Newborn screening for cystic fibrosis: pros and cons

Educational aims

- To give an insight into the arguments that have led to the implementation of newborn screening for cystic fibrosis in routine screening programmes.
- To understand the drawbacks specifically associated with newborn screening for cystic fibrosis and how these can be handled or avoided.

Summary

During the past decade, newborn screening for cystic fibrosis has been introduced in many countries. However, the disadvantages associated with newborn screening for CF hamper its introduction in other areas.

Within the routine heel-prick programmes implemented around the world, there is no other disease that elicited such a debate about whether or not to include it in these programmes as cystic fibrosis (CF).

The reasons for this debate are manifold: the diagnosis of CF is not as simple as it looks on first sight; a robust screening test suitable for universal use does not exist; and the effects of early management on the outcome are doubted by many. The improvement in survival of patients with CF observed during the last few decades [1] has been attributed to an improvement of the traditional therapies for CF. Examples include: pancreatic enzyme replacement therapy, high caloric intake and supplementation of fatsoluble vitamins for CF gastrointestinal disease; early eradication of airway infection with Pseudomonas aeruginosa (inhalation therapy with antibiotics when the infection becomes chronic); and improved therapies for mucociliary clearance using dornase alfa and hypertonic saline. However, therapies aimed at the basic defect have not yet been developed.

Newborn screening for CF allows early diagnosis, often when the child is still asymptomatic, or only has a few symptoms related to poor growth but no pulmonary problems. However, the evidence that starting treatment before respiratory problems have arisen leads to a better outcome on the long term is not very strong, but there is ample evidence that in the first decade of life, CF patients identified by screening have a better clinical condition than patients diagnosed clinically.

In this article we shall discuss the most common arguments for and against newborn screening for CF (NBSCF).

Arguments for NBSCF

The best arguments for screening can be found when observing cohorts of patients with CF diagnosed clinically in areas where no newborn screening programme for CF is running.

Newborn screening leads to diagnosis at a very young age. In most countries without newborn screening, the median age of diagnosis is ~1 yr, and when patients with a meconium ileus are excluded, the median age is even higher; for example, in Canada, the
median age at diagnosis is >2 yrs and in Sweden, it is ~6 months [2, 3], while with a newborn screening programme, the median age at diagnosis decreases to 29 days [4].

In regions without screening, clinical symptoms finally lead to the diagnosis. Observational data from Canada show that 81% of clinically diagnosed CF patients have respiratory symptoms, many show failure to thrive (37%) and reduced serum concentrations of fatsoluble vitamins due to malabsorption, and 95% have gastrointestinal complaints at the time of diagnosis [2]. The mean head circumference of patients with a vitamin E deficiency was significantly lower, at the 32nd percentile (95% CI 24–41%), than that of patients with a normal vitamin E concentration, of whom the mean head circumference was at the 63th percentile (95% CI 47–78%) [2].

Hospital admissions are not uncommon during the period in which children develop symptoms: 79% of the children were hospitalised, with a mean length of stay of 5 days (range 1–30 days) in a tertiary care hospital before the diagnosis CF was made. Some children already had severe bronchiectasis at the time of diagnosis [5].

Many of these problems associated with a late diagnosis can be prevented. Evidence for beneficial effects of screening has been reported in two randomised clinical trials [6–8], in analyses form large clinical databases [9–11] and from a few cohort studies [12–17].

Treatment of pancreatic insufficiency
Gastrointestinal problems in CF are mostly related to pancreatic insufficiency. Most patients with CF are pancreatic insufficient at birth or develop pancreatic insufficiency during the first year of life [18]. With an early diagnosis, pancreatic disease can be treated from the start with pancreatic enzyme replacement therapy.

Growth
In a randomised clinical trials [6, 8], a large clinical database [9] and four observational cohort studies, a significant improvement of height and weight was observed compared with clinically diagnosed CF [11, 14, 16, 17]. With an early diagnosis, children with CF can maintain a growth pattern that approximates that of healthy children [6].

Nutritional deficiencies
Malnutrition by protein deficiency and significant morbidity due to deficiencies of fatsoluble vitamins have been described in clinically diagnosed patients [19, 20], and newborn screening can prevent this morbidity. Vitamin deficiency can occur very early in life. Many infants diagnosed by screening already show low concentrations of vitamin A (60%), D (37%) and E (16%) [21], underlining the need for early diagnosis and start of treatment.

There is indirect evidence that early malnutrition and nutritional deficiencies early in life are related to lung growth. Patients with CF with pancreatic insufficiency who achieved early growth recovery within 2 years of diagnosis had fewer cough symptoms, higher lung function and better chest radiography scores at 6 years of age [22].

