Technical report: ethical and policy issues in genetic testing and screening of children

Laine Friedman Ross, MD, PhD1-2, Howard M. Saal, MD3, Karen L. David, MD, MS4,6 and Rebecca R. Anderson, JD, MS5; and the American Academy of Pediatrics; American College of Medical Genetics and Genomics

The genetic testing and genetic screening of children are commonplace. Decisions about whether to offer genetic testing and screening should be driven by the best interest of the child. The growing literature on the psychosocial and clinical effects of such testing and screening can help inform best practices. This technical report provides ethical justification and empirical data in support of the proposed policy recommendations regarding such practices in a myriad of settings.

Key Words: carrier identification; disclosure; genetic screening; genetic testing; newborn screening; predictive testing

INTRODUCTION

Two major events occurred in the 1950s that forever changed the influence of genetics in medicine: Watson and Crick1 described the double-helix model of DNA structure in 1953, and in 1956 Tjio and Levan2 established that the typical human carries 46 chromosomes. The goal of mapping and sequencing the human genome began in 1990, and a working draft was presented in 2000, with a more complete edition published in 2003.3 Knowledge of genetics and genomics continues to grow rapidly, as does consumer interest in genetic testing. As a result, statements about genetic testing and screening of children in the United States written in the past two decades need to be updated to consider the ethical issues that arise with the new technologies and expanded uses of genetic testing and screening.4,5 The growing literature on the psychosocial and clinical effects of genetic testing and screening can help inform us about best practices regarding diagnostic genetic testing, pharmacogenetics, newborn screening, carrier screening, testing asymptomatic children for genetic conditions that present later in childhood or adulthood, and how to respond to direct-to-consumer testing and the potential of genomic profiling.

Genetic testing and screening of minors are commonplace. “Genetic screening” denotes assays undertaken on a population-wide basis to identify at-risk individuals. “Genetic testing” denotes assays designed to provide a definitive diagnosis; these are performed because of positive screening results, family history, ethnicity, physical stigmata, or other reasons. Every year, approximately 4 million infants in the United States undergo newborn screening with state-chosen panels of rare metabolic, hematologic, and endocrine abnormalities for which early treatment may prevent or reduce morbidity or mortality. Most of the genetic conditions included in the state screening panels are autosomal recessive disorders, and some assays identify heterozygote carriers (e.g., hemoglobinopathies). Future screening may expand to X-linked conditions (e.g., Duchenne muscular dystrophy) and autosomal dominant conditions. In addition, universal newborn hearing screening allows for early identification of both acquired and hereditary hearing loss.

Outside of newborn screening, pediatric genetic testing is much less common. Diagnostic genetic testing may be performed on a child with physical, developmental, or behavioral features consistent with a potential genetic syndrome or for pharmacogenetic drug selection and dosing decisions. Predictive genetic testing may be performed on a child with a positive family history for a specific genetic condition, particularly if early surveillance or treatment may affect morbidity or mortality. Minors may also undergo histocompatibility testing as potential organ or tissue donors. This technical report provides ethical justification and empirical data in support of the recommendations enumerated in the accompanying policy statement.6 Genetic research (including the use and retention of dried blood spots and whole-genome sequencing) is beyond the scope of this technical report and accompanying policy statement.

DIAGNOSTIC GENETIC TESTING

When performed for diagnostic purposes in a child with symptoms of a genetic condition, genetic testing is similar to other medical diagnostic evaluations. Parents or guardians should

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1Department of Pediatrics, University of Chicago, Chicago, Illinois, USA; 2Department of Medicine, MacLean Center for Clinical Medical Ethics, University of Chicago, Chicago, Illinois, USA; 3Division of Human Genetics, Cincinnati Children’s Hospital Medical Center, Cincinnati, Ohio, USA; 4Department of Pediatrics, Metropolitan Hospital Center, New York, New York, USA; 5College of Public Health, University of Nebraska Medical Center, Omaha, Nebraska, USA; 6Department of Medicine, New York Methodist Hospital, Brooklyn, New York, USA. Correspondence: Laine Friedman Ross (lross@uchicago.edu)

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be informed about the potential benefits and potential harms of testing, and their permission should be obtained. Medical benefits include the possibility of preventive or therapeutic interventions, decisions about surveillance, the clarification of diagnosis and prognosis, and recurrence risks. Medical harms occur if parents or guardians respond to the results by pursuing unproven treatments or preventive measures, particularly if they are ineffective or have significant adverse effects (e.g., megadoses of vitamin A have been proposed for children with developmental disabilities but can cause liver toxicity).7 Discovery of misattributed parentage8 is a potential risk of certain genetic tests. Other “incidental” findings can be life-saving (e.g., identifying a gene that predisposes to a preventable cancer) or psychologically disruptive (e.g., identifying a gene sequence variant of unknown clinical significance). If the medical benefits of a test are uncertain, will not be realized until a later time, or do not clearly outweigh the medical risks, the justification for testing is less compelling.

