Detection of a 22q11.2 Deletion in Cardiac Patients Suggests a Risk for Velopharyngeal Incompetence

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ABSTRACT. Objective. Conotruncal cardiac anomalies frequently occur in patients with DiGeorge or velocardiofacial syndrome. Additionally, these patients may have overt or submucousal cleft palate, as well as velopharyngeal incompetence (VPI). Previous studies have demonstrated that the majority of these patients have a submicroscopic deletion of chromosome 22q11.2. We hypothesized that a subpopulation of newborns and children with congenital heart defects caused by a 22q11.2 deletion are at a high risk for having unrecognized palatal abnormalities. Therefore, we proposed to evaluate a cohort of patients with conotruncal cardiac malformations associated with a 22q11.2 deletion to determine the frequency of palatal abnormalities.

Methods. We identified 14 deletion-positive patients with congenital cardiac defects who had no overt cleft palate. Of the 14 patients evaluated for the 22q11.2 deletion, 8 patients were recruited from a previous study looking for deletions among patients with isolated conotruncal cardiac anomalies. Informed consent was obtained in these cases. The remaining patients had the deletion study on a clinical basis, ie, conotruncal cardiac defect and an absent thymus, immunodeficiency, or minor dysmorphia appreciated by the clinical geneticist. These patients were evaluated by a plastic surgeon and speech pathologist looking for more subtle palatal anomalies such as a submucousal cleft palate, absence of the muscular uvuli, and VPI. Some patients underwent videofluoroscopy or nasendoscopy depending on their degree of symptoms and age. VPI was not ruled out until objective evaluation by a speech pathologist and plastic surgeon was obtained. In addition, the child had to be old enough to provide an adequate speech sample.

Results. Of the 14 patients evaluated, 6 patients older than 1 year were found to have VPI. It is noteworthy that 3 of these patients were older than 5 years and had remained unrecognized until this study. The remaining 6 patients had inconclusive studies based on their age (younger than 26 months) and their inability to participate in adequate speech evaluations. Two of these patients, however, had histories of nasal regurgitation suggesting VPI and, in addition, had incomplete closure of the velopharyngeal mechanism during crying and swallowing observed during nasendoscopic examination—consistent with the diagnosis of VPI. Thus, 8 of 14 patients evaluated had evidence of VPI by history and examination. The remaining 6 patients will require further study when they are older before a definitive palatal diagnosis can be made.

Conclusions. A significant number of patients with a 22q11.2 deletion in a cardiac clinic may have unrecognized palatal problems. Recognition of such abnormalities will afford patients the opportunity for intervention as needed, ie, speech therapy and/or surgical intervention. Notably, two of our patients with findings suggesting VPI were infants and will, therefore, be afforded the opportunity for close follow-up and early intervention. Furthermore, three school-aged children had palatal abnormalities that were unrecognized until this study. Thus, we recommend 22q11.2 deletion studies in patients with conotruncal cardiac malformations, followed by extensive palatal and speech evaluations when a deletion is present. Pediatrics 1997;99(5). URL: http://www.pediatrics.org/cgi/content/full/99/5/89; chromosome 22q11.2 deletion, DiGeorge syndrome, velocardiofacial syndrome, conotruncal anomaly face syndrome, Opitz G/BBB syndrome, conotruncal cardiac anomalies, cleft palate, velopharyngeal incompetence.

ABBREVIATIONS. DGS, DiGeorge syndrome; FISH, fluorescence in situ hybridization; VCFS, velocardiofacial syndrome; VPI, velopharyngeal incompetence.

Cardiac anomalies are often observed in association with other malformations and as a feature of well-defined genetic syndromes. Frequently, the cardiac abnormality is the first presenting sign of a genetic disorder and may remain the predominant medical problem in the child’s early years. Because of this, attention to or recognition of other problems may be overlooked. Diagnosis of a syndrome may not be made until later in life, if at all. In many cases, the early diagnosis of a syndrome is beneficial in guiding the total treatment of the child as well as in providing the family with accurate genetic counseling.

