus on the clinical course of children with cystic fibrosis


ABSTRACT: The present study was designed to compare the clinical course of children diagnosed with cystic fibrosis (CF) in infancy due to the presence of meconium ileus (MI) with children diagnosed by way of a newborn screening programme (non-MI).

A matched case-control study design was used. Matching was performed on the basis of sex and date of birth. All children born in New South Wales, Australia after 1980 and who had attended the CF clinic at The Children’s Hospital at Westmead since diagnosis were included as possible cases or controls. Parameters pertaining to the clinical course were compared in 39 matched pairs.

MI children had a significantly worse pulmonary status. The forced expiratory volume in one second was 16.3±5.2% higher (p<0.001, n=21 pairs) and the forced vital capacity value 10.5±4.7% higher (p=0.05, n=21 pairs) in non-MI children. The difference between the pairs (18.6±4.4 MI and 20.5±3.4 non-MI) in the Shwachman chest radiograph score was statistically significant (p<0.05, n=39 pairs). There were no significant differences in any other assessed parameters, such as height, weight, the presence of liver function abnormalities, the frequency of hospitalization or airway microbial colonization.

Meconium ileus may be an early indication of a more severe phenotype of cystic fibrosis. This was suggested by the significantly lower pulmonary function found in children with a history of meconium ileus compared to age- and sex-matched children who did not have meconium ileus.

Diagnosis of cystic fibrosis (CF) is generally made on the basis of either the presence of meconium ileus (MI) in newborns, a positive newborn screening test, suspicion raised by a family history of CF, failure to thrive, or the presence of respiratory and/or gastrointestinal symptoms in childhood [1]. The mode of presentation has been shown to be a significant prognostic factor for survival of patients with CF, as patients presenting with predominantly gastrointestinal symptoms (presumably symptoms such as pancreatitis, steatorrhoea or rectal prolapse) have higher survival rates than those presenting with respiratory symptoms or MI [2].

MI is the presenting feature in ~18% of Australian children with CF, and this incidence is similar to that reported in other countries [3]. There appears to be a tendency for MI to occur in subsequent siblings with CF when the first child presents with MI [3]. Linkage studies indicate that predisposition to MI may be explained by genetic differences unassociated with mutations of the cystic fibrosis transmembrane regulator (CFTR) gene [4]. Adoption of novel surgical procedures over the last 30 yrs, such as the Bishop-Koop ileostomy, has resulted in a dramatic decrease in the postoperative mortality associated with MI, from >60% in the 1960s to ~0% in the 1990s [5, 6].

It has been suggested that infants with MI who survive the neonatal period have a long-term survival that is equivalent to children presenting with other symptoms [2]. The earlier diagnosis of CF in these infants has been suggested as conferring positive prognostic benefits [5]. Moreover, these children have normal lung function at the time of diagnosis [7], and it has been reported that children presenting with MI have less rapid progress of lung disease than children presenting with respiratory symptoms [2]. The absence of long-term effects of MI on nutritional, pulmonary, gastrointestinal and hepatobiliary status has been reported in a number of studies, although some investigators have shown an increase in hepatic abnormalities in patients with a history of MI [8–11].

Therefore, there is evidence that the presence of MI in the newborn period can influence the clinical course of CF. The present authors surmised that this influence may be negative. The absence of control CF groups composed of asymptomatic children also diagnosed in infancy, might have obscured the true long-term effects of MI. This has been reinforced by a recent study, demonstrating a clear association between MI and malnutrition when children diagnosed via newborn screening were used as the control population [12]. The purpose of the present study, which employed a matched case-control study design, was to compare the clinical course of children with CF.
presenting with MI to the clinical course of children with no history of MI, diagnosed in infancy via newborn screening.

