Congenital salivary fistula of an accessory parotid gland in Goldenhar syndrome

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Abstract

Objectives: We report two cases of congenital salivary fistula of an accessory parotid gland, and we discuss its occurrence in Goldenhar syndrome.

Methods: Two teenagers complained of a congenital cheek fistula with constant salivary discharge. Computed tomography fistulography and sialography were performed. The diagnosis of Goldenhar syndrome was established based on clinical and imaging findings. Previously reported cases are reviewed and the clinical and radiological features summarised.

Results: In these two patients, a salivary fistula of an accessory parotid gland was demonstrated on computed tomography fistulography, and did not communicate with Stensen’s duct. Deformity of Stensen’s duct and hypoplasia of the ipsilateral mandibular ramus were present. Tragal appendices have frequently been reported in such cases.

Conclusion: A congenital cheek salivary fistula of an accessory parotid gland should be considered indicative of Goldenhar syndrome.

Key words: Child; Goldenhar Syndrome; Parotid Gland, Abnormality; Fistula; Computed Tomography

Introduction

Goldenhar syndrome refers to a wide spectrum of congenital abnormalities in the head and neck which originate from developmental disturbance of the first and second branchial apparatus.1,2 Hypoplasia or aplasia of the zygoma, ear, parotid gland, mandible and masticatory muscles are usually present in this syndrome. Hemifacial microsomia often occurs secondary to the anomalies.3

Congenital salivary fistulae due to abnormal development of an accessory parotid gland have been documented in several articles.4–8 Their association with Goldenhar syndrome has not been fully studied. A congenital cheek fistula of an accessory parotid gland should be considered indicative of Goldenhar syndrome.

We present two such cases for further documentation.

Case report

Case one

A 16-year-old male was referred to our hospital because of a fistula on his left cheek, which had been discovered soon after birth (Figure 1a). Discharge of clear, serous fluid was reported by the patient, which significantly increased in quantity during eating. No purulent excretion had been noticed.

Physical examination showed a skin opening located approximately 2 cm lateral to the left commissure. No swelling or redness of the facial skin was found. Asymmetry of the lower face was noted, with the chin slightly shifted to the left (Figure 1a).

A spiral computed tomography (CT) scan, performed without contrast medium, revealed a cord-like soft tissue mass extending from the skin to the lateral side of the left masseter muscle. The left parotid gland was of relatively normal size (Figure 1c and 1d). Hypoplasia of the left masseter muscle and the left mandibular ramus was noted (Figure 1b and 1d). The left masseter muscle was separated from the ramus by an abnormal collection of fat tissue (Figure 1d).

Computed tomography fistulography was performed, involving injection of contrast medium (iopamidol, 370 mg I/ml) into the skin opening, and showed the soft tissue cord to be a fistulous tract ending in a mass of attenuated salivary tissue (Figure 1d and 1e). No contrast medium was seen within the left parotid gland. An X-ray sialogram

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showed a narrowed left Stensen’s duct (Figure 1c). A CT sialogram of the left parotid gland via the normal orifice showed leakage of the contrast medium outside Stensen’s duct (Figure 1f and 1g). No communication was found between the fistula and the Stensen’s duct system.

Resection of the fistulous tract and an accessory parotid gland involved an intraoral incision together with excision of an ellipse of skin containing the fistula orifice.

Histopathological examination showed that the surgical specimen contained abundant glandular tissue (Figure 2).

Thus, the cheek fistula was considered to have arisen from an accessory parotid gland. A diagnosis of Goldenhar syndrome was established, based on hypoplasia of the ipsilateral mandibular ramus and masseter muscle, Stensen’s duct abnormality, and a congenital salivary fistula from an accessory parotid gland.

**Case two**

A 12-year-old girl was referred to our hospital with a fistula on her right cheek, discovered after birth. The fistula constantly discharged clear liquid, more so while eating. No history of infection was reported by the patient.

Clinical examination revealed that the skin opening of the fistula was located approximately 1 cm lateral to the commissure on the right side (Figure 3a). No purulent secretion or skin erythema was found. Facial asymmetry was noted, with the chin shifted to the right (Figure 3a and 3c). Prominent deformity of the right tragus was also noted (Figure 3a).

Sialography of the right parotid gland showed a duplication deformity of Stensen’s duct (Figure 3d).