Pharmacogenetic testing to determine safety and efficacy of drugs is expanding.9 Currently, such testing is most commonly used in pediatrics for drug selection and dosing regulation in cancer therapies,10,11 psychiatric conditions,12–14 pain management,15–17 and asthma.18,19 The aim of pharmacogenomics is to improve therapeutic responsiveness and reduce the incidence of adverse drug reactions.20 If test results are expected to have clinical significance beyond drug selection and dosing (e.g., diagnostic or prognostic implications), then the additional implications of the genetic information should be disclosed before testing.

**NEWBORN SCREENING**

Virtually every infant in the United States has blood collected to screen for a variety of metabolic, endocrine, hematologic, and infectious conditions within the first week of life. State-administered newborn screening programs arose following Guthrie’s development of an assay in 1961 to detect phenylketonuria using dried blood spots collected on filter paper. Initially, some professional groups opposed the rapid population-based implementation of this assay, citing the lack of long-term data on the dietary intervention and correctly predicting that a subset of children could be harmed by overtreatment.21 However, the devastating nature of phenylketonuria, its treatability, and the availability of an inexpensive assay made widespread population screening both feasible and acceptable.

As public health screening continued to expand, the World Health Organization commissioned a study by Wilson and Jungner22, who in 1968 enumerated 10 criteria “to guide the selection of conditions that would be suitable for screening” (Table 1). Although not written specifically for genetic applications, the criteria have served as a policy standard for population-based genetic screening over the past four decades. In recent years, some authorities have advocated minor23 or substantial24 modifications to the criteria in response to evolving technologies. Despite variability in how the criteria are interpreted,25 there is broad consensus that to qualify for population-based screening, the natural history of a candidate condition should be understood, an acceptable intervention for affected individuals should be available, and cost-effective screening and confirmatory testing should be available.22,24

<table>
<thead>
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<th>Table 1 Wilson and Jungner classic screening criteria</th>
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<td>1. The condition sought should be an important health problem.</td>
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<td>2. There should be an accepted treatment for patients with recognized disease.</td>
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<td>3. Facilities for diagnosis and treatment should be available.</td>
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<td>4. There should be a recognizable latent or early symptomatic stage.</td>
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<td>5. There should be a suitable test or examination.</td>
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<td>6. The test should be acceptable to the population.</td>
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<td>7. The natural history of the condition, including development from latent to declared disease, should be adequately understood.</td>
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<td>8. There should be an agreed policy on whom to treat as patients.</td>
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<td>9. The cost of case finding (including diagnosis and treatment of patients diagnosed) should be economically balanced in relation to possible expenditure on medical care as a whole.</td>
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<td>10. Case finding should be a continuing process and not a “once and for all” project.</td>
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Data from ref. 22.

Debates about which conditions should be included in state newborn screening panels grew in importance as variations grew between states and advocacy groups began to express equity concerns.26 By the mid-1990s, some states were screening for more than 30 conditions, whereas others were screening for fewer than 5.27 In 2002, the American College of Medical Genetics and Genomics (ACMG) conducted a systematic evaluation of newborn screening programs, sponsored by the Maternal and Child Health Bureau of the Health Resources and Services Administration. Eighty-three conditions were assessed for possible inclusion in a uniform screening panel.28 After evaluating the empirical evidence for these conditions and gathering opinions from experts and advocacy constituents, the ACMG chose to consider candidate conditions within the context of multiplex platform technology, such as tandem mass spectrometry, which is capable of identifying many conditions simultaneously. Other professional groups had previously noted that multiplex testing was likely to reveal information about conditions that did not meet the Wilson and Jungner screening principles27. The ACMG evaluation also took into consideration the benefits of screening that may accrue to the family even if none accrued to the child being screened. Although family benefit as a justification for screening has been supported by one professional group statement,22 it has been rejected by others.4,5,27,29–34