Material and methods

Study subjects and study design

This study was approved by the Human Ethics Committee of the Royal Alexandra Hospital for Children, Westmead, New South Wales (NSW), Australia. The medical records of all children attending the CF clinic at this hospital were scrutinized to identify those that had undergone an annual check in 1999. These children were then further limited to those born in NSW between January 1, 1982–January 1, 1998, who had been followed by this clinic from their birth and who had been diagnosed with CF on the basis of MI (designated as an MI child). Each MI child (index child) was paired with a child of the same sex and with a birth date in the same month or 1–2 months before or after the index child’s birth date. The paired child (non-MI child) had also undergone an annual check in 1999, was also born in NSW during the same period, and had also been followed by the same clinic since diagnosis; however, the paired child was diagnosed as having CF as a result of newborn screening. Prior to 1994, the newborn screening test consisted of detection of a raised concentration of immunoreactive trypsinogen in a heel-prick blood spot, whereas the test used since 1994 detects the presence of the ΔF508 gene mutation. These newborn screening tests were performed on each child born in NSW since August 1981. A confirmatory diagnostic sweat test (Gibson-Cooke) was performed on all children detected via newborn screening, who did not have two copies of the ΔF508 gene mutation.

Methods

Information from the 1999 Annual CF Review, as well as the medical record, was collected for each child. Annual reviews were performed close to the child’s birthday and included a physical (respiratory and gastrointestinal) examination as well as an assessment of nutritional, biochemical (liver function tests) and haematological status, sputum cultures and physiotherapy refinement. The physical examination included recording the child’s height and weight, as well as the presence of hepatomegaly, splenomegaly, or digital clubbing. Nutritional parameters included serum vitamin A and E levels, vitamin supplementation, pancreatic enzyme replacement therapy and enteral feeding via gastrostomy. The number and duration of hospitalizations and intravenous antibiotic therapy were recorded, as were any episodes of distal intestinal obstruction syndrome and allergic bronchopulmonary aspergillosis. Spirometry was performed on all children aged >5 yrs and assessment of lung volumes (by plethysmography) was carried out in children >8 yrs. The Shwachman scores [13] for general activity, physical examination, nutrition and chest radiograph, were also recorded.

Height and weight were converted into percentiles, and z-scores (sd scores) and body mass index were calculated. If clubbing was present, the degree of clubbing was recorded using a scale from 1–3. In this scale, loss of the angle between the nail bed and the cuticle was awarded a score of 1, convex nails and an increase in the bulk of the terminal pulp was awarded a score of 2, and a score of 3 was awarded in cases where clubbing had progressed to the point where there was an increase in nail curvature and widening of the phalanges. Pancreatic enzyme replacement is expressed as the number of lipase units consumed per kg of body weight per day. Spirometric values (forced expiratory volume in one second (FEV1), forced vital capacity (FVC) and FEV1/FVC) and lung volumes are expressed as percentage predicted using the normal values published by Polgar and Promadhat [14]. Where known, the genotype was classified as ΔF508 homozygous, ΔF508 heterozygous or ΔF508 null.

Analysis

As this was a matched case-control study, paired data analysis was performed. Paired t-tests were used for parametric data and the McNemar’s test was used for nonparametric variables. A subgroup analysis was performed for the length of stay (in days) and the number of admissions for those patients admitted to hospital, for any reason, in the 12 months prior to their annual review. For this subgroup analysis, comparison was made on a group basis between MI and non-MI patients, using unpaired t-tests. All reported p-values are two-tailed and statistical significance was ascribed to p-values <0.05.

Results

In 1999, 341 patients were seen for an annual review in the CF clinic. Of these, 294 had been born between January 1, 1982–January 1, 1998, and 253 of them had been born in NSW. Of this cohort of 253 children, 39 (22 male) had been diagnosed with CF due to the presence of MI on the first or second day of life. The matched non-MI patients identified via the newborn screening programme were diagnosed by a positive sweat test at an average age of 60.6±9.5 days.

In the MI group, eight children had deceased or living siblings with CF. Two of the children were fraternal twins who both presented with MI. Two children had one younger CF sibling without MI, one child had two younger CF siblings without MI and one child had one older and one younger CF sibling who did not present with MI. One child had a deceased older CF sibling who had not presented with MI and another had a deceased younger CF sibling who had presented with MI. Only in two instances did both or all CF siblings of a family present with MI as newborns.

