Paediatric screening for hypercholesterolaemia in Europe

D M Kusters,1,2 C de Beaufort,3 K Widhalm,4 O Guardamagna,5 N Bratina,6 L Ose,7 A Wiegman1

1Department of Pediatrics, Academic Medical Centre, Amsterdam, The Netherlands
2Department of Vascular Medicine, Academic Medical Centre, Amsterdam, The Netherlands
3DECCP, Clinique Pédiastrique/ Centre Hospitalier de Luxembourg, Luxembourg, Luxembourg
4Division of Clinical Nutrition, Department of Pediatrics, Medical University Vienna, Vienna, Austria
5Department of Pediatrics, Turin University, Turin, Italy
6Department of Pediatric Endocrinology, Diabetes and Metabolic Diseases, University Children’s Hospital, University Medical Centre, Ljubljana, Slovenia
7Lipid Clinic, Oslo University Hospital Rikshospitalet, Oslo, Norway

Correspondence to
Albert Wiegman, Department of Pediatrics, Academic Medical Center, Meibergdreef 9, 1105 AZ Amsterdam, The Netherlands; a.wiegman@amc.nl

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ABSTRACT

Different screening strategies are currently recommended to identify children with (familial) hypercholesterolaemia in order to initiate early lipid management. However, these strategies are characterised to date by low adherence by the medical community and limited compliance by parents and children. In a literature review, the authors assess which children should undergo screening and which children are in effect identified through the currently recommended strategies. Furthermore, the authors discuss the different screening tools and strategies currently used in Europe and what is known about the negative aspects of screening. The authors conclude that currently recommended selective screening strategies, which are mainly based on family history, lack precision and that a large percentage of affected children who are at increased risk of future coronary artery disease are not being identified. The authors propose universal screening of children between 1 and 9 years of age, a strategy likely to be most effective in terms of sensitivity and specificity for the identification of children with familial hypercholesterolaemia. However, this concept has yet to be proven in clinical practice.

INTRODUCTION

Hypercholesterolaemia is one of the major modifiable risk factors for the development of atherosclerosis and cardiovascular disease (CVD).1 Since the process of atherosclerosis starts in early childhood,2 hypercholesterolaemia requires timely diagnosis and management. This is especially the case when marked elevated cholesterol levels are inherited and present from birth onwards. Individuals in early adulthood with untreated familial hypercholesterolaemia (FH) or familial defective ApoB (FDB) are nearly 100 times more likely to develop coronary artery disease than unaffected individuals.3 As children with hypercholesterolaemia generally do not present with any complaints or symptoms, paediatric organisations have developed different screening strategies to identify such children.4–7 However, the strategies that do exist are characterised by low adherence by the medical community and limited compliance from parents and children.

Screening for hypercholesterolaemia in children is worthwhile if the following three assumptions are true: children with hypercholesterolaemia are at increased risk of developing accelerated atherosclerosis and subsequent CVD; screening can accurately identify those at increased risk; and existing therapies can effectively and safely reverse the atherosclerotic process and reduce the CVD risk in these children. We can safely assume that, based on current evidence, the first and third requirements are met, and the remaining question therefore is: which screening strategy can best identify those at increased risk?5 In this review we will provide an outline of the currently recommended screening strategies and will discuss which children are identified with these strategies and which children are not and might be still at risk. Subsequently, we will present some suggestions for improving these strategies across Europe. As early diagnosis and management are especially important in children with FH, the emphasis will be on these children.

EVIDENCE FOR EARLY Atherosclerosis

It was shown that risk factors measured during the teenage years and in early adulthood predicted the presence of subclinical atherosclerosis that developed 15 years later much more accurately than the same risk factors measured at the time of lesion assessment.5–7 This indicates that the prevention of risk factors in childhood or adolescence is likely to be most effective and, in fact, this ‘the younger the better’ strategy is rapidly gaining acceptance vis à vis ‘the lower the better’ concept. Evidence for this new approach comes from statin trials in children with severe hypercholesterolaemia which show that atherosclerotic lesions are reversible if drug treatment is initiated really early.10

The first indications of atherosclerosis are already present in children with FH, despite the absence of complaints or symptoms. Increased inflammatory activity is reflected by elevated levels of high-sensitivity C-reactive protein (hsCRP) in children with FH as compared with their unaffected siblings.11 Additionally, functional and morphological changes in the arterial wall are observed in children with FH, which indicates that the atherosclerotic process has already begun. This is illustrated by impaired flow-mediated dilatation (FMD) of the brachial artery12 and increased intima media thickness (IMT) of the carotid artery,13 respectively. These findings have led to the hypothesis that treatment should be initiated early in life to reduce the incidence of CVD in later years.