The ACMG report recommended 29 primary targets for universal screening. It recognized 25 secondary conditions identifiable through multiplex testing on the core conditions, for which the natural history, the need for treatment, the duration of treatment, or the efficacy of treatment were insufficiently understood to recommend screening at the time of the report. Although many parent advocacy groups press for further expansion on the grounds that knowledge is beneficial to families,29 expanded screening carries potential harms. With every added condition, the frequency of false-positive results increases.35 Confirmatory
testing is likely to avert unneeded medical interventions, but other possible adverse effects include psychosocial and emotional distress and the potential distortion of parental perceptions about the child. Similarly, expanded screening may also give rise to “patients in waiting”: individuals with a genetic diagnosis who have no signs or symptoms and may remain asymptomatic for years or decades.

The uniform panel proposed by the ACMG study was endorsed by the US Department of Health and Human Services Secretary’s Advisory Committee on Heritable Disorders in Newborns and Children, and all states have adopted the panel despite some criticisms regarding its methodology. The Advisory Committee has since developed more stringent criteria for adding conditions to the uniform panel and in 2010 recommended the addition of severe combined immunodeficiency. Also needed is a process to delete conditions ultimately shown to be inappropriate for screening, although no conditions have been challenged, to date, in the United States.

The American Academy of Pediatrics (AAP) and ACMG support the mandatory offering of newborn screening for all children. Following education and counseling about the substantial benefits of newborn screening, its remote risks, and the next steps in the event of a positive screening result, parents should have the option of refusing the procedure, and an informed refusal should be respected.

A matter of ongoing debate is how best to involve parents in consent for newborn screening. Currently, most jurisdictions mandate newborn screening, with only Wyoming and the District of Columbia requiring active parental consent, although neither requires written consent. With the exception of Nebraska, all states allow parents to opt out, although they differ in what reasons parents may give for refusing. Given the number of conditions screened for, a consent process discussing each condition in detail is neither feasible nor desirable. Rather, studies consistently show that parents wish to be told why newborn screening is being performed, where they can obtain more information, and what they can expect in the event of an abnormal result. Studies show that most parents prefer to be informed about newborn screening during prenatal care, and a statement by the American College of Obstetricians and Gynecologists supports this practice.

Research on informed consent increasingly supports a model of shared decision making, with emphasis on conversation rather than documentation. The principle of respect for autonomy is operationalized in the process of informed consent. When the individual lacks decision-making capacity, as all newborn infants do, the principle typically is expanded to include surrogate permission for medical interventions. Legal principles of privacy and self-determination support considerable deference to child-rearing decisions made by parents or guardians, with state intervention generally confined to instances of abuse or neglect. Although virtually all physicians would agree that the benefits of newborn screening far outweigh the risks and harms, few would assert that refusing screening constitutes medical neglect, given that the likelihood of a true-positive screening result is quite low. Obtaining parental permission may also increase the likelihood that a parent who receives notice of a positive screening result will be better prepared to respond appropriately and in a timely manner.

Three main objections have been raised against requiring formal parental permission for newborn screening. One is that the benefits of early diagnosis and treatment so greatly outweigh the risks that the state is justified in requiring screening to preserve its own interest in healthy children. The second is that requiring parental permission will lead to a large number of refusals, leaving many children untreated and at risk of having a condition undiagnosed. The third is that formal consent will be superficial and time consuming. Although studies have shown high acceptance rates even when formal parental permission is required, the argument that the benefits greatly outweigh the harms loses some of its force as newborn screening expands to include conditions for which early diagnosis does not necessarily reduce morbidity.

The recommendation of the AAP and the ACMG that parental permission be obtained for newborn screening does not specify how permission should be obtained or documented. Traditionally, written informed consent has not been required, although there is recent support for written documentation of refusal. Given the importance of newborn screening to reduce morbidity and mortality, all parents should be offered newborn screening and all parents must be educated about the process and its purpose. The optimal nature of the consent and refusal process may emerge from different states electing different processes if systematic data about the strengths and weaknesses of the different methods are collected.