As expected for the study design, the MI and non-MI patients were well matched for age, as the mean
Data are presented as mean±SEM. BMI: body mass index; MI: meconium ileus. *: p<0.05, the mean difference between MI patients and matched non-MI patients is statistically significant using paired t-test, n=39 pairs in all cases.

difference in age between the pairs was only 6 days (p>0.05, n=39, paired t-test). The diagnostic sweat test results were not significantly different (sweat chloride: 101±4 mmol·L⁻¹ for MI patients; 100±3 mmol·L⁻¹ for non-MI patients, p>0.05, n=26 pairs); however, in 13 pairs of children, one or both of the children had not been sweat tested. Analysis of the children’s geno-type indicated no statistically significant difference in ΔF508 homozygosity between children presenting with and without MI. There were 19 pairs of children who were ΔF508 homozygous: nine pairs in which only the child with MI was ΔF508 homozygous, four pairs in which only the non-MI child was ΔF508 homozygous, and six pairs in which neither child was ΔF508 homozygous (p>0.05, n=38, McNemar’s test).

A similar absence of significant differences was seen in the analysis of ΔF508 heterozygosity and cases in which there was no copy of the ΔF508 mutation.

The nutritional status of the children was not affected by a history of MI (table 1). The MI children were significantly taller in terms of both height percentile (14.77% taller) and height z-scores (0.572 greater). The equivalence of the children’s nutritional status was also demonstrated by the absence of any significant difference in the nutrition component of the Shwachman score (fig. 1). All the children with MI were pancreatic insufficient and all but two of the matched controls were pancreatic insufficient, this difference being of no statistical significance (p>0.05, McNemar’s). The use of supplemental feeding via gastrostomy was equally distributed in children with and without a history of MI. The children with MI did not require significantly higher levels of pancreatic enzyme replacement therapy than their matched pairs in order to attain and maintain this equivalent nutritional status (mean±SEM 4,873±226 and 4,241±385 lipase units·kg body weight⁻¹·day⁻¹ respectively, p>0.05, t-test).

A history of MI did not predispose the children to many of the recognized complications of CF. There was no significant difference in the incidence of distal intestinal obstruction syndrome, hepatomegaly, splenomegaly, CF-related diabetes mellitus or allergic broncho-pulmonary aspergillosis (all p>0.05, McNemar’s, n=39) in children with MI as newborns and their matched pairs. In addition, there were no differences in plasma vitamin A, vitamin E, and nonfasting glucose levels or liver function tests (table 2). The mortality rate during 1999 was no different for children with or without a history of MI. One child died as a result of respiratory failure during 1999; the patient presented with MI as a newborn and was 16.5 yrs old at the time of their death.

Despite having a nutritional status equivalent to their matched pairs, the MI children had a significantly worse pulmonary status. The results of spirometric testing (fig. 2) showed a lower FEV₁ (71.32±3.86% and 87.58±4.77% pred, MI and non-MI respectively, n=21, p<0.001, paired t-test), a lower FVC (86.03±3.55% and 96.55±3.40% pred, MI and

| Table 1. – Physical characteristics of patients at 1999 annual review |
|------------------|------------------|------------------|
| **MI group**     | **Non-MI group** | **Mean difference** |
| Age yrs          | 9.39±0.74        | 9.40±0.75        | -0.02±0.03 |
| Height percentile| 43.33±5.30       | 28.56±3.69       | 14.77±7.09*|
| Weight percentile| 40.79±4.67       | 35.79±4.28       | 5.00±6.58  |
| Height z-score   | -0.23±0.197      | -0.81±0.142      | 0.572±0.261*|
| Weight z-score   | -0.34±0.159      | -0.47±0.157      | 0.133±0.236|
| BMI              | 16.81±0.39       | 17.54±0.63       | -0.731±0.576|
| Shwachman score  | 85.13±1.79       | 89.49±1.33       | -4.36±2.37 |

