Further delineation of the G syndrome: a manageable genetic cause of infantile dysphagia

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SUMMARY Three families including five subjects with the G or Opitz-Frias syndrome are added to 23 published cases who had dysphagia; characteristics of the two affected relatives were added to 19 well documented published reports. The data from index cases support the concept of the G syndrome as a constellation of midline defects, which include hypertelorism or telecanthus (89%), oesophageal dysmotility (69%), laryngotracheal clefts (44%), cleft palate or bifid uvula (34%), heart defects (29%), hypospadias (100% of males), renal or ureteral anomalies (42%), and mental retardation (38%). Affected relatives, often identified by hypertelorism, dysphagia, or hypospadias, had a much lower incidence of associated defects and mental retardation. They provide a more rounded but still biased view of a syndrome compatible with normal intelligence and life span. The data do not support a highly characteristic face in the G syndrome, which discriminates it from the phenotypically similar BBB syndrome. The variable expressivity and five cases of male to male transmission observed in 18 families are consistent with autosomal dominant inheritance. Vigilance for the morphological characteristics of G syndrome in patients with dysphagia is underscored by the potential for normal development with appropriate intervention.

Since Illingworth's categorisation of infantile dysphagia into structural or neuromuscular causes, numerous examples of clefts or of palsies affecting the gastrointestinal tract have been described. A malformation syndrome combining problems in both categories was first described in 1969 by Opitz et al in four patients from the G family. Subsequent publications added more patients from the G family and a second J family, defining hypertelorism, laryngotracheo-oesophageal clefts, oesophageal dysmotility, and hypospadias as cardinal features of the G syndrome. Paediatric consideration of the G syndrome in the evaluation of infantile dysphagia is important, since oesophageal function improves with age and appropriate measures to prevent aspiration can result in normal lifespan and intelligence. Diagnosis of the syndrome also allows genetic counselling with anticipatory management of pregnancy and delivery.

At the time of this review, 26 children with the G or Opitz-Frias syndrome from 16 families have been reported, three of whom did not have the cardinal feature of dysphagia. The predominance of affected males (23 cases) may in part reflect ascertainment bias due to hypospadias, and several cases of male to male transmission suggest autosomal dominant inheritance. The existence of mildly affected family members is consistent with variable expressivity and implies that the syndrome may be much more common than the number of reported cases would suggest. In addition, the expanded spectrum of the G syndrome as seen in mildly affected patients makes its phenotypic separation from related conditions such as the BBB or hypertelorism-hypospadias syndrome less clear. To clarify these issues further, we report an additional five cases of the G syndrome in three families, and consider the pattern of anomalies in mildly affected family members. An approach to infants with dysphagia which considers the expanded phenotype of the G syndrome is discussed.

Case reports

CASE 1
This six year old white female (fig 1b) weighed 2640 g at birth; the 34 week gestation was complicated by polyhydramnios and premature rupture of mem-
costal margin and a normal sweat sodium level was obtained. The child improved rapidly with erythromycin therapy, but required many admissions to hospital during her first year for suspected pneumonia, failure to thrive, and developmental delay. At the age of one year, extensive evaluation disclosed the following findings: height 69 cm (5th centile), weight 6-72 kg (below the 3rd centile), head circumference 44-5 cm (25th centile), anterior fontanelle open (1×1 cm), haemangiomata over the left flank and left inner thigh, telecanthus (3·1 cm, 90th centile) with hypertelorism (5·2 cm, 95th centile), shallow nasal bridge, midline clefts in the hard palate (posterior 1 cm), cleft epiglottis, bifid uvula, liver 4 cm below the costal margin, and general hypotonia with increased lower extremity reflexes (3+). Laboratory findings included incoordination of the soft palate without evidence of gastro-oesophageal reflux (barium swallow), left upper lobe cavitation and presumed lung abscess (chest x ray and bronchoscopy), normal culture results and blood chemistries except for Staphylococcus aureus from the nares. normal liver function tests including coagulation times and α1 antitrypsin, normal serum immunoglobulins, complement, haemagglutinin, candida reactivity, auditory evoked response, serum amino acids, thyroid, and lactate, and an EEG showing diffuse slowing without paroxysmal features. The child responded to aggressive pulmonary therapy and multiple antibiotics and was discharged on Enfamil formula.