CARRIER SCREENING

The AAP and the ACMG do not support routine carrier testing or screening for recessive conditions when carrier status has no medical relevance during minority. This recommendation accords with previous statements supporting the future decisonal autonomy of the minor, who will be able to make an informed choice about testing once he or she reaches the age of majority. Possible harms of early testing include labeling, stigma or discrimination, and a potential for misunderstanding the distinction between carrier status and affected status. The history of sickle cell screening in the 1970s exemplifies the risk of harm when information is poorly or inaccurately conveyed. Physician and lay confusion about the meaning of test results led to employment and insurance discrimination, fear, and stigma, compromising not only the interests of tested
individuals but also the credibility of the medical community and public health services.77,66,67

Possible benefits of carrier screening and disclosure in childhood include potentially greater acceptance and integration of status into life plans, avoidance of the shock and resentment that may accrue when disclosure is delayed, and greater opportunity for parental guidance in appreciating the nature of the genetic challenge and available management options. A recent review of empirical data regarding the effects of carrier testing on children concluded that "current research provides little evidence of a significant negative or positive impact on an array of indicators of psychosocial well-being."68 However, the authors acknowledged that "the methodological approaches used in quantitative studies might have been inadequate to detect important effects on children's emotions, self-perception, and social well-being."68

As noted previously, newborn screening may identify carriers for recessive conditions, such as hemoglobinopathies, galactosemia, and cystic fibrosis. There is broad consensus that when carriers are identified in newborn screening, carrier status should be disclosed to the child’s parents or guardians.27 Disclosure is supported primarily by the argument that carrier status is information about the child that belongs to the child and the parents are the appropriate surrogates.27 Carrier information about the child also has implications for the parents and their own reproductive risks, which may be perceived as a benefit of disclosure.

The rationale against routine carrier screening of minors is altered when carrier status has potential medical implications during minority. For example, the National Collegiate Athletic Association now requires sickle cell trait screening of all Division I athletes, proof of prior testing, or a written waiver from the athlete before participation in its programs. The justification for this policy is that adolescents and young adults with sickle cell trait have been shown to be at increased risk of exercise-related splenic infarct or fatal exertional rhabdomyolysis.69 However, others argue against screening because risk can be prevented, or at least reduced, by modifying training for all athletes—a policy that would minimize morbidity and mortality without stigmatization.70,71 Carriers of other conditions have also been found to be at risk of health conditions, for example, 20–30% of female carriers of Duchenne muscular dystrophy develop cardiomyopathies72 and fragile X premutation carriers are at risk of primary ovarian insufficiency and fragile X tremor–ataxia syndrome.73 As more knowledge is gained about genetics and genetic conditions, it may be discovered that a continuum of expression is the norm rather than the exception, and the distinction between "carrier testing" and assessment for clinical purposes may diminish.

Carrier screening may also be appropriate for adolescents who are pregnant or considering reproduction. The legal authority for pregnant adolescents to make reproductive health-care decisions independent of their parents or guardians varies from state to state. Health-care providers should be aware of the regulations in their jurisdictions before proceeding with genetic assays in pregnancy during minority.

The AAP and the ACMG do not support school-based genetic screening or testing because the school setting raises concerns about whether the uptake is informed and voluntary, whether privacy and confidentiality are maintained, and whether appropriate genetic counseling is provided before and after testing.74

PREDICTIVE GENETIC TESTING

Predictive genetic testing can occur in many contexts and can refer to predictive testing of either a childhood-onset or adult-onset condition, a distinction not made in previous statements about the genetic testing of children.77 It may involve testing an asymptomatic male infant with a positive family history for Duchenne muscular dystrophy or testing a child for a mutation associated with an adult-onset cancer syndrome. The former is an example of “predictive” testing: the presence of the mutation will almost certainly give rise to clinical manifestations. The latter is an example of “predispositional” testing: most (not all) cancer genes are incompletely penetrant and may never become manifest. Most predictive genetic testing for adult-onset conditions is predispositional.

Early professional statements recommended that predictive genetic testing of minors be considered only if effective medical interventions were available to treat, prevent, or retard the course of the disease.4,30 Since then, more than two dozen additional national and international guidelines have concurred.74 These statements identified a number of possible benefits and harms of predictive genetic testing for adult-onset conditions.4,30 Medical benefits include the possibility of evolving therapeutic interventions, targeted surveillance, refinement of prognosis, and clarification of diagnosis. Medical harms include misdiagnosis to the extent that genotype does not correlate with phenotype, ambiguous results in which a specific phenotype cannot be predicted (e.g., incompletely penetrant Huntington disease with 36–39 CAG repeats), and use of ineffective or harmful preventive or therapeutic interventions. Psychosocial benefits include reduction of uncertainty and anxiety, the opportunity for psychological adjustment, the ability to make realistic life plans, and sharing the information with family members. Psychosocial harms include alteration of self-image, distortion of parental perception of the child, increased anxiety and guilt, altered expectation by self and others, familial stress related to identification of other at-risk family members, difficulty obtaining life and/or disability insurance, and the detection of misattributed parentage. Reproductive benefits include avoiding the birth of a child with genetic disease or having time to prepare for the birth of a child with genetic disease. Reproductive harms include changing family-planning decisions on the basis of social pressures.

