The ultrasonographic diagnosis of cystic cervical lesions: a pictorial essay

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Abstract

The importance of ultrasonographic (US) examination in the evaluation of the superficial structures of the head and neck region is increasing due to the widespread availability of modern equipment (such as high-frequency transducers) and modern techniques. In the case of a cervical lesion, ultrasound is usually the first imaging method used to assess the changes. First of all, US is capable of differentiating between cystic and solid structures, and based on the US appearance correlated with the age of the patient, location and growing pattern, the appropriate diagnosis is possible with a high accuracy. The aim of the present paper is to describe the US features of the most common cystic lesions encountered in the head and neck region.

Keywords: ultrasonography, cyst, congenital, head and neck region

Introduction

Ultrasoundography (US) represents the first imaging method used for the evaluation of nodular lesions in the head and neck. For achieving the appropriate diagnosis, the examiner should have very good knowledge regarding the anatomy, the clinical status of the patient and the ability to recognize pathological changes [1,2].

Cystic lesions are frequently encountered in the cervical region, being divided into two separate categories: congenital and acquired. The data obtained at the clinical exam and the age of the patients narrow the differential diagnosis [3]. The congenital cystic masses are characteristic for pediatric patients and young adults. On the other hand, a newly appeared cystic lesion encountered in an adult, should be regarded as suspicious for metastatic node with necrosis [4].

Harnsberger et al [5] describe a more complex classification of cervical cystic lesions, according to the age and the frequency of the findings, as shown in table I.

Table I. The classification of cervical cystic lesions, according to the age and the frequency of the findings [5, adapted].

<table>
<thead>
<tr>
<th>Frequency</th>
<th>Pediatric age</th>
<th>Adult age</th>
</tr>
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<tbody>
<tr>
<td>Common</td>
<td>Infected lymph nodes (abcess)</td>
<td>Metastatic necrotic nodes</td>
</tr>
<tr>
<td></td>
<td>Cervical abcesses</td>
<td>Cervical abcesses</td>
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<tr>
<td></td>
<td>Thyroglossal duct cyst</td>
<td>Infected lymph nodes (abcess)</td>
</tr>
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<td></td>
<td>Lymphatic malformations</td>
<td>Lymphatic malformations</td>
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<tr>
<td></td>
<td>Ranula</td>
<td>Ranula</td>
</tr>
<tr>
<td></td>
<td>Second branchial cleft cyst</td>
<td></td>
</tr>
<tr>
<td>Less common</td>
<td>First branchial cleft cyst</td>
<td>Thyroglossal duct cyst</td>
</tr>
<tr>
<td></td>
<td>Thymic cyst</td>
<td>Second branchial cleft cyst</td>
</tr>
<tr>
<td>Rare but</td>
<td>Dermoid/epidermoid cyst</td>
<td>Cystic schwannoma</td>
</tr>
<tr>
<td>important</td>
<td>Teratoma</td>
<td>Zenker diverticulum</td>
</tr>
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</table>
**Congenital cervical cystic masses**

*Thyroglossal duct cyst* represents the most common congenital lesions of the cervical region, representing approximately 90% of nonodontogenic congenital cysts [6,7]. The diagnosis is relatively easy, pointed by the presence of a cystic mass located on the anterior midline compartment of the neck [8]. The thyroglossal duct is a tubular structure that connects the thyroid gland with the tongue that is closing in the 8th-10th gestational week. The superior end of the duct is foramen cecum, and the inferior end will become the pyramidal lobe of the thyroid gland [9]. Thyroglossal duct cysts appear due to a persistence of the duct [10]. The duct is passing through the hyoid bone, these two structures being closely associated during ontogenesis [11]. In consequence the thyroglossal duct cyst will be found around the hyoid level (20-25% suprahyoid, 15-50% hyoid, 25-65% infrahyoid), usually on the midline or no more than 2 cm away from the midline [9,12] (fig 1) and very rarely in the floor of the mouth [13].

The diagnosis is frequently established during childhood or young adulthood, but still in 15% of cases the thyroglossal duct cysts are diagnosed after the age of 50 [9].