Lung function
A randomised clinical trial showed better chest radiography scores early in life but at a later age, no difference in lung function between the screened cohort and the clinically diagnosed patients was found [23]. This may be related to the fact that the screened cohort had earlier P. aeruginosa colonisation than the clinically diagnosed cohort. In all cohort studies, significantly better chest radiography scores were observed in the screened cohorts. Cross-sectional studies with data derived from the US database found better lung function in screened patients aged 6–10 and 11–20 years [20]. Cohort studies from the Netherlands and Australia showed better lung function until 12 and 15 years of age in the screened cohorts, respectively [12, 15]. In the UK database, no differences in lung function was observed between screened and nonscreened cohorts, but the screened cohort needed significantly less treatment [24].

There is also indirect evidence that early diagnosis and treatment are related to better lung function, as a cohort study showed that in sibling pairs, the younger siblings show a significantly better lung function at adult age [25]. Moreover, while recruiting patients <12 years of age for the Early Pseudomonas Infection Control (EPIC) study, it was found that patients with a diagnosis through newborn screening had a significantly better forced expiratory volume in 1 s % predicted than clinically diagnosed patients [26].
It is possible to eradicate *P. aeruginosa* during the first infection in practically all patients, but practically impossible to eradicate *P. aeruginosa* when the infection has become chronic. *Pseudomonas* can colonise the airways of young infants with CF in the first months of life without causing symptoms but leading to infection, inflammation and structural lung disease [27]. The observation that an early diagnosis does not change the time to first acquisition of *Pseudomonas* is another argument for the need for NBSCF: with an early diagnosis and thorough monitoring of airway infection, it is possible to start eradication treatment of the first infections with pathogenic micro-organisms, which can postpone chronic infection for a long time.

**Survival**

In one randomised clinical trial, a higher risk was found for an early death in the clinically diagnosed patients [7]; a systematic review also reported a lower mortality risk in screened cohorts [28].

**Hospital admissions and burden of care**

Fewer hospital admissions were found in one randomised clinical trial [7] and in two cohort studies [16, 20]. In the UK database cohort study, less therapy was needed, and improved growth and reduced morbidity were found in the screened cohort [24].

**The opinion of parents**

Parents prefer an early diagnosis even when their child has an untreatable disorder [29] in order to prevent a long diagnostic odyssey. In a recent study in Sweden, where currently no NBSCF takes place, parents favoured screening [30]. Parents of children with CF with a diagnosis within the first 3 months of life had more confidence in the medical profession and less negative feelings than parents of children with a late CF diagnosis [31].

An early diagnosis of a genetic disease such as CF also renders the opportunity that parents receive genetic counselling, which is important for further family planning. In some countries with longstanding newborn screening programmes for CF, the prevalence of CF at birth has been steadily diminished during the last 20 years [32].

**Research needs**

Present treatment for CF cannot reduce the increased susceptibility of CF airways to bacterial infection, and once the airways have become infected it is extremely difficult to eradicate the invading microorganisms. There is a need for research that can elucidate the process of bacterial infection in the CF airway, which may lead to improved treatment options. Clinical trials that investigate interventions in young infants without pulmonary involvement can only be performed when infants with CF are identified early in life, before pulmonary involvement, *i.e.* by newborn screening.

**Economic arguments**

Newborn screening for CF seems cost effective and can lead to cost savings, as costs of treatment can be reduced by 10% [33, 34]. A more recent analysis assumed that the costs of screening, diagnosis and treatment in the first 3 years of life are about 71% of the costs of diagnosis and treatment without a newborn screening programme, which means that NBSCF can lead to savings [35].

**Arguments against NBSCF**

**Effect on parents due to abnormal results of newborn screening tests**

Most parents experience high levels of emotional stress during their wait for further diagnostic testing, mostly the sweat test. It may cause depressive symptoms that vary depending on their perceptions of how likely it is their child has CF [36]. A good protocol for handling abnormal screening test results with a minimum waiting period between the notification of the abnormal test result and the sweat test, and good information of the parents may reduce much of the parental stress [37].

**False-positive tests**

Practically all screening programmes lead to abnormal results that, during the following diagnostic process, appear to have been a false alarm. Such false alarms often lead to parental stress and anxiety. However, in the long-term, when the child appears to be healthy and with good parental education,
parental anxiety levels do not differ from parents of children who did not get such an alarm [37].