Multiple admissions for respiratory infection continued and the child was in hospital from the age of 16 months to 20 months for resection of the left upper lobe cavitation. A Hill-Nissen fundoplication and gastrostomy tube placement were performed for gastro-oesophageal reflux and oesophageal incoordination which were now apparent on barium swallow. The diagnosis of G syndrome was made and the child monitored for reflux by serial barium studies. Despite the absence of observable reflux after operation, the subsequent course included recurrent pneumonias from staphylococcus, pseudomonas, and candida, persistent left pneumatoceles, chronic otitis externa and interna, multiple staphylococcal cutaneous abscesses, pseudomembranous colitis from Clostridium difficile, failure to thrive, alopecia, and developmental delay. The gastrostomy was removed at the age of three years without subsequent reflux. Immune evaluation has shown normal T cell mitogens, neutrophil function, and staphylococcal killing. Turbinate biopsy suggested hypoactive cilia. Mild developmental delay may reflect over 50 hospital admissions during her six years. A family history (fig 1a) showed that the mother, who has mild hypertelorism and a high
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palate, required tube feeding because of swallowing
difficulties as a child.

CASE 2
This three month old white male (fig 2a, b) weighed
3 kg following an uncomplicated term gestation and
vaginal vertex delivery. The mother had two healthy
children by a previous marriage and there was no
history of dysphagia, hypospadias, or other birth
defects in the family. Severe cyanosis and respira-
tory distress after delivery prompted evaluation
which revealed tetralogy of Fallot with severe
pulmonic stenosis, hypoplastic pulmonic arteries,
and right sided aortic arch as documented by
echocardiography. Laboratory findings included
normal EEG, cranial ultrasound, and pneumogram,
severe oesophageal dysmotility with delayed gastric
emptying and gastro-oesophageal reflux, normal
testosterone response to HCG, and a 46,XY normal
male karyotype. The clinical course included severe
failure to thrive, developmental delay, and chronic
cardiopulmonary disease despite a left Blalock-
Taussig shunt, Nissen fundoplication, and gastro-
stomy. The patient died at the age of 10 months after
cardiorespiratory arrest due to probable but un-
documented sepsis. Necropsy findings included a
length of 61 cm (50th centile), weight 3-9 kg (50th
centile), head circumference 42 cm (50th centile).

FIG 2  Frontal (a) and lateral (b) view of case 2.
relative hypertelorism for a four month old with an interpupillary distance of 4.5 cm (50th centile for nine months), prominent forehead, prominent nasal bridge, low set ears, right choanal atresia, short neck, bilateral fifth finger clinodactyly, bilateral simian creases, seven of 10 arch dermatoglyphic patterns, ambiguous genitalia with small penis (1 cm), mild hypospadias, bilateral cryptorchidism, horseshoe kidney, marked underdevelopment of the white matter of the cerebral hemispheres and descending systems of the brain stem, bilateral rudimentary intra-abdominal testicles, normal cerebellum, and normal gallbladder. Brain histology showed reduced white matter throughout the cerebral hemispheres, cerebral peduncles, and cortico-

spinal tracts. There was no evidence of a demyelination process. The olfactory bulbs and tracts could not be recognised. There were normal ganglion cells in the oesophagus.

**CASE 3**

This 13 month old white male (fig 3b) weighed 2980 g at birth after a 36 week gestation complicated by hypertension and eclampsia. The mother was 24 years old with a previous eight week miscarriage. After caesarean section with Apgar scores of 9 and 9, cyanosis and tachypnoea were noted at the age of one hour and an oesophageal tube could not be passed. Tracheo-oesophageal fistula and oesophageal atresia were corrected surgically at five days and a feeding gastrostomy placed until two months of age. Two episodes of aspiration pneumonitis led to recognition of oesophageal stenosis, oesophageal incoordination, and gastro-oesophageal reflux; a Nissen fundoplication with gastrostomy was performed at three and a half months. Dilatation of the oesophagus was performed and the patient has fed orally since the age of five months with a single episode of severe aspiration occurring at 11 months.

