Lymphangiomas are rare congenital malformations of the lymphatic system that can appear at any age or site, but commonly seen in the cervical region and the majority are diagnosed at birth. Although benign, and commonly asymptomatic, they can sometimes cause pressure and life threatening complications especially massive lesions involving the neck and mediastinum in newborn infants or cause a diagnostic confusion as a result of sudden enlargement following hemorrhage or infection. This is a report of 22 infants and children with lymphangiomas at different sites, outlining aspects of diagnosis and management.

Methods. The medical records of all children with the diagnosis of either lymphangioma or cystic hygroma admitted to Qatif Central Hospital, Qatif, Kingdom of Saudi Arabia from August 1989 to July 2000 were retrospectively reviewed and the following informations were collected: age at diagnosis, gender, site of lymphangioma, method of treatment and outcome. The operative findings and histological features were obtained from the operative notes and histopathology report.

Results. Twenty-two children (12 females: 10 males) with the diagnosis of lymphangioma were treated surgically except 3 who were treated with intralesional bleomycin and showed complete disappearance of their lesions. There was recurrence in the child from mediastinal lymphangioma and a small recurrence in the child with bilateral lesions in the floor of the mouth.

Conclusion: Lymphangiomas are relatively rare, involving mainly the head and neck, but they can be rarely seen at other sites. An important observation is the sudden appearance of cervical lymphangioma as a result of hemorrhage, which should be kept in mind. Our experience in the treatment of lymphangiomas using bleomycin is limited to draw any conclusions. We therefore considered surgery as treatment of choice for lymphangiomas. However, sclerotherapy can be used when there is a risk of damaging surrounding structures, and also to obviate the poor cosmetic results.

treated at our hospital. Their ages at diagnosis ranged from birth to 12 years, but in the majority of them (73%), the diagnosis was made at 4 years of age or younger. In 10 (45.5%), the lymphangioma involved the neck (8 posterior triangle, one submandibular, one supraclavicular). Five of them without prior history presented with sudden appearance of neck swelling which caused diagnostic confusion. All 5 children were found to have sudden hemorrhage into a cervical lymphangioma (Figure 1) and the diagnosis was confirmed histologically. All the cervical lymphangiomas were treated surgically except in one who was treated with sclerotherapy (bleomycin 0.6 mg/kg). He required 2 injections of bleomycin with total disappearance of the lesion. One of our patients was seen as a newborn with multiple lymphangiomas involving the floor of the mouth, tongue and left parotid gland (Figure 2). She was treated conservatively and on follow-up, there was marked regression of the floor of the mouth lymphangioma except for a ranula swelling on the right side. This was treated surgically at the age of 3 years. It was opened, deroofed and marsupialized. The tongue lymphangioma decreased markedly and only multiple superficial cystic swellings were seen on the dorsum of the tongue at the age of 2.5 years. These were excised using diathermy. The left parotid lymphangioma increased in size, and she underwent excision, but there was a recurrence. This was subsequently treated with one injection of sclerotherapy using bleomycin, with total disappearance. Now she is 15-year-old with no evidence of recurrence. Two other patients with parotid lymphangioma were treated. One was a 9-year-old girl with left parotid lymphangioma treated surgically while the other was a 6-month-old female with right parotid lymphangioma treated with bleomycin sclerotherapy. Both of them showed complete disappearance of their lesions. The 2 children with parotid lymphangioma treated surgically were difficult to excise but fortunately, the lymphangiomas involved the superficial part of the gland only. Three patients had lymphangioma of the floor of the mouth. All were treated with marsupialization of the cysts. One of them had bilateral lesions. This patient developed a small recurrence on the right side and currently being followed up in the clinic. A 10-year-old girl presented with lymphangioma above the right knee, treated by excision, and a newborn with a large abdominal wall (lumbar region) lymphangioma is currently being followed up. She will subsequently require excision. Four of our patients presented with lymphangiomas at unusual sites including the mediastinum, the mesentery, left breast and scrotum. The child with scrotal lymphangioma was a 3-year-old boy who presented with right scrotal swelling that was diagnosed as a hydrocele. Intra
lymphangiomas. A 3-year-old female presented with chest pain and bulging of the right side of the chest, and chest x-ray revealed a soft tissue density on the right side with pleural effusion which was also confirmed by CT scan (Figure 3a and 3b). She underwent thoracotomy and excisional biopsy. This was followed by a subsequent thoracotomy and total excision of the tumor. Both biopsies proved this to be a lymphangioma. Postoperatively, she did well, but on follow up 15 months later, she was found to have a recurrence. The family was offered surgery but they refused. The other patient was also a 3-year-old male, who was admitted to the hospital due to abdominal pain and vomiting. Abdominal ultrasound and computerized tomography scan revealed a large multiseptate cystic swelling occupying most of the abdominal cavity. Laparotomy revealed a large cystic swelling arising from the mesentery of the small bowel and encroaching on a loop of ileum that was very adherent to the cyst wall. The cyst as well as this part of ileum were excised and histology proved it as a large lymphatic cyst.

