INTRODUCTION
Hereditary leiomyomatosis and renal cell cancer (HLRCC) is thought to be a rare, autosomal dominant cancer syndrome of incomplete penetrance, caused by mutations of the fumarate hydratase gene (FH) located on the 1q42.3-q43 region.\(^1,2\)

Several synonyms for HLRCC exist such as: multiple cutaneous and uterine leiomyomas 1 with or without renal cell cancer (MCUL1), and Reed’s syndrome (cutaneous and uterine leiomyomatosis).\(^1,3\)

HLRCC variably predisposes to: 1) cutaneous leiomyomatosis; 2) uterine leiomyomatosis (rarely with uterine leiomyosarcoma); and 3) an elevated risk of malignant renal tumors.

CASE REPORT
Patient 1, a 34-year-old woman with no previously known diseases, was evaluated for a long course of multiple papules and small nodules of brownish pink colour on the anterior aspect of the upper body, painful either spontaneously or on exposure to cold environments (Figure 1). In the past she had two lesions excised, albeit non-cognizant of their histological examination.

She denied any similar lesions on her immediate family, except for her mother who was also observed (Figure 2). The 61-year-old mother had similar lesions on her trunk and upper limbs and reported a total hysterectomy for symptomatic uterine...
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myomatosis at age 38. They both recalled a history of uterine myomatosis on the maternal grandmother.

Dermatoscopically, the lesions had a thin brownish pink pigment network (Figure 3). Renal and adrenal gland ultrasound identified a simple cortical renal cyst with 20 mm diameter, and pelvic and endovaginal ultrasounds two uterine myomas with 10 and 6 mm in patient 1. Cutaneous biopsies of three lesions from patient 1 and two from the mother revealed Piloleiomyomata (Figure 4 and 5).

A clinical diagnosis of HLRCC was made, corroborated by the identification of a heterozygous variant on exon 5 of the Fumarate Hydratase gene (c.578C>T p.T193I).

The patient decided to excise the symptomatic lesions.

![Figure 1: Patient 1: Several brownish pink lesions on this patient’s chest. Note recurrence of two lesions adjacent to scars belonging to past excision of lesions](image1)

![Figure 2: Patient 2: Several lesions on the dorsum of the mother of the patient](image2)

![Figure 3: Patient 1: Brownish pink reticular pattern, with no other local features except follicular ostia (image taken with Sony Cybershot® DSC-T77 digital camera and Dermlite® II ProHr handheld dermoscope, 3Gen, LLC; original magnification × 10)](image3)

![Figure 4: Patient 1: Non encapsulated dermal tumor, with Grenz zone and consisting of interlaced fibers of smooth muscle cells (H&E × 40)](image4)

![Figure 5: Patient 1: Interlaced fibers of smooth muscle cells, with abundant eosinophilic cytoplasm and elongated nuclei with blunt ends (H&E × 100)](image5)
DISCUSSION

Although cases compatible with HLRCC already existed, the term Reed’s Syndrome appeared after Reed et al recognized tumoral predisposition in a family afflicted by multiple uterine (UL) and cutaneous leiomyomas (CL).1

Multiple heterozygous mutations of the FH gene were later identified, however with unknown implications in the generation of the multiple different phenotypes of this syndrome.4 In leiomyomatosis individuals, the presumed tumor suppressor activity of the fumarate hydratase is reduced.1,2,3

The most prominent phenotypical characteristic of HLRCC is the appearance of piloleiomyomas, with presumable origin in the arrector pili muscle. The number and size of these tumors can vary considerably, giving rise to cutaneous settings that vary from those that are practically unnoticeable to others that are severe and extensive.1,3 The skin-colored, red or brown papules or nodules occur in groups, scattered or both, on the limbs, trunk or face. Typically they cause pain on palpation or at low temperatures. There is a low risk for skin leiomyosarcomas.3

Dermoscopically, CLs have not been extensively described in the literature. Judging by what has been published, the findings are unspecific, varying between structureless homogeneous areas and a thin reticular pattern.5,6

The differential diagnosis should include: 1) from a clinical point of view, angiolipoma, glomus tumor and neurofibroma; and 2) histologically, smooth muscle hamartoma, dermatomyofibroma and dermatofibroma.7

ULs derived from the uterine smooth muscle cells are common benign tumors even in healthy women. In HLRCC, they are more frequent (79-100% of cases), larger (up to 10 cm), more numerous (from 1 to 20), and appear at an earlier age (average of 30 versus 40 years in sporadic cases).1 Apart from menstrual irregularity or associated pain, a growing number of infertility cases have been reported. The association with uterine leiomyosarcoma has been described as very rare.3,9

Although only 20-25% of patients apparently develop renal cell cancer (RCC), we prefer the term HLRCC in order to highlight the importance of this association.1,3 It is noteworthy that the majority of the RCC in these patients is exceptionally aggressive, with early metastasis even when smaller than 1 cm.1 Papillary RCC type 2 is the most frequent subtype, followed by collecting duct RCC. When present, RCC is typically solitary and unilateral.1,3,8 Renal cysts are also more prevalent, in 36% patients versus 4.6% - 8.3% of the general population.1

Although consensual criteria are lacking, according to those proposed by Smit et al., our patient met one major criterion (cutaneous leiomyomatosis) and one minor criterion (first-degree relative treated surgically for UL, under age 40).9 We also point to the presence of multiple small leiomyomata and a cortical renal cyst on our patient.

When the clinical and histopathological setting is suggestive, direct gene sequencing of the FH codifying region should be made. Piloleiomyomata rarely demand treatment except for cosmetic purposes or for pain relief, which in most cases is tolerable. In isolated tumors surgery is frequently resorted to. For pain relief in multiple lesions, isolated or combination therapy with nitroglycerin, adrenergic receptor blockers, analgesics, antidepressants, cryotherapy or CO2 laser may be used.3

In the follow-up and treatment of UL, the same gynecologic recommendations available for the general population apply. Signs that suggest malignant transformation may be screened for in gynecology.5

Currently there are no published guidelines for the screening of RCC in HLRCC, although the main therapeutic focus of this disease should be its early detection. When a FH germline mutation is identified, the patient and relatives should be subjected to adapted screening programs. Computerized tomography (CT) at age 18 seems to us to be a reasonable course of action, followed by ultrasound, or either CT or magnetic resonance for suspicious lesions. In the first case, the surveillance interval can be annual or every 2 years. Even though RCC is rare during childhood, this possibility should be considered in families with a history of RCC at a young age, with the screening test adapted to the child’s age.10

Dermatological evaluation of patients with HLRCC is mandatory. The histological confirmation of cutaneous tumors selected by their typical clinical appearance make the diagnosis easier, potentially leading to prompt cancer detection in affected patients and their families. Q
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