Increased inflammatory activity and impaired endothelial function indicating enhanced atherogenesis is also present in obese children.14–15 Furthermore, children with type 1 diabetes exhibit impaired FMD and increased carotid IMT.16


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### Screening in Europe

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### Table 1 Screening strategies in five European countries

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<th>Country</th>
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<td>Children of FH families</td>
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CVD, cardiovascular disease; FH, familial hypercholesterolaemia.

In the Netherlands, there is an ongoing cascade screening programme to identify patients with FH, in which all first degree family members of an index patient are identified and offered a DNA test. Cascade screening then is offered to more distant relatives using the inheritance pattern across the pedigree. To date, >20 000 individuals with FH carrying more than 550 different mutations have been identified, which is estimated to be 50% of the total number of FH patients in the Netherlands. A literature review reported that screening of the relatives of index cases with FH is the most cost-effective screening strategy.20

In Norway, as in the Netherlands, all genetic FH tests are performed by a single centre which has been working closely with the Lipid Clinic in Oslo. The total number of FH patients has been estimated at 16 000 and approximately 5000 individuals with >140 different mutations have been identified so far. A cascade screening program has been established,21 but as the national health system in Norway has not provided funding for other areas of the country, laboratory investigations will remain dependent on the Lipid Clinic and the Medical Genetic Laboratory of the University of Oslo.

The Slovenian Pediatric Society decided to initiate general cholesterol screening in preschool children at the age of 5 in 1995. Since then, more than 15 000 children have undergone screening annually. A blood sample is taken at a routine examination and total cholesterol (TC) values are measured. If TC values are elevated, a fasting lipid profile is performed. According to national guidelines, the child is referred to University Children’s Hospital in Ljubljana if TC levels are >6 mmol/l or >5 mmol/l and there is a positive family history of premature CVD. The data of these children are used for research and from 2011 onwards selective genetic screening for FH will also be carried out.

In Italy selective screening is delegated to general practitioners or paediatricians who follow Italian guidelines.22 Children considered at risk and suspected of having an inherited lipoprotein disorder (showing a positive family history of premature CVD or exhibiting dyslipidaemia with TC >6.2 mmol/l, HDL-C <0.9 mmol/l or triglyceride (TG) >5.4 mmol/l) are invited to attend lipid units where they undergo biochemical and genetic analysis. However, only a limited number of children complete this program, accounting for less than 10% of the total number of children with possible FH. To bridge the gap between the number of children detected and the number theoretically affected, an extensive program of recruitment, diagnosis and registration is currently in progress. This program is aimed at promoting awareness of the benefits of screening, extending this practice throughout Italy, defining the genetic mutations involved in FH (LDLR, ApoB, PCSK9) and providing an Italian register of genetically detectable lipoprotein disorders.

In the UK, the National Institute of Healthcare and Clinical Excellence (NICE) has recommended a family cascade
testing approach for FH, starting with adults who have a clear (preferably genetic) diagnosis. The NICE is regarded as the main organisation that promotes evidence based clinical practice in the UK. Testing of children in families known to have a clear diagnosis of FH is recommended from the age of 10 years. Recently, this approach has been fully implemented (2010) in Wales, which is one of the devolved countries within the UK.25

Wald and colleagues recently proposed the alternative strategy of screening children in the UK when they visit their general practice for routine vaccination at around 15 months of age. In children aged 1–9 years, the use of TC levels as a cut off yields a false positive rate of 0.1% (1.53 multiples of the mean) and would identify 88% of cases with FH. Measurement of cholesterol levels in the parents of an affected child would determine which parent was affected. Applying the rule that the parent with the highest cholesterol levels has FH gives a detection rate of above 96%.24

**Screening tools**

Once children are selected for screening, lipid levels are measured. A raised LDL-C level is the trigger for initiating treatment and determining therapy goals in children2 7 and is generally calculated according to the Friedewald formula. However, because this is dependent on TG levels, a fasting sample is required. The advantage of a TC measurement is that it is not necessary to obtain a fasting sample. In the Bogalusa Heart Study, increased TC levels detected elevated LDL-C levels with 44–50% sensitivity and 90% specificity.29

It was shown that the apoB/apoA-1 ratio measured in adolescents (12–18 years) was superior to the LDL/HDL ratio for predicting increased IMT in adulthood.26 A high apoB/apoA-1 ratio indicates the ratio between pro- and antiatherogenic lipoprotein particles. Therefore, measuring these apolipoproteins might have additional value in risk assessment and screening in children.

