valve closure on ultrasound, and systolic anterior movement of the anterior mitral cusp. Hypertrophic cardiomyopathy is seen in infants of diabetic mothers, Pompe's disease, and Friedreich's ataxia, and there is often a strong familial tendency. Maron et al.5 reported 46 cases in childhood of asymmetric septal hypertrophy and noted the high incidence of sudden death (30%) especially in those with overt heart disease. They could not predict those at risk on the basis of echocardiogram, ECG, left ventricular function, or outflow obstruction. Beta-blockers or surgery in selected patients did not reduce mortality.

Rigid spine syndrome is of unknown aetiology. In earlier reports it was associated with a good prognosis. The association with hypertrophic cardiomyopathy indicates that the prognosis may not always be good and may support the idea that the syndrome is a disorder of muscle rather than of bone, nerve, or connective tissue.

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Screening for cystic fibrosis by a stool trypsin method

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SUMMARY Mass screening for cystic fibrosis by a trypsin assay of stool dried on filter paper was evaluated in 20 000 5-day-old babies. Sweat tests were required in only 7 babies. Three of them had cystic fibrosis. The test gave a false-negative result in at least 2 babies, but each had normal pancreatic function. This is not an ideal screening test for cystic fibrosis, as it misses cases with normal pancreatic function, but it is very cheap, highly specific, and appears to be the best currently available screening test.

A screening method for cystic fibrosis has long been sought. Of methods based on an assay of pancreatic function, those using meconium are inconvenient: a second sample cannot be obtained, the test cannot be performed at a central laboratory, and all have had reports of high false-positive and false-negative rates.1–2 The method of Crossley et al.3 which assays stool trypsin content, seemed sufficiently promising for a large-scale evaluation, as samples could be sent by post, and repeat samples obtained noninvasively.

We conducted a study in New South Wales, Australia to evaluate Crossley's method of screening, and this paper reports the results.

Samples and methods

Stool samples from 20 000 5-day-old babies were collected on to filter paper, dried, and posted to the laboratory. Five Sydney and 20 country hospitals took part in the project which lasted a year and so included the hot summer months.

The method of Crossley et al.3 was used to assay trypsin activity. Discs 6 mm in diameter were punched from faeces-impregnated filter paper, incubated overnight with benzoyl-l-arginine-p-nitroanilide hydrochloride (1-BAPNA). Trypsin present in samples cleaves a yellow p-nitroanilide molecule from the 1-BAPNA. Samples deficient in trypsin remain colourless or very pale yellow. The test result can be read by eye. If the yellow colour was
absent or not strong, the optical density (OD) was measured at 410 and 460 nm, as described by Crossley et al., and samples with OD_{410} = 2 (OD_{460}) of less than 0.3 were reassayed using two 6 mm discs. If samples gave a positive result (lack of trypsin activity) on re-assay, a second stool sample was obtained by writing to the mother. If the second stool sample showed lack of trypsin activity too, the baby’s doctor was asked to arrange for a sweat test to be carried out.

In addition, samples from 17 newly diagnosed and untreated patients with cystic fibrosis were obtained and tested in the same manner.

To assess the stability of trypsin activity in the samples, 30 samples were assayed before and after being exposed for 5 days to temperatures of either 18°C or 34°C in either ambient humidity, or humidity close to 100% saturation.

**Results**

**Newborn population.** Resampling was necessary in 0.53% of cases because each first sample was repeatedly positive. 93% of second samples were received, but the remaining 7% could not be obtained despite three requests.

Seven of the second samples had deficient trypsin activity, and sweat tests were performed on these babies. Two sweat tests were positive, and both the babies had cystic fibrosis. In one the sweat test was inconclusive: there were high electrolyte levels but insufficient sweat was collected. This baby, a sibling of a patient known to have cystic fibrosis, died with symptoms that could have been due to cystic fibrosis. The 4 remaining sweat tests gave negative results.

Two babies whose stool trypsin screening test was normal were subsequently diagnosed as having cystic fibrosis. Neither required pancreatic enzyme supplementation.

The false-positive rate in our survey was thus 0.53% after the first sample, and 0.02% after the second sample, in 19 997 cases, and the false-negative rate at least 40% (2 of at least 5 cases). However, no false-negative result has yet been known to have been obtained in a patient whose pancreatic function is deficient.

**Cystic fibrosis cases, before treatment.** Of 17 stool samples from known cystic fibrosis patients, a positive test result was obtained in 13, indicating deficient trypsin activity. The 4 samples which gave a negative (normal) result came from patients whose pancreatic function was adequate at the time.