Conotruncal cardiac anomalies, including interrupted aortic arch type B, truncus arteriosus, and tetralogy of Fallot, are seen as part of DiGeorge syndrome (DGS). Additionally, some patients may have a conoventricular septal defect or a right-sided aortic arch with aberrant subclavian arteries.1 DGS is a developmental field defect of the third and fourth pharyngeal pouches, which often includes thymic and parathyroid gland aplasia or hypoplasia, mild facial dysmorphism, and palatal abnormalities.23 The

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etiology of DGS is heterogeneous. There have been reports of autosomal dominant, autosomal recessive, and X-linked inheritance as well as an association with maternal diabetes. In addition, cytogenetic abnormalities have been observed in patients with DGS. Many of the chromosomal abnormalities observed in affected children have been unbalanced translocations, which result in monosomy 22pter→q11. More recently, high-resolution cytogenetic analysis and molecular studies using DNA dosage analysis or fluorescence in situ hybridization (FISH) have demonstrated that the vast majority of patients with DGS have interstitial or submicroscopic deletions within chromosomal region 22q11.2. We have detected microdeletions of 22q11.2 in approximately 90% of patients with DGS referred for molecular analysis. Studies have shown that FISH using chromosome 22q11.2-specific probes is an efficient method for the detection of 22q11.2 deletions and can be recommended as an adjunct to routine cytogenetic analysis.

Conotruncal defects, predominantly tetralogy of Fallot and conoventricular septal defects, have also been seen in association with velocardiofacial syndrome (VCFS). VCFS has been described as an autosomal dominant disorder in which, in addition to having cardiac anomalies, patients may present with overt or submucousal cleft palate, velopharyngeal incompetence (VPI), facial dysmorphia, and learning disabilities. Overlap between the features of DGS and VCFS, including cardiac anomalies, cleft palate, hypocalcemia, and immunodeficiency, suggested a common etiology and pathogenesis.

Because facial dysmorphia is either not appreciated or evident in the newborn period, the diagnosis of DGS or VCFS is frequently dependent on the presence of other findings, such as hypocalcemia, immunodeficiency, and overt cleft palate in patients with conotruncal cardiac malformations. In the absence of these clues, the diagnosis may be difficult. Furthermore, a submucousal cleft palate or VPI without overt cleft palate is often unrecognized in the preverbal child and may remain undetected in the older child, particularly if there is a learning disability. Therefore, children who continue to have these unrecognized problems may remain untreated, missing the opportunity for early speech intervention and/or early surgical correction.

We hypothesized that a subpopulation of newborns and children with congenital heart defects and a 22q11.2 deletion are at high risk for having unrecognized palatal abnormalities. Therefore, we proposed to evaluate a cohort of patients with conotruncal cardiac malformations associated with a 22q11.2 deletion.
deletion to determine the frequency of palatal abnormalities.

METHODS

Patients with conotruncal cardiac anomalies underwent studies for the detection of a microdeletion of chromosome 22q11.2 (Fig 1) as previously described.²⁰ Children with an overt cleft palate were excluded from the present study. The first 14 cardiac patients with 22q11.2 deletions were referred for palatal evaluations (Table 1). Eight patients were recruited from a previous study looking for deletions among patients with isolated conotruncal cardiac anomalies.³² The remaining patients had the deletion study on a clinical basis, ie, conotruncal cardiac anomaly and an absent thymus, immunodeficiency, or minor dysmorphia appreciated by the clinical geneticist. These patients are also enrolled in a long-term prospective study to examine phenotype-genotype correlations.

Palatal evaluations were performed by a plastic surgeon and speech pathologist using standard history, physical examination, and speech evaluation. As indicated in Fig 2, patients underwent videofluoroscopy and/or nasendoscopy (Fig 3). VPI was not ruled out until objective evaluation by a speech pathologist and plastic surgeon was obtained. In addition, the child had to be old enough to provide an adequate speech sample.

RESULTS

Of the 14 patients evaluated, 7 were older than 1 year. Of these 7 patients, 6 were found to have VPI (Table 2). It is noteworthy that 3 of these patients, older than 5 years, were unrecognized until this study. One of the 3 patients subsequently has undergone a pharyngoplasty, whereas surgical intervention has been recommended for the other 2 patients. Although it is generally difficult to diagnose VPI in children younger than 1 year, 2 patients in this age group had histories of nasal regurgitation suggesting VPI. In addition, incomplete closure of the velopharyngeal mechanism during crying and swallowing was observed during nasendoscopic examination—consistent with the diagnosis of VPI. However, these 2 patients will still require complete speech evaluations before the initiation of treatment plans, which most likely will include surgical correction and speech therapy.