An important issue is the accuracy of paediatric lipid levels to identify children at increased risk. Approximately 40–55% of children with elevated lipid levels will continue to have elevated lipid levels in adulthood.27 On the other hand, subjects with multifactorial/polygenic causes of hypercholesterolaemia often exhibit normal lipid levels during childhood. Therefore, those with a strong family history of premature CVD in particular should be retested after puberty. At least two measurements are necessary to ensure that the mean measurement reflects the true value to within 10%. Children with very high LDL-C levels (≥4.1 mmol/l (≥160 mg/dl)) are likely to have substantially lower levels (mean decrease 0.5–0.75 mmol/l) at re-examination, which is attributable to regression to the mean.28 Furthermore, as childhood is a time of rapid growth and development, especially during puberty, and cholesterol is involved in the structure of cellular membranes and hormones, lipids vary with age among children.29 This must also be taken into account when interpreting cholesterol levels.

The criteria for a clinical diagnosis of FH are LDL-C levels above the 95th percentile for age and gender, a family history positive for CVD, and physical symptoms of cholesterol deposits in the skin, eyes or tendons known as xanthelasmas, arcus cornea and tendon xanthomas, respectively, which are typical for FH although rare in children.30 Once there is a clinical diagnosis, a definite diagnosis may be made using DNA based mutation screening methods. Once the causative mutation in the index patient has been identified, reasonably cheap molecular testing in relatives is possible. Molecular testing is simple in countries with only a few different mutations causing FH. On the other hand, in the majority of countries with full-blown genetic heterogeneity, DNA sequencing could fail to identify a mutation, with detection rates ranging from only 20% to 90%.31–35 However, as mentioned previously, extreme hypercholesterolaemia in children is less likely than in adults to be polygenic, which allows a higher mutation detection rate. In fact, in a recently published study it was found that in 95% of children with an FH phenotype, a functional gene mutation could be found in the LDL receptor or ApoB genes.36

Identification of the FH-causing mutation in a family permits a definite diagnosis, and the diagnostic problem caused by the overlap in cholesterol levels between the general population and the FH patient is thereby eliminated. A single test, performed once only, will ascertain FH status, and early diagnosis in these children, at least, becomes a reality.

Neonatal screening has been examined in some studies, but none followed abnormal results with mutation analysis, which makes it difficult to determine the value of such screening.27 The usefulness of IMT measurement as a screening tool in children must be further explored.

**WHO TO SCREEN?**

Lipid screening could identify three groups of children: those with monogenic hypercholesterolaemia (mainly heterozygous FH and FDB), secondary causes of hypercholesterolaemia and multifactorial (polygenic) hypercholesterolaemia. The last group is less common in children. Once children with hypercholesterolaemia are identified, lipid management can be started. The recently updated treatment recommendations of the AAP37 are summarised in box 2.

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**Box 2**  **AAP recommendations for the treatment of hypercholesterolaemia in children, 2008**

1. **Diet/lifestyle**
   - a. All children >2 years of age: healthy diet according to Dietary Guidelines for Americans
   - b. Children between 12 months and 2 years of age with family history of obesity, dyslipidaemia or CVD, or for whom overweight is a concern: use low-fat milk
   - c. Children and adolescents at high risk of CVD: diet with <7% saturated fat, cholesterol <200 mg/day and <1% transfats
   - d. For overweight or obese children with high triglyceride or low HDL-C levels: weight management, including dietary counselling and increased physical activity

2. **Pharmacology**
   - a. For patients ≥8 years of age, pharmacological intervention should be considered if:
     - LDL-C ≥4.9 mmol/l (190 mg/dl) OR
     - LDL-C ≥4.1 mmol/l (160 mg/dl) AND family history of premature heart disease OR ≥2 additional risk factors (obesity, hypertension, cigarette smoking) OR
     - LDL-C ≥3.4 mmol/l (130 mg/dl) AND diabetes mellitus
   - b. When there is a strong family history of CVD, especially with other risk factors, target levels for LDL-C as low as 3.4 mmol/l (130 mg/dl) or even 2.8 mmol/l (110 mg/dl) should be considered

AAP, American Academy of Pediatrics; CVD, cardiovascular disease; HDL-C, high-density lipoprotein cholesterol; LDL-C, low-density lipoprotein cholesterol.
Because FH is characterised by markedly elevated levels of LDL-C from birth onwards, leading to premature atherosclerosis and CVD, the importance of early diagnosis and management is well established. Underdiagnosis of (heterozygous) FH is a worldwide problem and therefore screening for FH is strongly advocated. The problem in that both TC and LDL-C levels show considerable overlap between FH and normal individuals plays a lesser role in children as compared with adults. It was recently shown in a meta-analysis that cholesterol levels discriminate best between people with and without FH at the ages of 1–9 years. Therefore, a diagnosis of FH should always be considered in children with elevated lipid levels.