**Cost of the test.** The cost of testing each baby was 11 cents (about £0.06) for clerical and technical time, chemicals, and glassware.

**Stability of trypsin activity.** Optical densities at 410 and 460 nm of samples before and after heating for 5 days at 18°C or 34°C in moist or relatively dry conditions were not significantly altered.

**Discussion**

The purpose of this study was to evaluate the sensitivity and specificity of the stool trypsin method of screening for cystic fibrosis, and to discover how easily it could be implemented in our health organisation.

The specificity of the test was fairly satisfactory. The false-positive rate on the initial assay, 0.53%, was much higher than the 0.1% found by Crossley et al., but the false-positive rate on the second sample was lower (0.02 versus 0.05%). Thus, although fewer unnecessary sweat tests had to be performed, the rate of resampling was rather higher. Crossley et al. noted that most false-positive results on the first sample appeared to be associated with prematurity in the babies, but our data did not give sufficient information about this. If the initial test could be carried out efficiently at, say, age 2 weeks, with sample collection arranged by local baby health centres, the specificity of the test would be likely to be much higher.

In our study, the sensitivity of the test was not good. We identified only 3 cases of cystic fibrosis in 20 000 patients, and 2 cases ascertained because of symptoms had had false-negative results. This total of 5 patients was about half that expected, and our false-negative rate may well prove to be higher. However, those cases we missed were patients whose pancreatic function at the time of diagnosis was adequate, and we would therefore have expected the test result to be negative in such cases.

No problems were encountered in setting up this pilot screening programme, and hospital staff had no difficulties in following sample collection instructions. The cost of performing the test was very low—$A 0.11 per baby—but no account was taken of costs associated with collection or dispatch. Nevertheless, it is certain that in Australia the cost would compete very favourably with that of using any commercial kit for meconium testing currently available on the Australian market.

Crossley et al. have recently described a blood spot method of screening for cystic fibrosis. If, when larger numbers of newborn infants have been studied, this proves to be a test with high specificity and sensitivity, it will be a significant advance in screening for cystic fibrosis. In the meantime although the stool trypsin method is not an ideal screening test, missing as it does those patients with normally functioning pancreata, in view of its cheapness, its
noninvasiveness, its low false-positive rate, its robustness, and simplicity it should be seriously considered as a possible method of screening for cystic fibrosis. It appears to be the best method currently available.

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References

Upper airways obstruction

Presentation with systemic hypertension

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SUMMARY Of 14 patients whose final diagnosis was upper airways obstruction associated with heart failure, 3 presented with systemic hypertension (up to 200/100 mmHg). In 2 the hypertension was so severe that at first it had to be considered as a possible cause of the presenting symptoms. The subsequent history indicated that it was an effect of the upper airways obstruction with heart failure.

Upper airways obstruction can lead to pulmonary arteriolar vasoconstriction, pulmonary arterial hypertension, cor pulmonale, and right heart failure.1–4 We describe 3 children who were part of a group of 14 children who presented consecutively with upper airways obstruction associated with heart failure.

Case report

Case 1. A 2-year-old boy was admitted to hospital, semi-comatose, with a 3-day history of upper respiratory infection. Blood pressure (BP) was 160/120 mmHg* and there were signs of severe congestive heart failure. Chest x-ray film showed gross cardiomegaly and signs of pulmonary oedema. Electrocardiogram (ECG) showed a normal QRS axis of +70°, right atrial P-waves, and left ventricular hypertrophy pattern. He was treated with intravenous diuretics and digitalis, and was given artificial ventilation using a respirator. Before artificial ventilation arterial Pco2 was 72.5 mmHg (9.6 kPa), Po2 63.6 mmHg (8.5 kPa), and pH 7.23. After 3 hours of artificial ventilation, Po2 was 46 mmHg (6.1 kPa), Pco2 149.5 mmHg (19.9 kPa), and pH 7.45. BP remained high (150/120 mmHg) and he remained semi-comatose. His fundi were normal but a lumbar puncture showed an opening pressure of 40 mm cerebrospinal fluid. BP fell after antihypertensive medication. By day 3 he was awake with no respiratory distress and his BP had stabilised (about 110/60 mmHg) without treatment. Renal arteriogram and vanillyl-mandelic acid (VMA) levels (0.7 mg/24 h) were normal. Because of large tonsils and adenoids it was felt that upper airways obstruction might be responsible for the symptoms, and removal of tonsils and adenoids was planned but the appointment was not kept.

The boy did not return until age 3½ when he was again admitted with a history of upper respiratory infection, in severe respiratory distress, semi-comatose, and in congestive heart failure. Again he recovered, removal of tonsils and adenoids was again planned, but again the appointment was not kept. He