Computed tomography performed without contrast medium showed a cord-like mass extending from the skin, with a large area of attenuated salivary tissue lying anterior and lateral to the masseter muscle (Figure 3e). Computed tomography fistulography via...
the skin fistula orifice showed that this attenuated area corresponded with the fistulous tract (Figure 3f and 3g). Volume-rendered images showed hypoplasia of the right mandibular ramus and mandibular asymmetry. (d) Combined right parotid gland sialogram and fistulogram, showing duplication deformity of Stensen’s duct (red arrows; sialography) and opacification of the accessory parotid gland (yellow arrow; fistulography via the cheek fistula opening). (e) Axial computed tomography scan without contrast medium, showing a small mass (red arrow) located anterior and lateral to the masseter muscle; note also the blue and yellow arrows indicate the masseter and internal pterygoid muscle separately. (f), (g) Computed tomography fistulography scans showing the fistulous tract extending posteriorly from the skin opening to the lateral side of the masseter muscle (arrow in part f); the salivary mass shown in part (e) is enlarged and filled with contrast (arrow in part g).

FIG. 3
(a) Clinical photograph of case two, showing the fistula skin opening lateral to the right commissure of the lip (red arrow, with a small naevus below); note the ipsilateral tragal deformity (yellow arrow) and the chin shifted towards the right. (b), (c) Volume-rendered images showing hypoplasia of the right mandibular ramus and mandibular asymmetry. (d) Combined right parotid gland sialogram and fistulogram, showing duplication deformity of Stensen’s duct (red arrows; sialography) and opacification of the accessory parotid gland (yellow arrow; fistulography via the cheek fistula opening). (e) Axial computed tomography scan without contrast medium, showing a small mass (red arrow) located anterior and lateral to the masseter muscle; note also the blue and yellow arrows indicate the masseter and internal pterygoid muscle separately. (f), (g) Computed tomography fistulography scans showing the fistulous tract extending posteriorly from the skin opening to the lateral side of the masseter muscle (arrow in part f); the salivary mass shown in part (e) is enlarged and filled with contrast (arrow in part g).

Discussion
Approximately during the fourth week of embryonic development, the ectodermal lining of the stomodeum or primitive mouth gives rise to buds or branches which form solid cords with round ends, and which subsequently develop into ducts and acini. Accessory parotid glands are derived from a similar pattern of branching and glandular proliferation, arising anterior to and separate from the main parotid tube. Aberrant buds which lose their communication with the main parotid gland may give rise to an abnormal, separate, accessory parotid gland.

The normally developed accessory parotid gland is a flattened nodule of salivary tissue separated from the main parotid gland, which lies superficial to the masseter muscle and connects to Stensen’s duct via one or occasionally two (or more) ducts; such accessory parotid glands are present in approximately 21 per cent of healthy people. Congenital malformation and acquired tumours of accessory parotid glands have been documented. Fistulae deriving from abnormal development of an accessory parotid gland in patients with Goldenhar syndrome are rare.

Goldenhar syndrome occurs due to abnormal embryogenesis of the first and second branchial apparatus, structures which give rise to most of the important facial structures. The first branchial arch contributes to the formation of the maxilla, zygoma, temporal bone, mandible, malleus, incus and masticatory muscles. The second branchial arch gives rise to portions of the ossicles, the styloid process, the hyoid bone and the muscles of facial expression. The terms oculo-auriculo-vertebral syndrome and first and second branchial arch syndrome have also been used to refer to this condition.

