A 76-year-old male presented to the Outpatient Department of General Surgery at Sikkim Manipal Institute of Medical Sciences with complaints of a pedunculated mass on his right cheek since seven years. The base of the peduncle was surrounded by a bluish red skin lesion, present since 30 years. There was a disproportionate growth of the skin lesion, particularly at the upper one third of the right nasolabial fold since seven years that gradually assumed a pedunculated shape. He denied the lesion being present since birth. Medical history included involution with infection of some areas of the skin lesion in the past. The patient denied having any significant medical problems or any significant allergic history. He was not on any medications. Family history disclosed no similar lesion in the family tree and he belonged to a middle class family.

On physical examination, he was well nourished, oral temperature was 98.6°F (37.0°C), blood pressure was 140/70 mmHg, pulse regular with a rate of 70 beats per minute, and respiratory rate of 22 per minute. Systemic examination was within normal limits. A complete body examination revealed no similar lesion or the presence of any other skin lesion. Peripheral arterial pulses of both the upper and lower limbs were within normal limits.

On local examination (Figure 1 and 2), the pedunculated mass was noted to originate from the upper 1/3 of the right nasolabial fold with extension to the right side of the upper lip. This was a single mass, bluish red in color, globular in shape, the translucency was bluish red and auscultation revealed no bruits. The swelling was nontender and nonpulsatile.

All the routine laboratory investigations were within normal limits. Finally, a histopathological examination confirmed the diagnosis.

What is your diagnosis?
See the next page for diagnosis.
The term hemangioma refers to the common angiomatous tumor of infancy that exhibits rapid postnatal growth and slow regression during childhood. The history did not reveal any congenital or acquired etiological factors. The patient had no symptoms other than mild local discomfort and obvious cosmetic disfigurement.

Approximately 80% of hemangiomas grow as a single tumor and 20% proliferate in multiple sites. Hemangiomas are more common in females with a female to male ratio of 3 – 5:1. The incidence in Caucasian infants is 10 – 12% and is 22% in preterm infants who weigh less than 1000 gm. The incidence is lower in infants with darker skin. Prenatal associations include older maternal age, placenta previa and pre-eclampsia. Basic fibroblast growth factor (bFGF) and vascular endothelial growth factor (VEGF) messenger RNA’s are upregulated in proliferative hemangiomas. VEGF is localized predominantly in pericytes and endothelial cells during the proliferative phase. bFGF is found in endothelial cells in both the proliferative and early involutional phases. Other studies indicate that the angiogenic peptide, bFGF, is elevated in the urine of infants with proliferating hemangiomas.

Hemangiomas appear in the neonate, usually within the first two weeks of life. Deep subcutaneous or visceral hemangiomas may not manifest until two to three months of life. Approximately 30% to 40% of hemangiomas are nascent at birth, presenting as a premonitory cutaneous mark. The lesions are composed of proliferating blood vessels and although benign, have a potentially destructive character. Hemangiomas undergo a proliferative and an involution stage. Congenital hemangioma is a rare variant that grows in utero and presents completely formed at birth. They do not proliferate. Sometimes the infant presents with neonatal hemangiomatosus, which refers to an infant with multiple cutaneous hemangiomas. Such patients may have intrahepatic hemangiomas that can cause congestive heart failure, hepatomegaly and anemia. Hemangiomas may be associated with other underlying conditions; lumbosacral hemangiomas may be overlying an occult spinal dysraphism as a tethered cord, lipomeningocele and diastematomyelia. PHACES syndrome is the association of the following: P, Dandy–Walker or other cystic malformations in the posterior cranial fossa; H, large facial hemangioma; A, arterial abnormalities; C, cardiac defects; E, eye anomalies; and S, spinal cleft. Subglottic hemangioma should be suspected in an infant with a facial hemangioma who presents with dyspnea and stridor. Diffuse hemangioma of the perineum and the lower limb is also seen with urogenital and anorectal anomalies.

The differential diagnosis includes deep lymphatic or venous vascular malformations. The presence of the lesion at birth supports the diagnosis of a vascular malformation although congenital hemangiomas are observed at birth. Pyogenic granuloma is also confused with hemangioma. These typically arise in the central face, they are small (average diameter: 6.5 mm) and rarely appear before six months of age (mean age: 6.7 years). Pyogenic granulomas grow rapidly, erupt through the skin and form a stalk or pedicle. Epidermal breakdown and crusting are common along with recurrent bleeding. Other differential diagnoses include: kaposiform hemangioendothelioma, tufted angioma, myofibromatosis, and fibrosarcoma. Ultrasoundography or magnetic resonance imaging (MRI) can confirm the diagnosis when in doubt. The patient underwent surgical excision under local anesthesia. An elliptical incision was made around the base of the pedicle. The feeding vessel was transfixed, the mass excised and the skin was repaired by interrupted sutures. He was followed up at one month and then one year with no additional complaints.

Additionally, a number of other therapies, depending on individual cases are as follows: observation, local wound care, local pressure, local or systemic steroids, recombinant interferon alfa-2a, flash lamp pulsed dye laser, intralesional laser [using potassium-titanyl-phosphate (KTP) alone or in combination with Nd:YAG lasers], and carbon dioxide laser in addition to suprascapular catheterization and embolization.

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References