Physical findings included a height of 79 cm (80th centile), weight 11.3 kg (80th centile), head circumference 48.5 cm (80th centile), shallow nasal bridge with intercanthal distance of 2.2 cm (80th centile), interpupillary distance 4.5 cm (90th centile), prominent forehead, highly arched palate, thin upper lip, small and simplified but normally posi-
tioned ears, normal palmar creases, mild hypospadias, and normal intellectual development. Echocardiography had identified a patent ductus arteriosus at the age of three months without left to right shunting.

A family history (fig 3a) showed that the father (fig 3c) had feeding problems with an ‘immature oesophagus’ during his youth. His formula was switched several times because of vomiting and persistent diarrhoea. As an adult he still gags easily and chokes on large pieces of food. His height is 180 cm (75th centile), weight 102 kg (97th centile), head circumference 59.5 cm (97th centile), interpupillary distance 6 cm (97th centile), intercanthal distance 3.1 cm (75th centile), and outer canthal distance 9 cm (75th centile). He has a highly arched palate but no hypospadias. As shown in the pedigree, a paternal brother died a ‘crib death’ at six weeks and a paternal nephew had surgically corrected pyloric stenosis. The couple has subsequently had a stillborn female fetus at 32 weeks’ gestation after CorGard® and Dyazide® treatment for hypertension. Fetal necropsy at the Teratology Unit, University of Michigan, revealed micrognathia, questionable small ears, and tapered fingertips: the tissue was too autolysed for karyotype analysis. Placental examination revealed haemorrhagic endovasculitis.

**Results**

The clinical manifestations of our cases 1 to 3 are summarised in the table with other reported patients with the G syndrome. The data from the definitive patients support a strong influence of the G syndrome gene upon diverse systems including the face (hypertelorism or telecanthus in 89%), respiratory tract (laryngotracheal clefts in 44%, aspiration pneumonitis in 85%), gastrointestinal tract (demonstrable oesophageal incoordination in 69% of those studied, swallowing difficulties in 81%), genitourinary tract (hypospadias in 100% of males, renal or ureteral anomalies in 42% of patients studied), and heart (29% assuming complete ascertainment). In addition to their typical clinical or family histories, our cases 1 to 3 each had more than six of the above nine features as evidence for the diagnosis of G syndrome.

**TABLE Manifestations of G syndrome patients and their relatives.**

<table>
<thead>
<tr>
<th>Clinical manifestations</th>
<th>Published cases</th>
<th>Present cases</th>
<th>Total</th>
<th>%</th>
<th>Relatives</th>
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<tr>
<td></td>
<td></td>
<td>1</td>
<td>2</td>
<td>3</td>
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</tr>
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<td></td>
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<td>+</td>
<td>+</td>
<td>25/29</td>
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<tr>
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<td>-</td>
<td>-</td>
<td>25/29</td>
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<tr>
<td>Gestation</td>
<td>Mean 38 wk</td>
<td>34</td>
<td>40</td>
<td>36</td>
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<td>Mean 29.61 kg</td>
<td>2640</td>
<td>3000</td>
<td>2980</td>
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<td>17/20</td>
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<td>12/27</td>
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<td>6/28</td>
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<tr>
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<td>+</td>
<td>+</td>
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<td>+</td>
<td>8/28</td>
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<tr>
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<td>6/18</td>
<td>+</td>
<td>-</td>
<td>+</td>
<td>8/21</td>
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</tbody>
</table>

* Numerator: number of patients affected. Denominator: number of patients living long enough to manifest feature and in whom feature specifically included or excluded.

†Includes cleft plate, uvula, tongue.

‡Includes cleft epiglottis, larynx, trachea, oesophagus.
The data do not support the existence of a highly characteristic facies for G syndrome patients as cited by Funderburk and Stewart as a means of distinguishing the condition from the BBB hypertelorism-hypospadias syndrome. Thus, small or narrow palpebral fissures were found in three of nine patients for which the feature was mentioned, mongoloid slant to the eyes in three of 10, antimongoloid slant to the eyes in four of 10, prominent nasal bridge in five of 11, shallow nasal bridge in six of 11, antverted nares in six of 12, epicantal folds in one of 12, central groove to the nasal tip in two of nine, and prominent occiput in three of eight. A prominent forehead and parietal bossing were present in all eight patients for which the feature was mentioned. While several of the six of 25 patients with mental retardation might have sustained secondary damage because of severe dysphagia, the presence of EEG abnormalities and the severity of case 2 suggests a still undefined incidence of congenital neurological disease. The cerebral hypomyelination shown at necropsy in case 2 needs additional confirmation for this to be considered a part of the syndrome. A normal karyotype has been documented in 12 cases.