Although scant empirical data existed when earlier statements were written, the concern for harms was paramount, and the general consensus was to discourage if not proscribe predictive genetic testing of minors for late-onset conditions. In the intervening decades, some empirical data have emerged.68 They suggest less harm than anticipated, with considerable resiliency and ability of minors to successfully incorporate these risks
into their self-concepts and life plans. However, these studies disproportionately represent white individuals of higher socio-economic status; the effects on lower educated and underserved populations are largely unknown.77

Data show that in hypothetical situations, adults express a high interest in genetic testing for cancer predisposition (84–98%); however, of those actually at risk, only half proceed with testing.78 Uptake of genetic testing is even lower for untreated conditions, such as Huntington disease, with <20% of at-risk individuals undergoing testing.79–82 Given the wide variation in preferences among adults, commentators argue that allowing parents or guardians to test their minor children unfairly preempts the future choices of those children. The AAP and the ACMG continue to support the traditional professional recommendation to defer genetic testing for late-onset conditions until adulthood. However, predictive genetic testing may be appropriate in limited circumstances.83 In deciding whether a child should undergo predictive genetic testing for late-onset conditions, the focus must be on the child’s medical best interest; however, parents and guardians may also consider the potential psychosocial benefits and harms to the child and the extended family.84 Extending consideration beyond the child’s medical best interest not only acknowledges the traditional deference given to parents about how they raise their children55–57 but also recognizes that the interest of a child is embedded in and dependent on the interests of the family unit. In some families, the psychosocial burden of ambiguity may be so great as to justify testing during childhood, particularly when parents and mature adolescents jointly express interest in proceeding. Some parents may seek predictive genetic testing for adult-onset conditions even when children are unable to participate in the decision-making process because of immaturity or cognitive impairment. After careful genetic counseling, it may be ethically acceptable to proceed with predictive genetic testing to resolve disabling parental anxiety or to support life-planning decisions that parents sincerely believe to be in the child’s best interest.83,85

Thorough genetic counseling before predictive testing is essential to ensure that parents, guardians, and maturing minors fully understand the limits of genetic knowledge and treatment capabilities as well as the potential for psychological harm, stigmatization, and discrimination.85,88,89 Characterizing predictive genetic testing as an elective medical procedure helps to frame issues of consent. In general, an elective medical procedure is conditioned on the child’s assent as well as parental permission.87 If an adolescent is not interested in testing, and the clinical benefits of knowing will not be relevant for years to decades, the adolescent’s dissent should be final. If a minor is young or immature, delaying testing until the minor can actively participate shows respect for the developing capacities of the maturing minor.

Adolescents occasionally seek predictive genetic testing without parental involvement. Health-care providers should be cautious about providing such testing to minors without the collaboration of their parents. Results may disclose information about parental status, thus compromising parental privacy. In addition, data show that adults have difficulty understanding the full implications of genetic information, and they often involve other adults in their decision making.88–91 Permitting an adolescent to make similar choices without the benefit of parental guidance is problematic. It is also not clear by what authority minors would be able to avoid parental participation. Although some genetic testing may be relevant to pregnancy management for adolescents, predictive information about adult-onset conditions does not often enter into prenatal decision making.92,93 Predictive genetic testing does not typically affect management of emergency medical conditions. If clinical benefits will not accrue for years to decades, testing should be deferred until adulthood or should require parent or guardian permission as well as adolescent assent.87

In the case of predictive testing for childhood-onset conditions or conditions for which childhood interventions will ameliorate future harm, the balance of interest may be altered in favor of early testing (e.g., children at risk of familial adenomatous polyposis are tested before adolescence to determine whether annual colonoscopy should be initiated or if there is a risk of hepatoblastoma and a need to initiate ultrasonographic screening). In such cases, parental authority to determine medical treatment supersedes the minor’s preferences with regard to liberty and privacy.