**Identification of carriers**

Screening approaches that include CF transmembrane conductance regulator (CFTR) mutation analysis also identify healthy infants carrying one CFTR mutation. An advantage of this finding may be that parents are offered genetic counselling, and occasionally both parents may turn out to be carriers. This can be important information for further family planning. However, in most cases, only one parent will be identified as a carrier, which can lead to anxiety and stress. From the child’s perspective, the knowledge of being a carrier is not of direct and immediate benefit. Moreover, as the child could not decide whether or not he/she wished to be tested, this can be considered as a violation of the “right not to know”.

**Inconclusive results of NBSCF**

In most screening programmes, quite a number of infants are found by newborn screening in whom the diagnosis CF cannot be confirmed nor excluded, because they do not meet the diagnostic criteria for CF. Mostly, they have elevated immunoreactive trypsinogen (IRT) concentrations, normal or borderline sweat chloride concentrations, and carry one or two CFTR mutations with an unclear or unknown pathogenic effect on phenotype. Some of these infants will develop a complete or partial CF phenotype later in life, but many will never develop any symptoms. The uncertainty about prognosis is difficult for genetic counselling and requires that such infants get a regular further follow-up at CF centres. This may have a lifelong impact on the child and his or her family.

**Risk for early colonisation with P. aeruginosa**

Various studies showed less chronic airway infection with *P. aeruginosa* [10, 13, 20, 24]. However, in one randomised clinical trial, infants identified by screening developed colonisation and chronic infection with *P. aeruginosa* at an earlier age than clinically diagnosed patients with CF [23]. This may relate to cross-infection. The finding underlined the fact that patients with CF have an innate increased susceptibility for early airway colonisation with opportunistic micro-organisms, such as *Staphylococcus aureus* and *P. aeruginosa*. Infants identified by newborn screening will mostly visit CF clinics at an earlier age than clinically diagnosed patients; special hygienic measures to prevent cross-infections from older CF patients should be in place in every CF clinic.

**Risk of ethnic discrimination**

The use of a screening strategy including CFTR mutation analysis carries the risk that very rare mutations will not be identified. In countries and cities with large multiethnic populations, this may lead to more missed cases in infants originating from non-Western countries. A European survey of mutations in CF patients of North-African or Turkish descent showed that only 63% of patients of Turkish origin would be identified by newborn screening [38]. It may be possible to compensate by using procedures such as a second IRT test if the first level was very high, or by performing DNA scanning of the CFTR gene in samples with very high IRT concentrations without mutations in the common panel.

Many of the hazards associated with running a newborn screening programme for CF can be reduced. Notification of abnormal test results should lead to prompt referral and diagnostic testing according to an established protocol with solid parental information. Parents should be directed to safe and reliable sources of information on internet. Novel screening strategies can drastically reduce the number of false positives and equivocal diagnosis [35]. Infants with a CF diagnosis should not be exposed to older patients with CF to prevent cross-infection with *P. aeruginosa*.

**Discussion**

Most arguments that favour NBSCF seem to be dependent on the management and treatment that can be given to the child with CF identified by screening. NBSCF leads to diagnosis before lung disease and malnutrition have appeared. Whilst after a clinical diagnosis, all efforts are primarily directed to treating disease in the affected organs, after a diagnosis by newborn screening, the main challenge is to keep the infant as healthy as possible. This should lead, and often does lead, to a change to proactive, preventive medicine rather than reactive damage limitation. Recent evidence supports the notion that very early treatment of pulmonary infection and inflammation is a key factor for further improving prognosis of patients with CF.
and further research in this area is urgently needed.  

Most arguments against screening seem to be related to the imperfect properties of the available screening tests for CF; therefore, as tests improve, most of the arguments against screening will disappear. Since May 2011, the Dutch newborn screening programme has been extended with screening for CF after a pilot study assessing the characteristics of two novel screening strategies [35, 39]. The screening test now in use in the Netherlands differs from screening tests used in other countries: it has a very high specificity and positive predictive value, and detects only a very small number of carriers and patients with an equivocal diagnosis [35]. With this screening programme, most of the arguments against screening, as discussed in this article, are no longer relevant.

Finally, costs of screening cannot be used as an argument against newborn screening for CF. Although the savings of an early diagnosis of CF are not as clear cut as in newborn screening for congenital hypothyroidism or phenylketonuria, NBSCF and treatment appear to cost less than a clinical diagnosis and treatment [25].

Conclusion

The benefits of NBSCF are now generally accepted. The benefits of NBSCF amply compensate for the disadvantages. Moreover, NBSCF is cost-effective and there is growing evidence that it leads to cost savings. Many of the disadvantages of NBSCF are reduced to practically nil with the novel screening strategy that has recently been introduced in the Netherlands.

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