Fig. 1.– Lung function in children with cystic fibrosis with and without a history of meconium ileus (MI). The mean values measured in 21 children with MI in the newborn period (MI, □) and their matched pairs with no MI (non-MI, ◇) are shown and the error bars represent the SEM values. FEV₁: forced expiratory volume in one second; FVC: forced vital capacity; % pred: percentage of predicted values. *: p<0.05; ***: p<0.001, as assessed by performing paired t-tests.

| Table 2. – Plasma biochemistry results at 1999 annual review |
|------------------|------------------|------------------|
| **MI group**     | **Non-MI group** | **n**            |
| Vitamin A μmol·L⁻¹| 1.2±0.1          | 1.2±0.1          | 39               |
| Vitamin E μmol·L⁻¹| 11.6±1.1         | 14.2±1.1         | 39               |
| Nonfasting glucose mmol·L⁻¹| 5.6±0.2 | 5.3±0.3 | 35               |
| ALPase U·L⁻¹     | 329±20           | 306±19           | 39               |
| Transaminase U·L⁻¹| 37±4             | 31±3             | 39               |
| Gamma glutamine transaminase U·L⁻¹| 22±4 | 21±3 | 39               |

Data are presented as mean±SEM. MI: meconium ileus; n: number of pairs. There were no statistically significant differences for any measured parameter.
non-MI respectively, n=21, p<0.05, paired t-test) and a lower FEV1/FVC ratio (74.58±2.72% and 80.58±1.49%, MI and non-MI respectively, n=21, p<0.05, paired t-test). The chest radiograph and the general activity components of the Shwachman score were also significantly lower in these children than that of their matched pairs (fig. 1), but the presence and degree of finger clubbing was not associated with MI in the newborn period. There were 21 pairs of children >8 yrs, who underwent lung volume assessment. While there was evidence of a greater residual volume in the MI child in each pair (181.1±15.7% and 161.4±11.2% pred MI and non-MI, respectively), the difference was not statistically significant (p=0.05, t-test, n=21). The total lung capacity of children with MI was not statistically different from that of non-MI children (105.9±6.0% and 106.8±2.2% pred MI and non-MI respectively, p=0.05, t-test, n=21).

Airway bacterial colonization patterns were not affected by a history of MI. Sputum cultures from six pairs of children grew normal respiratory flora. There were six instances where only the MI child had normal respiratory flora and another eight where only the non-MI child of the pair was free from colonization with MI. However, these differences were not statistically significant (p=0.06 and 0.07 respectively, paired t-tests, n=39). Subgroup analysis to compare length of stay and the number of admissions in children who had required an admission in the previous year did not reveal statistically significant differences related to a history of MI.

Table 3. – Sputum culture results for 1999 for children with and without a history of meconium ileus (MI) as newborns

<table>
<thead>
<tr>
<th>Culture</th>
<th>Both children</th>
<th>MI child only</th>
<th>Non-MI child only</th>
<th>Neither child in pair</th>
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<tbody>
<tr>
<td>Staphylococcus</td>
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<td>7</td>
<td>9</td>
<td>21</td>
</tr>
<tr>
<td>aureus</td>
<td>8</td>
<td>9</td>
<td>4</td>
<td>18</td>
</tr>
<tr>
<td>Pseudomonas</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td>37</td>
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<tr>
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<td></td>
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<td>Burkholderia</td>
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<td>cepacia</td>
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Discussion

This study confirms that MI in the newborn period does not affect the long-term nutritional status of children with CF. MI may, however, be an early indication of a more severe form of CF. This is suggested by the detection of significantly lower pulmonary status in children with a history of MI than in age- and sex-matched children who did not have MI as newborns.

The observation of a decreased pulmonary status (as measured by lung function testing and chest radiography) in the children presenting with MI in this study, supports the suggestion that MI may be an indicator of a more severe variant of CF [15]. Although the residual volumes were not significantly higher in the children with a history of MI, this may reflect the small number of pairs of children who were old enough to perform these manoeuvres. Nevertheless, the spirometric measures and chest radiograph scores were all lower in children who had MI. This finding is contrary to that reported in the study of FUCHS and LANGER [9], which showed no significant difference in respiratory status in patients who had presented with MI, and age- and sex-matched patients.

