CASE REPORT

Management of an unusual case of iatrogenic parotid sialocele using an infant feeding tube: a novel approach

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SUMMARY

Injuries to the parotid duct and parenchyma resulting in swelling in the cheek region commonly present in maxillofacial practice. Sialocele is a discrete collection of saliva in a subcutaneous cavity that can be iatrogenic, idiopathic or the result of post-traumatic injuries to the duct or parenchyma, presenting as a salivary pseudocyst or retention cyst. We present an unusual case of parotid sialocele following surgery for sialolithiasis which presented with progressive painless swelling in the preauricular region of 9 months duration. Treatment included surgical exploration under local anaesthesia and draining the cavity by keeping the opening patent using an infant feeding tube.

BACKGROUND

Sialocele or salivary pseudocyst is an uncommon complication following trauma or surgery involving the parotid gland.1,2 A sialocele is an extraglandular collection of saliva in a subcutaneous cavity following disruption of the parotid duct or parenchyma.3,4 Clinically, sialocele appears as an asymptomatic soft mobile swelling, occasionally fluctuant, which, if left untreated, may rupture and develop into a salivary fistula.5,6 Diagnosis can be made by fine needle aspiration cytology, sialography, contrast CT and MRI. The various treatment options recommended include repeated aspirations and compression dressings, antisyloplagues, parasympathetic denervation (tympanic neurectomy), percutaneous sclerotherapy, sialography, sialendoscopy, botulinum toxin7 and injection of hypertonic saline into the gland.8

CASE PRESENTATION

We report a case of iatrogenic parotid sialocele in a 47-year-old man who presented to our maxillofacial unit with a chief complaint of painless swelling over the right side of the face of 9 months duration (figure 1). The patient was apparently normal 1 year previously but started experiencing a dull pain in the right preauricular region which worsened after having food. Past dental history revealed that he was diagnosed with right parotid sialolithiasis 10 months earlier, for which he underwent surgery.

The sialolith was located inside the parotid duct but the dimensions were not known as the patient had undergone surgery at another specialty centre for oral and maxillofacial surgery. The sialolith was removed from the duct by an extraoral approach by making a skin incision parallel to the parotid duct and blunt dissection was performed to expose the duct. The sialolith was removed in piecemeal fashion. One month after surgery the patient observed swelling in the cheek region which was asymptomatic. It gradually increased in size, with occasional reduction followed by a further increase. The medical and family history was non-contributory. On examination there was a 3×4 cm diffuse non-tender swelling in the right preauricular region which was fluctuant. The skin over the swelling did not show any discolouration, erythema or purulent discharge but there was a scar which showed the incision placed during the previous surgery (figure 2); no associated lymphadenopathy was detected. Intraorally, the mucosa was normal without evidence of any infection (figure 3). A diagnosis of sialocele was made by ultrasonography and analysis of aspirate for salivary amylase.

Figure 1 Extraoral swelling in the preauricular region on the right side.
INVESTIGATIONS
- Fine needle aspiration cytology
- Ultrasonography
- Sialography
- Analysis of aspirate for salivary amylase

TREATMENT
As a primary modality of treatment, aspiration and compression dressings were used but the swelling reappeared after 2 weeks so surgical decompression was planned. After baseline investigations the patient underwent intraoral surgical exploration under local anaesthetic using lignocaine 2% with adrenaline 1:100 000. A horizontal incision was placed 1 cm below the ductal orifice and the site was carefully explored to reach the cavity (figure 4). Once the lining was incised, straw-coloured serous fluid was visible. The cavity was irrigated with saline and an infant feeding tube no. 6 was taken and both ends were cut to achieve a length of 4 cm. The tube was kept in the cavity and secured to the cavity ends by suturing it with 4/0 Vicryl sutures. The mucosal incision was closed after proper haemostasis, by keeping approximately 1 cm of the tube outside (figure 5). This created an intraoral fistula as one end of the tube remained within the sialocele while the other end drained saliva into the oral cavity. Follow-up revealed no signs of sialocele formation with scanty flow of saliva from the tube and the patient was discharged on the third postoperative day (figure 6).

OUTCOME AND FOLLOW-UP
The wound healed uneventfully without any complication and the tube was removed after 10 days once it had stopped draining saliva. The patient was followed up at weekly intervals for the first 2 months and then once a month for the next 6 months. No signs of recurrence were noted during the follow-up observation period.

DISCUSSION
Iatrogenic parotid sialocele is an uncommon complication usually seen after surgery in and around the parotid gland. The causes of iatrogenic parotid sialocele include injury to the main duct, accessory duct, parenchyma of the gland and stricture of the duct. A thorough knowledge of the anatomy of the parotid gland and its relationship with other structures in the cheek are
essential when treating the pathology. The diagnosis of this condition has become easier as a result of recent investigative methods such as biochemistry of the aspirate for salivary amylase, Doppler ultrasonography, contrast CT and MRI. Most sialoceles are managed conservatively by percutaneous needle aspiration. However, resistant cases are managed either by diverting the parotid secretions into the mouth by reconstruction of the duct or creation of a controlled internal fistula or by depressing the parotid secretions by duct ligation, injection of sclerosants, hypertonic saline and botulinum toxin. We believe this is the first case of iatrogenic parotid sialocele to be managed by using an infant feeding tube. The purpose of placing an indwelling tube in the cavity is to drain the sialocele completely, and it is thought that the sialocele resolves spontaneously after 4 weeks due to formation of granulation tissue around the margins of the salivary parenchyma to seal any further extravasation of saliva. The infant feeding tube used in this technique did not cause any untoward hypersensitivity reactions. However, long-term randomised studies are needed to prove the efficacy of this technique compared with other methods for the management of sialocele.

Learning points

▸ A sialocele is an extraglandular collection of saliva in a subcutaneous cavity following disruption of the parotid duct or parenchyma.

▸ A sialocele can be iatrogenic, idiopathic or caused by post-traumatic injuries to the duct or parenchyma.

▸ Clinically, a sialocele appears as an asymptomatic soft mobile swelling, occasionally fluctuant, which, if left untreated, may rupture and develop into a salivary fistula.

▸ Diagnosis can be made by fine needle aspiration cytology, sialography, contrast CT and MRI.

▸ Among various treatment modalities, the present case of iatrogenic sialocele was managed by a novel approach using an infant feeding tube.

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Contributors AK performed the surgical procedure; SPS followed up the case during recall visits; SC monitored the overall progress of the patient; BSN prepared the manuscript.

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