Significant deference should be extended to parents regarding the timing of predictive genetic testing for childhood-onset conditions. Parents may want this information early for life-planning purposes, such as choosing an appropriate dwelling in a community with appropriate services. Parents may also desire this information for reproductive planning, including the number and spacing of children, the use of assisted reproductive technology, or the use of prenatal diagnosis. Deferred testing is also acceptable as long as delayed diagnosis does not compromise the well-being of the child.

HISTOCOMPATIBILITY TESTING
The human leukocyte antigen (HLA) system comprises the major histocompatibility complex in humans. Genetic testing for HLA matching is most important for bone marrow and less important for solid organs. Because siblings have a 25% chance of being HLA identical, they are an important potential source of hematopoietic stem cells. HLA-matched siblings often are preferred as stem cell donors because of reduced risks of graft rejection, graft-versus-host disease, and other transplant-related complications compared with unrelated donors. Although HLA testing serves no clinical benefit for the child per se, the AAP and the ACMG believe that tissue compatibility testing of minors of all ages for stem cell donation for stem cell donation is permissible to benefit immediate family members.94 Such testing should be conducted only after thorough exploration of the psychosocial, emotional, and physical implications for the minor serving as a potential donor.94 Previous AAP statements support the participation of a donor advocate for all minors when tissue or, more rarely, when solid organ donation is contemplated.94,95
The role of HLA testing on embryos as part of preimplantation genetic diagnosis and in vitro fertilization goes beyond the scope of this technical report and accompanying policy statement.

**ADOPTION**

Some individuals planning adoption request preadoption genetic testing of the child. A joint policy statement of the American Society of Human Genetics and the ACMG issued in 2000 concluded that, in general, the same guidelines developed for children in biological families should apply for adopted children and children awaiting placement for adoption.96 However, a Working Party of the Clinical Genetics Society (United Kingdom) asked “whether there are particular considerations that might justify the genetic testing of a child being considered for adoption...”96 The Working Party noted that, although most adoptive parents are involuntarily infertile, some seek adoption to avoid a particular heritable condition. Their personal experiences are likely to affect their views on genetic conditions in a prospective adoptive child96 and warrant closer consideration. Clearly, prospective adoptive parents have an interest in obtaining as much family medical history as possible and should be informed of any clinical genetic concerns identified in a comprehensive physical and developmental examination. Whether prospective adoptive parents may undertake broader genetic investigation is more controversial.97-102 Admittedly, biological parents have substantial discretion in prenatal genetic testing, the results of which can inform decisions about postnatal management as well as pregnancy interruption. In the case of adoption, similar information may be requested to help an individual or couple decide whether they have the capacity and resources to raise a child with a specific known condition. However, testing may leave the child with a potentially stigmatizing diagnosis that interferes with permanent placement.99-102

Issues of genetic testing of children placed for adoption are even more complicated today, when many adoptions involve some degree of an open relationship with the child’s birth parents. Testing results may have implications for the extended family members of the birth parents, and such issues should be discussed as part of the process. If the decision is to allow for preadoption testing, consideration of disclosure to the extended family of the birth parents must be addressed prospectively. Likewise, some children are adopted by biological relatives (e.g., grandparents, aunts, and uncles). Testing in these families may also have far-ranging implications that need to be addressed before testing is performed.

An adoption agency may decide that it will give preference to individuals or couples who are willing to accept an at-risk child without testing, although prospective parents will require extensive anticipatory guidance. One could argue, however, that a child facing a serious medical prognosis is best served by having a condition diagnosed early and being placed with a family equipped to deal with the challenges ahead.96-98

The adopted child or child born of donor gametes has a medical interest in knowing his or her family genetic history.103 Despite debate about whether such knowledge is in the child’s overall best interest104 and whether the child’s interest always trumps other familial interests,104-106 there is a persistent call for access to genetic parentage information on reaching adulthood, even if donors or relinquishing parents were promised anonymity.104,107-110 For discussion about the role of the physician in facilitating communication about these issues, see the AAP statement on the topic.111

**DISCLOSURE**

Parents or guardians occasionally request that genetic test results not be disclosed to a child.112,113 Similar requests are sometimes encountered in conditions such as HIV and cancer.114-118 With respect to HIV, the AAP supports disclosure on the grounds that it improves self-esteem, ensures that the child learns in a positive environment, and promotes the child’s growing role in the health-care process.118,119 More generally, the AAP supports disclosure because children often sense that something is wrong, and nondisclosure may lead to feelings of abandonment.118