The US characteristic appearance of the thyroglossal duct cyst is a homogeneous, anechoic, thin walled lesion, with posterior enhancement, but this pattern is seen only in 42% of the cases [4,8]. Quite often these cysts may contain some internal debris or have a very heterogeneous appearance (fig 2), probably due to infection or hemorrhage. Sometimes the appearance may mimic a solid structure, more probably due to the proteinaceous content [3,4,14,15].

The acoustic enhancement is often difficult to depict due to gas in the neighboring pharynx and larynx. The cystic lesion may have thick walls (especially when infected). Usually the lesion can be compressed with the transducer and also displaced [4].

Identification of a solid component within the cyst raises the suspicion of a thyroid carcinoma, which may appear in 1% of the cases, 80% being of the papillary type [4,16]. The assessment of the thyroid gland is important because, very rarely, these cysts may contain the only functioning thyroid tissue [4].

**The branchial cleft cysts**

The term *cyst* is used when no communication is present between the inner pharyngeal mucosa and the outer cervical skin. *Sinus* anomalies have one communication open (at the pharyngeal mucosa or the skin) and mesenchymal tissue at the other end. The term *fistula* is used when both openings are present [17].

First branchial cleft cyst represents 5-8% of all branchial anomalies, more frequently encountered in middle-age females. They develop between the external auditory canal, the parotid and submandibular spaces [8]. Clinically, these cysts usually manifest as recurrent abscesses or inflammation around the ear or at the mandibular angle. Other symptoms may appear, such as otorhea (in case of communication between the cyst and the external auditory canal) and facial nerve paralysis [8,16]. Work [18] described two types: type I: derives from the ectoderm, being located medially to the concha of the ear and type II: derives from the ectoderm and mesoderm, containing skin appendages (sebaceous and sweat glands, hair follicles); it affects the external auditory canal and the cartilaginous pinna and includes very often parotid lesions [18,19].

The second branchial cleft cyst represents 95% of all branchial anomalies and occurs between 10-40 years of age, with no predilection for gender [8]. Bailey [20] classifies these anomalies in four types: type I – located superficially, along the anterior margin of the sternocleidomastoid muscle and deep to the platysma muscle; type II – the most...
common, along the anterior margin of the sternocleidomastoid muscle, lateral to the carotid space and posterior to the submandibular gland; type III – extends between the internal and external carotid arteries, lateral to the pharyngeal wall; type IV – situated in the pharyngeal mucosa.

Clinically these cysts are painless, fluctuant, slow-growing lateral neck masses. If infected, pain and tenderness may appear [8].

The third branchial cleft cyst is rare (3%) and is located in the posterior cervical space [8].

The fourth branchial cleft cyst represents approximately 1% of the cases. It extends from the pyriform sinus, passes through the thyrohyoid membrane and descends along the tracheoesophageal groove [3].

On US examination uninfected cysts are well defined, with thin walls, anechoic, homogeneous, with acoustic enhancement (fig 4).

Sometimes a pseudosolid appearance may be found, due to the proteinaceous content [21].

If infection occurs, the walls become thicker, irregular, the cyst is ill-defined and the internal structure is inhomogeneous, with debris and septa (fig 5-7) [21].

These features can mimic cystic nodal metastasis [22]. The diagnosis between these two entities is challenging, both for the clinician and the radiologist. In the adult population, cystic neck masses should be considered primarily as metastatic nodes until the benignity is proven [23, 24].

Lymphangiomas are congenital abnormalities arising from the lymphatic system, with an infiltrative pattern, representing around 5-6% of all benign global lesions in pediatric patients. Most of them (80-90%) are detected in the first 2 years of life [8, 9]. There are three major theories proposed in order to explain the development of lymphangiomas: the failure of the primordial lymphatic sac to drain into the veins, abnormal sequestration of lymphatic tissue, and abnormal budding of the lymphatics [9].

There are four histological types of these anomalies: cystic hygroma (lymphangioma – the most common), cavernous lymphangioma, capillary (simple) lymphangioma and vasculolympathic malformation (lymphangiohemangioma) [9].

