Multicentric, Synchronous and Metachronous Liposarcoma

Liposarcomas (LS) are the most frequent type of soft tissue sarcoma in adults. Their etiopathogenesis is unknown and, although they can develop in any adipose tissue, their primary location shows a predilection for certain anatomical regions.

Occasionally, more than one synchronous or metachronous LS may present in a patient in non-visceral regions. The unusual peculiarity of the natural history of some LS has led us to ponder the possibility of a multifocal or multicentric origin (MLS). This has been and continues to be a question of controversy that is difficult to resolve because of the rarity of this entity. The main clinical interest lies in its therapeutic approach and prognosis.

We present a new case of synchronous and metachronous MLS. The patient was a 57-year-old male, with no prior medical history of interest, who came to our Emergency Department due to progressive abdominal distension and general malaise that had been developing over the previous 5 months. He reported a simultaneous thickening of the left calf region. Physical examination revealed that the abdomen was occupied by a large mass. Likewise, the lower left extremity was presented with an increased perimeter in the calf region, which was not painful but hard. CT confirmed an abdominal mass (Fig. 1) suggestive of LS and a tumor formation measuring 15 cm in diameter in the left leg that was displacing the calf muscles (Fig. 1). Needle aspiration cytologies of both tumors were positive for LS.

The patient underwent resection of the tumor in the calf region as well as the abdominal tumor, which weighed 4100 g and originated in the retroperitoneum. The pathology diagnosis of both tumors was myxoid liposarcoma. The surgical treatment was complemented with radiotherapy.

During the first six months of follow-up, the patient presented a myxoid LS in the right dorsolumbar region (Fig. 2), left spermatic cord (Fig. 2) and the left axillary, right scapular, subxiphoid, right inguinal, sub-mammary and right infraclavicular regions, which were treated surgically. One year later, there were recurrences in the dorsolumbar area and lower left leg, which were also removed. The cytogenetic study showed an XY karyotype with several chromosomal anomalies. The most frequent anomaly was the presence of a long-armed isochromosome of chromosome 7. Likewise, mutations were identified in chromosomes 1, 3, 6, 12, 16 and 19.

Three years after the initial intervention, 2 small nodules were identified in the left lung and one in the right, compatible with metastases. Several months later, recurrences were diagnosed in the abdomen and the left calf region, as well as new primary tumors in non-visceral regions whose tumor replication time was significantly shorter than that of the lung nodules. The patient died after 6 years follow-up.

Discussion

The first case of multifocal synchronous LS in non-visceral regions was reported in 1934 and received the name of “sarcomatosis”.2 Ten years later, Ackerman presented a new case and referred to another 4, differentiating this entity with the name of “multicentric liposarcoma” (MLS).3 Case reports of MLS are rare. In a review of the literature from 1992, 35 cases were referenced and another new case was presented.4 Since then, we have witnessed the publication of sporadic cases,5,6 only one of which was found within the Spanish literature.7

The type of MLS that is most commonly identified is myxoid and expresses the chromosomal translocation t(12;16), a fact that was confirmed in the case that we present. The rarity of MLS makes it difficult to clarify whether they are synchronous or metachronous multicentric primary tumors, or whether it is metastatic disease, which involves disseminated neoplastic disease. Classically, in favor of the “multicentric” type, there has been emphasis made to the fact that synchronous or metachronous MLS appear in anatomical regions that are not commonly metastasized, generally in adipose tissue, and sometimes symmetrically. In addition, these areas usually coincide with the common locations of single LS: retroperitoneum, lower extremities and pectoral and pelvic girdles or even the spermatic cord.1,7 Since systemic metastatic dissemination requires us to consider lung involvement, a relevant fact has been the definition of MLS as LS whose presentation is evident in at least 2 different sites before pulmonary metastases.7 In the case we present, the tumor replication time of the pulmonary M1 has been clearly different than that of the non-visceral tumors, which could support the hypothesis of early subclinical microscopic lung disease. Recently, a clonal relationship in these tumors has been established, thus proving a common, and therefore metastatic, origin.10 Nonetheless, this does not explain why the anatomical locations of the metastases are areas where lung metastases are not usually located.

REFERENCES


Fig. 2 – (A) Liposarcoma in the right dorsolumbar region; (B) liposarcoma in the left spermatic cord.
Hydrocele After Peritoneal Dialysis: Persistence of the Peritoneal-Vaginal Duct

Hidrocele tras diálisis peritoneal: Persistencia del conducto peritoneo-vaginal

Persistent peritoneal-vaginal duct can lead to the appearance of four pathologies, depending on the degree of its obliteration: inguinoscrotal hernia, communicating hydrocele, simple hydrocele or spermatic cord cyst. Likewise, it is associated with cryptorchidism and anomalies in the development of the epididymis.3

We present the case of a 40-year-old male patient with a history of renal failure secondary to renal polycystosis. He initiated a program of peritoneal dialysis, and we immediately observed passage of liquid to the right scrotal region. Physical examination ruled out the existence of an inguinal hernia. A scrotal/abdominal ultrasound showed evidence of a right communicating hydrocele. With the initial diagnosis of permeable embryonic vaginal process, we decided on surgery to perform its resection. The procedure was begun with a right inguinal incision. We proceeded with the dissection of the spermatic cord with individualization of its elements and the observed existence of the permeable peritoneal-vaginal duct that measured 0.5 cm in diameter (Fig. 1). The duct was resected with ligation and proximal dissection at the internal inguinal orifice and distal ligation and dissection at its scrotal end. Hernioplasty was carried out with a mesh plug in the internal inguinal ring. The patient was discharged and followed-up in the outpatient clinic without incident; peritoneal dialysis was re-initiated after one month, with no complications.

The formation of the peritoneal-vaginal duct is completed in the third month of gestation with evagination of the coelom on each side of the midline toward the scrotal fold. Between the third month and the end of gestation and directed by the fibers of the gubernaculum testis, the testicle descends from its primitive retroperitoneal location to the scrotal region. Afterwards, the vaginal process of the peritoneum is obliterated, forming a fibrous cord called the Cloquet ligament. The distal part of the duct becomes the tunica vaginalis of the testicle. In 40% of cases, this obliteration is completed in the last weeks of gestation. In the remaining 60%, it closes before the end of the first year of life.2

Fig. 1 – Intra-operative image demonstrating the elements of the cord and the peritoneal-vaginal duct.

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