With respect to predictive genetic testing, a request for nondisclosure may indicate ambivalence on the part of the parent or guardian regarding the significance of the test results and, thus, a potential for harm either from the parent’s interpretation or from the child’s eventual discovery of the concealment.118 As the child matures, justifying such a request may become more difficult, even if the health-care provider agrees that disclosure might not promote the well-being of the child. The health-care provider should consider deferring testing if not clinically necessary, pending a detailed discussion of these issues.4

Ideally, health-care providers and parents or guardians should address disclosure issues before genetic testing. The adolescent candidate for testing ordinarily should be a participant in these discussions and should concur with the proposed plans. During pretest genetic counseling, parents should be made aware that the genetic information belongs to the child being tested and that they should be prepared to share the results with the child either at the time of testing or at a defined time in the future.4

If genetic testing occurs before the request for nondisclosure to the child, the health-care provider may wish to defer a decision about disclosure until after the issues have been explored fully. Factors such as the age of the child, the need for medical interventions, and the need for the child to participate in therapeutic plans must be considered. The AAP and the ACMG believe that a request for the results of a genetic test by a mature adolescent should be given priority over his or her parents’ requests to conceal information. When a younger child is tested and the parents request that the provider not reveal results, the provider should engage the parents in an ongoing discussion about the benefits and harms of nondisclosure, the child’s interest in the information, and when and in what manner the results will be disclosed. Ideally, a system or plan would be developed such that when the child reaches adulthood, the individual could be made aware of the existence of the test results and be given the option to know the results, but how to achieve this in

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**ACMG POLICY STATEMENT**

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a fragmented health-care system is unclear, underscoring the importance of a medical home for every child.120

As noted previously, genetic testing may produce evidence that the biological relationships among family members are different from the stated relationships. Pretest counseling should alert parents or guardians to this possibility,27,65 thus providing an opportunity to forgo testing or seek alternatives to the proposed test to avert undesired consequences. Even when parentage is not the issue, genetic testing of an individual may have implications for other family members. Whether physicians or patients have a legal duty to warn extended family members about known genetic risks is unsettled. It is well established that the physician’s duty to preserve patient confidentiality may give way to a duty to report or take other action, including warning third parties, in the event of contamination or credible threats of serious violence. Genetic risk differs from infectious disease risk in its lack of temporal urgency, among other things.121–124 But the public policy interest in averting foreseeable harm has led some courts to find negligence on the parts of physicians who failed to make a timely genetic diagnosis or to alert family members about potential genetic risks.

When diagnosis of a childhood-onset genetic condition is negligently delayed, physicians may be held liable to the index patient for delayed therapeutic intervention125 and to parents for loss of reproductive options and the added costs of raising a subsequently born affected child.126–129

More recently, courts have been asked whether the diagnosis of a hereditary cancer syndrome in an adult creates duties to other family members. In parallel cases, the supreme courts of Florida and New Jersey held that the physician’s duty to recognize and disclose the heritable nature and familial implications of a cancer syndrome must be measured by the standard of care at the time of the original physician–patient encounter.130,131 Both appellate courts allowed relatives to go forward at the trial level to establish whether an oncologist had a duty, in the early 1960s, to disclose the heritable nature of familial adenomatous polyposis (Safer v. Estate of Pack, New Jersey) or an otolaryngologist had a duty, in the early 1990s, to disclose the heritable nature of medullary thyroid carcinoma (Pate v. Threlkel, Florida). Although the Florida court declared that any such duty would be satisfied by informing the patient of risks to other family members, the New Jersey court declined to foreclose a potential duty to warn a relative over the objections of a patient.130,131

Previous commentators have supported a limited warning to relatives at risk of serious harm that could be avoided by prompt action, when attempts to persuade the patient to disclose have failed.122–124 Others argue that few, if any, genetic risks are so urgent as to justify breach of confidentiality.125 Most conclude that continuing dialog with the patient, including offers to assist in disclosure, is the preferable course; however, a disclosure to third parties that is limited to the information required to take action is permissible.133 Disclosure may be warranted if the consequences of a delayed diagnosis are severe, interventions are available and time is of the essence, a risk is not likely to be appreciated without disclosure, or misinformation has been conveyed in the past.134

PATERNITY TESTING

Historically, in the United States, a child born to a married woman was presumed by law to be the biological child of her husband. With the advent of surrogate, donor gametes, and same-sex couples, the legal recognition of parentage has become more complex. Moreover, 40% of children in the United States are born to nonmarried parents.135 Typically, in the absence of marriage, paternity is legally established either by a written acknowledgment or by genetic testing. Genetic paternity testing may be performed to establish or challenge child support, custody, or visitation rulings. Although testing is best performed on a triad (mother–father–child), it is possible using samples from a dyad (father–child). Assays can be performed using blood, cheek swabs, or other tissue.