Lymphangiomas are most frequently present in the cervical region (75%), especially in the posterior neck and oral cavity in children and in the sublingual, submandibular...
dibular and parotid spaces in adults. Uncommonly, they are also described in other sites, such as the mediastinum (5%), axilla (20%), abdominal cavity (in the colon, liver and spleen), retroperitoneum (in the kidneys), scrotum and, very rarely, in the skeletal system [25,26].

The patient present with a painless, soft tissue cervical mass. After trauma or when hemorrhage or infection occurs, usually a sudden enlargement of the mass is noticed. More rarely, symptoms such as facial nerve paralysis, dysphagia, or feeding difficulties have been described [27-29]. In very rare cases, if the mass is very large, an extension and compression of the airways may appear, leading to the death of the patient [27].

The US examination reveals a multiloculated cystic lesion, often extending across several anatomic compartments, with septa of different thickness (fig 8). If hemorrhage is associated, then fluid-fluid levels are seen [3].

**Teratomas, epidermoid cysts and dermoid cysts**

The term teratoma defines a lesion that is containing multiple tissues not normally found in the region where these neoplasms occur. Teratomas are very rare and approximately 10% of them affect the cervical area [30]. *Dermoid cysts* represent the most common form of teratomas. They contain squamous cell epithelium, a fibrous wall and skin appendages (such as sebaceous glands, hair follicles). By comparison, the *epidermoid cyst* contains only squamous cell epithelium with a fibrous wall and teratoid cyst tissue derived from all three germinal layers [31].

Dermoid cysts most commonly involve the orbit, followed by the oral cavity (especially the floor of the mouth, but may also appear in the lips, tongue or buccal mucosa) and the nasal region [32]. Clinically these lesions appear as painless slow-growing masses. During pregnancy they may rapidly enlarge [9]. At US, the epidermoid cyst usually appears as a simple cystic structure, but pseudosolid appearance is also possible. Meanwhile, the dermoid cyst, due to the presence fat, fluid and calcification (represented by the osseo-dental structures), produces more commonly a mixed appearance (fig 9,10). Sometimes, the dermoid cysts are difficult to differentiate from thyroglossal duct cysts [4,21].
Acquired cervical cystic masses

*Ranula* is a cystic lesion that occurs in the sublingual gland, secondary to the obstruction of the sublingual duct. They are divided into simple ranulas (located on the sublingual space) and “plunging” or “diving” ranulas (located in the sublingual and submandibular space, appearing secondary to the rupture of the wall of the simple ranula) [4].

On US ranula appears as an anechoic, cystic lesion, unilocular, avascular, well delineated by thin walls (fig 11); in infections, the walls may become thicker [3].

Necrotic nodal masses

Cystic (liquefaction) necrosis appears as a fluid area within an enlarged lymph node. It is highly suggestive for malignancy in cases of known primary squamous cells [33] or papillary carcinoma [34, 35] (fig 12).

Very rarely, lymphomatous lymph nodes may present areas of cystic necrosis, more often encountered after treatment or in advanced stages [36] (fig 13).

Intranodal liquefaction can also be identified in benign conditions, such as tuberculosis (fig 14) or abscess formation (fig 15, 16) [37].
Doppler US shows no vessels inside the lesion.

Therefore, even if not specific for malignancy, this aspect should always be considered pathological [36].

Neurogenic tumors are more commonly located in the head and neck region in the carotid (related to the vagus nerve or sympathetic chain) or posterior cervical space (related to the spinal nerve or brachial plexus). Cystic areas within these tumors develop secondary to mucinous degeneration, hemorrhage or necrosis. Usually the tumors are well delineated, with a fusiform shape. When appearing in the carotid space, they are located posterior to the vessels (this feature helps in the differential diagnosis from paragangliomas which appear between the internal and external carotid arteries). Clinically, neurogenic tumors are asymptomatic and no suspicious lymphadenopathy is associated [3].

In conclusion, in cases of tumoral cervical lesions of unknown origin and etiology, ultrasonography represents the imaging method of choice, as it is able to differentiate between solid and cystic masses. The clinical examination and US appearance (location, extension, internal architecture, and vascularization) are extremely useful criteria for establishing the proper diagnosis, the optimal treatment and preoperative planning of cystic lesions in the head and neck region.

Conflict of interest: none

References