Children with diseases known to possibly cause secondary dyslipidaemia, such as nephrotic syndrome, HIV treated with protease inhibitors, systemic lupus erythematosus or conditions after solid organ transplantation, should also be screened.

Screening for hypercholesterolaemia (and/or hypertriglyceridaemia) is of special importance in children with disorders that are characterised by the presence of other cardiovascular risk factors, such as diabetes and obesity. According to the 2006 AHA statement, children with type 1 diabetes are classified in tier 1, the highest cardiovascular risk category. Screening guidelines within the European Union for children and adolescents with type 1 or type 2 diabetes have been recently published in the SWEET report.

**Potentially Harmful Aspects of Screening**

Children with FH or other causes of severe hypercholesterolaemia do not have clinical disease, but the knowledge of future risks and the awareness of having the disorder and taking medication every day, raises the question of whether this influences their quality of life. There is limited evidence on this subject, but it was shown that children diagnosed with FH did not have elevated anxiety levels, that taking medication made the children feel safer and that they did not ‘mind’ taking this medication for the rest of their lives. Another study showed that children generally coped well with their carrier status and its implications. Studies in adults suggested that relatives usually believe that genetic information is beneficial and reports on the impact of receiving a diagnosis of FH showed that the proportion of individuals experiencing anxiety was no higher than in the general population. A study involving participants of the Dutch cascade screening program reported that 87% of parents from FH families wanted their children to undergo a genetic test. Another study showed that (treated) individuals with FH who apply for life or disability insurance should be accepted at standard rates if their TC and LDL-C level is below 4 mmol/l and no additional cardiovascular risk factors are present.

**The Importance of Adequate Follow-Up**

Screening is only valuable if adequate follow-up is guaranteed. The full implementation of cascade screening requires an infrastructure that allows for both family tracing and the increasing clinical workload as new cases are identified. For example, in the Netherlands, cascade testing is coordinated through national centres supported by the government, and patient care is delivered by existing lipid clinics. However, such arrangements are still limited for children. In the ideal situation, all identified children should be linked to a recognised specialist network managed by a paediatrician. This should ensure that all children are referred to a specialised paediatric lipid clinic and are adequately informed and treated. More specialised paediatric lipid clinics should be established to provide sufficient capacity to admit identified paediatric patients with FH and other types of dyslipidaemia.

**Conclusions and Recommendations**

Screening for dyslipidaemia in children is only rational if screening tests and strategies can accurately identify those at increased risk. In 2007, the US Preventive Service Task Force concluded that the evidence is insufficient to recommend for or against routine screening for lipid disorders in infants, children, adolescents or young adults. Based on the mainly observational studies discussed above, we conclude that there is good evidence that currently recommended selective screening strategies mainly based on family history are not accurate and that an important proportion of children at risk are not being identified. Furthermore, the need for fasting samples and repeated measurements is likely to affect feasibility and compliance with cholesterol screening, even in a targeted approach. There is also evidence that enhanced atherogenesis is present in some children, and that this process is reversible when early treatment is initiated. We will make some recommendations based on the current evidence, although clinical trials are needed to determine the best screening approach.

Paediatricians should screen children with disorders known to possibly cause secondary dyslipidaemia, such as obesity and diabetes. The most difficult but important challenge, however, is to identify children with FH, as they experience no symptoms while the importance of early disease management is well established. The imprecision of clinical screening strategies for FH emphasises the importance of genetic testing for a definite diagnosis of FH. Although cascade screening has been very successful in the Netherlands, this strategy may not be cost-effective in larger countries where families are small and geographically dispersed, and genetic heterogeneity is prevalent.

The proposed strategy of universal screening of children when they visit their general practitioner for routine vaccination at about 15 months of age, or at 5 years in Slovenia, is promising. However, this strategy is based on theoretical considerations and has not been proven in practice. Furthermore, as drug treatment is currently not initiated before the age of 8, the correct age for screening needs to be discussed. If this approach is as effective as claimed, it should be followed by mutation analysis to establish the diagnosis of FH. Once a child has been identified as having FH, cascade screening of more distant relatives should be performed using the inheritance pattern across the pedigree.

Once children with FH or other causes of hypercholesterolaemia are identified, lipid management can be started. Clinicians should strongly consider statin therapy in children with FH and elevated LDL-C, when LDL-C cannot be adequately reduced by lifestyle modification alone. However, there are no data on the safety of life-long lipid-lowering therapy and the exact age at which therapy should be initiated is also uncertain. Future research should address these questions. Nevertheless, an adequate screening program with sufficient follow-up is very likely to improve the management of children with hypercholesterolaemia, leading to better control of cardiovascular risk.
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