Discussion. Most lymphangiomas, which are benign developmental malformations of the lymphatic system, occur in the head and neck with the cervical region being the commonest site. In 16 (72.7%) of our patients, lymphangiomas occurred in the head and neck, and in 10 (45.5%), the cervical region was affected. The majority of lymphangiomas being congenital were diagnosed at birth and over 90% were diagnosed by the fourth year of life. In our series, 73% were diagnosed by the age of 4 years. Lymphangiomas can however appear at any age. This is specially at sites other than the head and neck including the abdomen and mediastinum where they can attain a large size before being clinically apparent. This was the case in 2 of our patients. Sometimes they can attain a large size as a result of infection or hemorrhage. This can cause pressure symptoms or life-threatening complications at sites such as the mediastinum. An important observation we noted in 5 of our patients was the sudden appearance of cervical lymphangioma as a result of bleeding. This caused diagnostic confusion and the final diagnosis was confirmed only intraoperatively and by histology. The possibility of lymphangioma should always be considered in children with sudden appearance of a cervical swelling. The lymphangiomas are rare lesions, and their exact incidence is not known. Gupta reported an incidence of one case in approximately 4,000 live births. Nicholls et al. reported an incidence of one in 6,500 live births. The exact incidence in Saudi Arabia is not known. Kennedy has classified lymphangiomas into 5 groups as follows: (1) superficial cutaneous lymphangioma (lymphangioma simplex and lymphangioma circumscriptum), (2) cavernous lymphangioma, (3) cystic hygroma, (4) diffuse system lymphangioma and (5) mixed lymphangioma. Although lymphangiomas occur most commonly in the head and neck area, they can be seen rarely at other sites including the mediastinum (5), mesentery (6), scrotum (7), spleen (8), breast (9), retroperitoneum (10) and bones (11). Four of our patients had lymphangiomas at rare sites including the breast, scrotum, mediastinum and mesentery. Although rare, lymphangioma should be considered in the differential diagnosis of children presenting with cystic swellings at these unusual sites.

The treatment of lymphangiomas continues to be a challenge. Although spontaneous regression of lymphangiomas has been reported as in one of our patients, it is however rare. It may be reasonable to treat small asymptomatic lymphangiomas expectantly. The treatment of lymphangiomas is early surgery as this is technically easier before further invasion of normal tissue or scarring secondary to infection has occurred. This however needs to be a complete radical resection, as with an incomplete resection the danger of a recurrence with tendency to invasive growth. Although surgical excision has been considered the treatment of choice by most surgeons, sclerotherapy with OK-432 (produced from the low virulent strain of type 3, group A Streptococcus pyogenes) and bleomycin has gained popularity during recent years. Rantio et al. found OK-432 safe and effective in the treatment of lymphangiomas, and Claesson and Kuylenstierna proposed OK-432 to be the first choice of treatment of lymphangioma. Banieghbal and Davies found that macrocystic lymphangiomas respond almost universally to OK-432 injections, whereas patients with microcystic lesions generally do not respond and should not therefore be injected with OK-432. Hall et al. reported that OK-432 injection was effective in approximately one third of children with lymphangiomas and cysts larger than 5 cm are unlikely to respond to this therapy. They also advised against using this form of therapy for lesions outside the head and neck and for lymphangiomas surrounding the airways.

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treatment of choice for lymphangiomas. Sclerotherapy has a place in the treatment of lesions where there is risk of damaging surrounding structures as well as to obviate the poor cosmetic results. Sclerotherapy can also be used as an adjunctive therapy in the treatment of wide spread and incompletely excised lymphangiomas.

References