Approximately 21 relatives of the previously and presently reported G syndrome cases have been described in sufficient detail to provide an estimate of the incidence of cardinal features. The predominance of affected female relatives (14/21), the lower incidence of swallowing difficulties (5/8), hypertelorism (16/20), and hypospadias (4/7), and the presence of only one patient with mild mental retardation gives a less biased view of a syndrome compatible with a relatively normal face and life span. This additional evidence for variable expressivity and the five reported cases of male to male transmission now available provide strong support for autosomal dominant inheritance of the G syndrome and necessitates revision of its listing as an X linked disorder.

Discussion

Opitz and Gilbert have discussed the vertebrate midline as a developmental field which seems particularly susceptible to morphogenetic variation. Defects of the G syndrome which have midline origin include hypertelorism, widow’s peak, bifurcated nasal tip, short lingual frenulum, clefts of the palate, tongue, uvula, larynx, trachea, and oesophagus, hypospadias, urethral duplication, bifid scrotum, and imperforate anus. Of interest in view of midline problems is the apparent immune defect in patient 1, with recurrent otitis and sinusitis reminiscent of Kartagener patients with situs inversus and immotile cilia.

The dominance of midline problems in the G syndrome explains its phenotypic overlap with a related developmental field defect, the BBB or telecanthus-hypospadias syndrome. While certain families with G syndrome have a face which is quite distinct from that of typical patients with the BBB syndrome, compilation of facial features does not support a definitive separation of these two conditions. Many G syndrome patients will have obvious hypertelorism with a flattened bridge of the nose, frontal and parietal bossing, prominent occiput, narrow palpebral fissures, antverted nares, malformed ears, and a highly arched or cleft palate. BBP patients are more likely to have cranial asymmetry, epicantal folds, strabismus, and a prominent nasal bridge without the prominent forehead or occiput or parietal eminences.

Perhaps more important than discrimination of related midline defect syndromes is an appreciation of the risks which these conditions confer upon neonatal feeding and development. Patients with the G syndrome may have diverse causes for dysphagia which range from oesophageal dysmotility to severe defects in closure of the embryonic laryngotracheal groove. These may be difficult to demonstrate even by special evaluation and the infant with hypertelorism, dysphagia, and particularly aspiration pneumonia should be regarded as a medical emergency until a safe means of feeding can be established. Urgent evaluations should include direct laryngoscopy under general anaesthesia by an experienced endoscopist in addition to oesophagrams with a water soluble contrast media. In addition to repair of laryngotracheal clefts, significant gastro-oesophageal reflux or oesophageal incoordination or both should prompt consideration of a Nissen-Hill fundoplication with feeding gastrostomy. Since aspiration of saliva may still occur after these measures in severely affected infants, cervical oesophagostomy should also be considered. Failure to institute timely management has severe and deadly results, as indicated by the nine of 26 infants in the table dying before the age of one year and the severe pulmonary problems in case 1. The improvement in oesophageal function with age and the normal mentality of most G syndrome patients provides strong incentives for aggressive management.

The information now available on 18 families with G syndrome is strongly suggestive of autosomal dominant inheritance with variable expressivity. The high number of affected males may reflect ascertainment bias due to hypospadias or greater severity with increased abortion in affected females.
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The predominance of female relatives of G syndrome cases as listed in the table would favor ascertainment bias as an explanation. The existence of apparently normal heterozygotes, along with the low rate (three of 26 cases at most) of spontaneous mutation, indicates both a higher incidence and lesser severity of the syndrome than reported cases would suggest. The latter fact should be considered during genetic counselling: even apparently normal couples, such as the parents of case 2, might have their pregnancies followed for fetal anomalies or polyhydramnios by ultrasound. Since as many as 59 to 82% of patients with tracheo-oesophageal fistula have gastro-oesophageal reflux after operative repair, more attention to the nature of laryngotracheal clefts and craniofacial features in this population may reveal a subset with the genetic and pulmonary risks characteristic of the G syndrome.

References


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