Misattributed paternity may be suspected, for example, when a person offers to be an organ donor for a close relative and an ABO or HLA mismatch is revealed.136,137 However, a finding suggestive of nonpaternity must be verified by formal testing.138 Mismatches at a single locus or chromosomal region are insufficient to establish nonparentage. Other possible explanations include sampling error, labeling error, interpretive error, new mutation, and uniparental disomy. The loci sampled during formal paternity testing are selected for the express purpose of providing a thorough and statistically reliable assessment of parentage.139,140

DIRECT-TO-CONSUMER GENETIC TESTING

Direct-to-consumer (DTC) companies currently offer a wide variety of genetic assays, including paternity testing, carrier testing, nutritional profiling, and multiplex panels claiming to assess risk of monogenic and complex multifactorial conditions.141 To avoid regulatory requirements, some are marketed not as medical assays but as “recreational” tests. Their sensitivity, specificity, accuracy, and interpretive reliability are difficult to assess. Although direct access to medical testing increases consumer autonomy and supports self-determination, it raises a number of questions.

In 2004, the ACMG issued a statement concluding that “due to the complexities of genetic testing and counseling, the self-ordering of genetic tests by patients over the telephone or the Internet, and their use of genetic “home testing” kits, is potentially harmful. Potential harms include inappropriate test utilization, misinterpretation of test results, lack of necessary follow-up, and other adverse consequences.”142

In 2010, the US General Accounting Office reported the results of its investigation on DTC testing to the Subcommittee on Oversight and Investigations, Committee on Energy and Commerce, House of Representatives.143 The investigation uncovered evidence of unfounded genetic predictions, misleading test results, deceptive marketing, and other questionable practices.143
The effects of DTC genetic testing on children have not been evaluated by any professional body. Only a few DTC companies specify that their websites are not directed to children; regardless, there is no practicable way to prevent minors from engaging in DTC testing without parental knowledge. A survey of social networkers in the United States found that 6% of respondents had engaged in personal genome testing, and an additional 64% indicated an interest in doing so. Of these respondents, the majority were interested in carrier testing for someone other than themselves, including their progeny, even though less than half were confident that they understood the risks and benefits of personal genome testing.145

Tabor and Kelley146 suggest that DTC companies have a moral responsibility to educate parents about the risks of testing their children, to discourage testing of minors for rare genetic traits “particularly if they have no reason to be concerned about increased family risk,” and to provide genetic counseling to avoid misunderstandings.

The AAP and the ACMG strongly discourage the use of DTC and home-kit genetic testing of children. In addition to the risks of inaccurate results, inaccurate interpretations, potentially harmful interventions, and altered family dynamics are possible negative consequences.85,147 DTC testing raises issues of privacy, self-determination, and disclosure vis-à-vis parents and children, as discussed previously in the section on Disclosure. Professional involvement is indicated in any type of genetic testing on minors.

**GENETIC SERVICES**

Genetic counselors and medical geneticists are too few in number to take primary responsibility for managing all genetic testing and counseling.148–150 Primary and subspecialty pediatric care providers must have a working knowledge of the genetic issues likely to affect their patient populations150–152 and either have sufficient expertise to prepare families adequately before ordering genetic testing65 or refer such testing to a genetics professional.

The primary care provider often will serve as care coordinator and medical home for children with genetic conditions.52,153–155 Resources such as the AAP newborn screening fact sheet156 and the ACMG newborn screening action plans, known as ACT sheets,157 can help primary care providers interpret and manage genetic diagnoses in collaboration with a child’s metabolic or genetic specialist, hematologist, endocrinologist, neurologist, or infectious disease specialist. Greater familiarity with genetics is of special importance for physicians serving minority patients, who are less likely to be appropriately referred for genetic testing and counseling.158 The AAP and the ACMG support the expansion of educational opportunities in human genetics for medical students, residents, and practicing physicians and the expansion of training programs for genetics professionals.64

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DISCLOSURE
All authors have filed conflict of interest statements with the American Academy of Pediatrics. Any conflicts have been resolved through a process approved by the board of directors. Masamichi Ito, a member of the SELI Committee, works for Athena Diagnostics, a company that performs genetic testing in minors.

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