In addition to the two cases presented, five cases of congenital salivary fistula of an accessory parotid gland have been previously documented (Table I). The prominent clinical sign in all cases was a punctate skin orifice lateral to the commissure of the lips, with
<table>
<thead>
<tr>
<th>Case no</th>
<th>Study</th>
<th>Sex</th>
<th>Age (y)</th>
<th>Ext opening site</th>
<th>Discovery time</th>
<th>Secretion</th>
<th>Other defects</th>
<th>Sialography</th>
<th>Fistulography</th>
<th>Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Present</td>
<td>M</td>
<td>16</td>
<td>L cheek (2 cm lat to commissure)</td>
<td>At birth</td>
<td>Clear, serous liquid; more during eating</td>
<td>Mandibular asymmetry (deviated to R) R tragal deformity; mandibular asymmetry (deviated to R)</td>
<td>Main duct fistula; no communication with cheek fistula (CT) Double main duct malformation; no communication with cheek fistula (X-ray plain film)</td>
<td>Extending from skin opening to ant part of parotid gland superficial to masseter (CT)</td>
<td>Surgical excision</td>
</tr>
<tr>
<td>2</td>
<td>Present</td>
<td>F</td>
<td>12</td>
<td>R cheek (1 cm lat to commissure)</td>
<td>At birth</td>
<td>Clear, serous liquid; more during eating</td>
<td>Skin tag on R tragus</td>
<td>Not done</td>
<td>Extending from R ant cheek skin opening to accessory gland ant to R parotid gland (CT)</td>
<td>Chemocauterisation with botulinum toxin</td>
</tr>
<tr>
<td>3</td>
<td>Hah et al.⁶</td>
<td>F</td>
<td>1</td>
<td>R cheek (1 cm lat to angle of mouth)</td>
<td>At birth</td>
<td>Clear, serous salivary discharge</td>
<td>Not done</td>
<td>Soft tissue opacification lat to masseter, thought to be ectopic parotid gland (CT) Opacification of Stensen’s duct; no communication with fistula (CT)</td>
<td>Soft tissue nodule of salivary gland tissue; no communication between fistula &amp; Stensen’s duct system (CT)</td>
<td>Surgical excision</td>
</tr>
<tr>
<td>4</td>
<td>Moon et al.⁵</td>
<td>F</td>
<td>5</td>
<td>R cheek (2 cm lat to commissure)</td>
<td>At birth</td>
<td>Clear, serous fluid; more during eating</td>
<td>Periaural appendix; ectopic parotid gland Periaural appendix</td>
<td>Soft tissue opacification lat to masseter, thought to be ectopic parotid gland (CT) Opacification of Stensen’s duct; no communication with fistula (CT)</td>
<td>Surgical excision</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>Gadodia et al.⁴</td>
<td>M</td>
<td>8</td>
<td>L cheek</td>
<td>At birth</td>
<td>Serous discharge; more during eating</td>
<td>None</td>
<td>Small accessory parotid duct &amp; gland lat to masster, which opacified with contrast (CT)</td>
<td>Ongoing follow up</td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>Yamasaki et al.⁶</td>
<td>M</td>
<td>4</td>
<td>12 mm post to L lip commissure</td>
<td>At birth</td>
<td>Salivary outflow from fistula opening; +ve for amylase</td>
<td>None</td>
<td>Lobulated L parotid gland in normal position (X-ray plain film)</td>
<td>Fistula distinct from &amp; approx parallel to Stensen’s duct, which converged within a glandular body like density separate from parotid gland (X-ray plain film)</td>
<td>Translocation of fistula to oral cavity (Delore’s method)</td>
</tr>
<tr>
<td>7</td>
<td>Zhao et al.⁷</td>
<td>F</td>
<td>14</td>
<td>Small pit in L cheek</td>
<td>At birth</td>
<td>Constant salivary flow; more during eating</td>
<td>Ipsilateral preauricular appendices</td>
<td>Normal L. parotid gland (X-ray plain film)</td>
<td>Tract led to salivary tissue; very close to but no communication with parotid gland (X-ray plain film)</td>
<td>Translocation of fistula to oral cavity; excision of preauricular appendices</td>
</tr>
</tbody>
</table>

*Patient refused surgery. No = number; y = years; ext = external; M = male; L = left; lat = lateral; CT = computed tomography; ant = anterior; F = female; R = right; +ve = positive; approx = approximately
salivary discharge. This condition usually causes only minor cosmetic and functional morbidity, and patients may delay their treatment until their teenage years. Although the diagnosis of Goldenhar syndrome had not been clearly presented in previous articles, tragal deformity\textsuperscript{4,5,8} and parotid gland malformations\textsuperscript{5} have been documented, which are considered important markers for Goldenhar syndrome.\textsuperscript{14}

Mandibular hypoplasia on the ipsilateral side was notable in the present two cases, and contributed to diagnosis. In our second patient, a duplication deformity of Stensen’s duct revealed developmental disorder of the main parotid gland duct system. Thus, we suggest that congenital salivary fistula should be included in the spectrum of disorders which make up Goldenhar syndrome, due to its origin from an abnormal accessory parotid gland.

Computed tomography assisted fistulography and sialography can identify abnormal duct systems and aid in both the diagnosis and clinical management of such patients. From our experience and our review of the literature, the fistulous tracts do not usually communicate with the main parotid gland system.

- Congenital salivary fistula from an accessory parotid gland has been described in Goldenhar syndrome
- This study reports two further cases
- This abnormality should be added to the description of this syndrome
- Such cases often have deformity of the tragus, Stensen’s duct, the main parotid gland and the mandibular ramus
- Computed tomography fistulography and sialography are valuable for diagnosis
- These patients’ fistulous tracts do not communicate with the Stensen’s duct system

As congenital salivary fistula of an accessory parotid gland is a very rare clinical entity, there is a paucity of data on its treatment. Surgical excision of the lesion should be radical to prevent recurrence, and should be performed with aesthetic considerations, as patients are usually children or adolescents. We recommend surgical excision of the accessory parotid gland, using an intra-oral approach, and of the fistula and skin orifice, via a small skin excision, in order to maximise both therapeutic and aesthetic outcomes.

Translocation of the fistula to the oral cavity using Delore’s method has also been documented: a tunnel is made through the cheek and the orifice is implanted into the oral buccal mucosa.\textsuperscript{5} The more conservative treatment of chemocauterisation with botulinum toxin has also been reported to have a satisfactory outcome.\textsuperscript{5}

References
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