There are a number of reasons for the lack of consensus between the results of the present study and that of FUCHS and LANGER [9]. In the present study, the children with MI were matched with children who had also been diagnosed in infancy by way of a state-wide newborn screening programme. The matched children in the study of FUCHS and LANGER [9], however, were presumably diagnosed on the basis of symptomatology, as the mean age at diagnosis was 7.7 yrs. As it has been shown that the early detection of CF reduces morbidity [15, 16] and promotes superior nutritional status later in life [17], it is likely that the matched control children in the present study had benefited from early detection of their disease.

Abnormalities in pulmonary function have been reported in very young infants with CF diagnosed on the basis of clinical symptoms [7], but lung function appears to have been normal in asymptomatic infants [18]. Thus, it is conceivable that these early abnormalities partly explain why previous studies have shown that MI in the newborn period had no effect upon lung function later in life. Abnormalities in pulmonary function in the non-MI population used as controls, may have masked the true impact of MI.
This assertion is supported by the work of Waters et al. [19], who demonstrated superior pulmonary function at both 5 and 10 yrs of age in children diagnosed via newborn screening programme. By using matched controls also diagnosed in infancy, but by way of newborn screening before symptoms could develop, the present study has been able to eliminate these masking effects. The matched control children were diagnosed at approximately the same age as the children with MI, were all born and grew up in the same state, and have been treated within the same clinic using the same treatment strategies. By adopting this study design, some of the confounding variables have been eliminated, thus allowing the effects of MI on the clinical course of children with CF to be more accurately delineated.

It is possible that the impaired pulmonary status associated with a history of MI could be explained by the CF genotype of the patients. Relationships between the CF genotype and the presence of MI have been found. The G551D and R117H mutations are associated with a decreased incidence of MI, while there is a positive association between MI and the ΔF508 and G542X mutations [20–22]. Moreover, very young children, homozygous for the ΔF508 mutation and diagnosed via newborn screening, have been shown to have early evidence of airway obstruction when compared with children of other genotypes [23]. Despite this, studies of older children have shown that while poor pulmonary function may be related to pancreatic insufficiency [24], it is not influenced by the CF genotype [20, 21]. Therefore, the CF genotype is unlikely to explain the present results. No significant difference in pancreatic insufficiency or genotype was found between the matched pairs. There were, however, nine pairs in which the MI child alone was ΔF508 homozygous. Therefore, the genotype of the second mutation in the non-MI children may, in part, explain the differences in lung function. The limited population available for study and the disparity in second genotypes precludes firm conclusions in this regard.

Studies of sibling pairs with CF and identical CFTR genotypes have indicated the presence of modifier genes for MI on chromosome 19 [25]. Hence, it is possible that modifier genes could explain the presence of more severe pulmonary disease in children with a history of MI. Future studies may, therefore, affirm or eliminate a role for modifier genes in determining pulmonary status of patients with CF, with and without a history of MI.

Tepper et al. [7] showed that at the time of diagnosis, infants presenting with MI have a lung function that is indistinguishable from healthy control infants. Therefore, the relative decrement in pulmonary status associated with MI is not likely to have been a result of any in utero pulmonary abnormalities associated with the development of bowel obstruction in these children. This contention is supported by evidence of normal pulmonary anatomy and histology in children with a history of MI [26, 27]. The point at which the relative decrement in pulmonary status first becomes obvious is yet to be revealed by serial lung function studies in children with MI and matched controls. Because the present study included lung function assessments on children ≥5 yrs of age, the authors conclude that the decrement in lung function must develop sometime between infancy and 5 yrs of age.

The results of the present study suggest that meconium ileus is an indication of a more severe phenotype of cystic fibrosis, which may be conferred by the presence or absence of particular modifier genes. While nutrition is unaffected by meconium ileus later in life, adverse pulmonary status is related to the presence of this complication. Greater surveillance of, and early attention to, the pulmonary status of these patients may assist in preserving their lung function.

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References