Patients younger than 26 months were considered too young to assess to our satisfaction, based on our inability to perform complete speech evaluations. Thus, a total of 8 of 14 patients evaluated had evidence of VPI by history and examination. The remaining 6 patients will require further study when they are older before a definitive palatal diagnosis can be made.

<table>
<thead>
<tr>
<th>Table 1. Cardiac Findings in 14 Patients With 22q11.2 Deletion Referred for Palatal Evaluation</th>
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<tbody>
<tr>
<td>Finding</td>
</tr>
<tr>
<td>Truncus arteriosus</td>
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<tr>
<td>Interrupted aortic arch</td>
</tr>
<tr>
<td>Tetralogy of Fallot</td>
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<tr>
<td>Ventricular septal defect</td>
</tr>
<tr>
<td>Right-sided aortic arch</td>
</tr>
<tr>
<td>Interrupted aortic arch with truncus arteriosus</td>
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<tr>
<th>Table 2. Results of Palatal Evaluations in 14 Patients With Cardiac Anomalies and 22q11.2 Deletion</th>
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<tr>
<td>Age, y</td>
</tr>
<tr>
<td>Present</td>
</tr>
<tr>
<td>&lt;1 (n = 7)</td>
</tr>
<tr>
<td>1–5 (n = 4)</td>
</tr>
<tr>
<td>&gt;5 (n = 3)</td>
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* Determined by history of nasal regurgitation and nasendoscopy, with full speech evaluation to follow.
Facial dysmorphia was discernible in some of the infants and was subtle, consisting of prominent ears, a bulbous nasal tip, and malar flatness, features seen with a 22q11.2 deletion (Fig 4). Dysmorphia was more apparent in some of the preschool-aged children (Fig 5) and school-aged children (Fig 6), in particular the bulbous nasal tip and prominent nasal root.

**DISCUSSION**

Studies by us and others have reported an association among DGS, VCFS, and chromosome 22q11.2 deletions. The commercial availability of a FISH assay for the detection of this deletion has made definitive diagnosis possible in most patients with clinical histories of DGS or VCFS. In addition, such studies may now be helpful in patients with conotruncal cardiac anomalies for the purpose of treatment and recurrence risk counseling. If a 22q11.2 deletion is detected, patients may be offered clinical evaluations before a problem becomes evident and, thus, may be offered early intervention. For example, it is noteworthy that two of our patients with findings suggesting VPI were infants and will, therefore, be afforded the opportunity for close follow-up and early intervention as indicated. These two patients, along with the other five in the younger age group, will provide us with unbiased prospective data to determine the efficacy of early recognition of and intervention for palatal abnormalities. Furthermore, it is of interest that the three school-aged children (older than 5 years) had unrecognized palatal abnormalities until this study.

With regard to recurrence risk, all 14 patients had de novo deletions of chromosome 22q11.2. Therefore, the recurrence risk to the parents of these 14 children is extremely small, assuming a very low but undefined risk for germ line mosaicism. The affected individuals have a 50% chance of passing the deletion chromosome 22 to their offspring. As adults, they will have the option of prenatal diagnosis, because this deletion may be detected in cultured cells obtained by either amniocentesis or chorionic villus sampling. In addition, some of the structural abnormalities associated with 22q11.2 deletions, including cleft palate and cardiac malformations, may be detected by prenatal ultrasonography and echocardiography.

In summary, this study indicates that there may be a significant number of patients with conotruncal cardiac anomalies, within the cardiac clinic, in whom unrecognized palatal problems exist. Deletion studies of patients with conotruncal malformations seem warranted, followed by extensive palatal and speech evaluation when a 22q11.2 deletion is present. Long-term prospective studies of these patients will be required to determine the outcome of early recognition and intervention.
Fig. 6. Three school-aged children with the 22q11.2 deletion have bulbous nasal tips and prominent nasal roots with varying degrees of ear malformations.

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