A 25-year-old clinically healthy female presented to the Oral Medicine Clinic at the University of the Western Cape, with a swelling on her lower lip. The swelling started following a knife injury to the affected area and was present for almost three years. No treatment was sought for the initial injury and the lesion has slowly increased over time. The lump was asymptomatic but the patient requested that it be removed for cosmetic reasons.

She had no significant medical history and her extra-oral examination revealed bilateral submandibular lymphadenopathy. Intra-orally, a 15x15mm exophytic soft tissue lesion was seen on the lower labial mucosa, across the midline. The lesion firm, broad based, non-tender on palpation and was the same colour as the surrounding mucosa (Figure 1).

In addition, a small scar was noted across the midline of the lesion possibly from her previously reported assault. She had poor oral hygiene with visible soft plaque and calculus deposits detected on all her molars as well as multiple carious lesions and missing teeth.

The palpable bilateral lymph nodes could be attributed to the multiple lower carious teeth. The differential diagnosis included a mucocele, salivary gland neoplasm, benign soft tissue neoplasm (based on rate of growth and clinical appearance), traumatic fibroma, or a swelling secondary to the initial trauma.

An excisional biopsy was performed under local anesthesia. A curvilinear incision was made superficially across the midline of the lesion. Thereafter, blunt dissection was performed to separate the lesion from the surrounding tissue. Following complete enucleation, primary closure was obtained using vicryl sutures.

Postoperative instructions were given and a follow up appointment scheduled one week later. The healing was uneventful at the follow up appointment and the patient was referred to the relevant departments for prophylaxis and management of the carious lesions.

The macroscopic examination revealed an oval, firm well-circumscribed mass measuring 13 x 8.5mm (Figure 2). Microscopic examination showed a well circumscribed, encapsulated neoplasm comprising bland spindle-shaped cells with hyperchromatic nuclei, exhibiting a palisaded arrangement surrounding acellular eosinophilic areas (Verocay bodies) (Figure 3). A predominant “Antoni A” pattern was demonstrated throughout; with, focal “Antoni B” areas also present (Figure 4).

Immunohistochemical staining with S100 protein showed nuclear and cytoplasmic positivity in the spindle-shaped cells, confirming the Schwannian origin of the neoplastic cells (Figure 5).
Schwannomas, also referred to as neurilemmoma or neurinoma, are rare encapsulated benign neural neoplasms of Schwann cell origin. Schwann cells form a thin barrier around each extracranial nerve fiber, and wrap larger fibers with an insulating membrane, forming the myelin sheath in order to enhance nerve conductance. First described by Verocay in 1910, these tumors were initially referred to as 'Neurinomas'. Thereafter, in 1935, the term 'Neurilemmoma' was proposed due to the nerve sheath elements associated with these tumors.

Schwannomas do not arise from cranial nerves I and II, because these nerves lack Schwann cells. Schwannomas arise in association with a nerve trunk and as it grows, displaces the nerve. Nerve impingement can become symptomatic. Approximately 25%-45% of all extracranial Schwannomas occur in the head and neck region with only 1% seen intra-orally. Intra-oral Schwannomas are rare, with the tongue being the most common site affected, followed by the palate, floor of the mouth, buccal mucosa and lip. Furthermore, Schwannomas of the lip is the rarest reported clinical entity and is therefore generally not included in the differential diagnosis of a lower lip swelling.

Schwannoma of the lip was first described in 1969 and since then only a few number of Schwannoma cases of the lip have been reported. Moreover, as the Schwannoma of the lip enlarges, it also causes obvious aesthetic disfigurement for the patient and can lead to emotional distress. On occasion, it can arise centrally within bone, most commonly in the posterior mandible with concomitant bone expansion with associated pain and paresthesia. The tumour can affect individuals of all age groups but is most commonly found in the 2nd and 3rd decades of life, with no gender predilection.

The aetiology remains unknown. Some causative factors such as rare genetic predisposition, spontaneous growth, external injury, and chronic inflammation have been postulated. Schwannomas are not usually associated with a history of trauma, even though post traumatic cases have been reported. Moreover, with the history of trauma in our case as well as the location, a Schwannoma was not considered as a differential diagnosis.

Clinically, Schwannomas are typically asymptomatic solitary, freely mobile, submucosal mass characterized by slow growth and a smooth surface. Clinical symptoms are variable depending upon the location, other symptoms may arise, such as dysphonia, nasal obstruction, dyspnea, dysphagia and oral pain.

Schwannomas are usually solitary lesions but infrequently may be multifocal. Multifocal lesions also occur in 1) multiple localised neurilemmomas; 2) in association with neurofibromas, in von Recklinghausen's
disease and in 3) Schwannomatosis, a non-hereditary disease characterized by multiple subcutaneous and intradermal Schwannomas along with variety of intra-crural tumours.16,21

Histologically Schwannomas are usually encapsulated and shows proliferation of spindle shaped-cells, which assume two different patterns. The hypercellular areas, so-called Antoni type A, consists of spindle shaped cells organised in a palisaded pattern usually around eosinophilic areas, forming the so-called Verocay bodies. The hypocellular areas, so-called Antoni type B, are less organised and the spindle cells are randomly arranged within a loose myxomatous stroma.2

The prognosis of Schwannoma is favourable and no malignant transformation has been reported, local recurrences are rare and this has been associated with incomplete enucleation of the tumour.1,10,16

The preoperative diagnosis of intra-oral Schwannomas are challenging. The most commonly applied imaging method is magnetic resonance imaging (MRI).8 MRI or computed tomography (CT) may be significant in outlining tumour margins, composition and identifying any associated tumour infiltration. These imaging techniques are useful guide for surgical mapping prior to biopsy, which is required to arrive at a definitive diagnosis.12,20

Surgical resection of Schwannomas located on the lips is difficult. The intricate neuronal innervation of the lower lip requires precision during surgical removal, to prevent damage. Minimal damage during excision may cause significant morbidity such as impaired speech, aspiration, dysarthria, dysphagia, and paresthesia.20 A diagnosis of lip Schwannomas of the lip is very rare and is often misdiagnosed in early lesions. A differential diagnosis of lip Schwannomas is often only regarded until a more progressive stage of growth and subsequent cosmetic disfigurement occurs. To our knowledge there have been 9 previous reports of Schwannomas arising in a previous site of trauma.7,13,15

Even though Schwannomas are not bluish in colour, a differential diagnoses of a mucocele should be included, as the colouration of a mucocele can vary depending on the lesion size, proximity to the surface and the elasticity of the overlying mucosa.16,18 Albeit rarely encountered, it is our recommendation that intra-oral Schwannomas be included in the differential diagnosis for asymptomatic, well-circumscribed nodules or masses of the lower lip.7,13,17,22

References