Hidradenitis suppurativa: literature review and case report

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ABSTRACT: Hidradenitis suppurativa (HS) is a chronic, recurrent and debilitating disease, affecting mainly women, especially in their second and third decades of life. Its most common incidence is in the axillary, inguinal, perianal and inframammary regions. Its complications include chronic fistulizing processes, with involvement of important adjacent structures, such as the sacrum and coccyx, the anal sphincter, urethra and great-caliber vessels, such as the groin vessels. The proportions of some cases of HS requiring extensive surgical procedures at several moments and the application of flaps and grafts, justify unusual cases reports, like this one. The authors present a case of extensive involvement of the perianal and gluteal regions, which required extended resection with flap in the first approach and fistulectomy in a second surgical moment, with good result for the patient.

Keywords: hidradenitis suppurativa; buttocks; surgical flaps.

INTRODUCTION

Hidradenitis suppurativa (HS) is a chronic, recurrent and debilitating disease, affecting mainly women, especially in their second and third decades of life¹–⁸. Its most common incidence is in the axillary, inguinal, perianal and inframammary regions³,⁹,⁷. Its cause has been attributed to apocrine duct occlusion caused by keratin plugs, involving some triggering factors – friction of adipose tissue, sudoresis, heat, stress, tight clothes – and some facilitating factors – genetic and hormonal factors¹,²,⁴,⁶,⁸,¹⁰,¹¹. Its chronic complications include chronic fistulizing processes, with involvement of important adjacent structures, such as the sacrum and coccyx, the anal sphincter, urethra and great-caliber vessels, such as the groin vessels¹⁰. The proportions of some cases of HS requiring extensive surgical procedures at several moments and the application of flaps and grafts, justify unusual cases reports, like this one².

Report a case of perianal and gluteal hidradenitis suppurativa treated with radical resection and rotation flaps.
CASE DESCRIPTION

Clinical examination: MAMN, female, 45 years old, with three years of purulent secretion drainage in the gluteal and perianal regions.

Proctologic examination: at the inspection, several fistulous orifices are observed, with palpable courses through the skin with purulent secretion drainage, located in the gluteal and perianal regions (Figure 1). No anomalies were detected with rectal touch examination and rectosigmoidoscopy.

Colonoscopy: performed until the terminal ileum, without alterations.

Intestinal flow: no alterations.

Blood exams: routine exams, as well as hepatic function tests, VHS and PCR, did not show alterations.

Diagnostic hypothesis: hidradenitis suppurativa.

Conduct: radical excision of skin and subcutaneous tissue affected by the disease.

First surgery: a radical excision of affected skin and subcutaneous tissue was performed using seton of a perianal fistula. The bleeding area reconstruction was performed using rotation of subcutaneous skin and muscular fascia flaps (Figures 2 and 3A and B). The final aspect of the surgery was according to expected patterns, with proper skin approximation (Figure 4).

Histopathology examination (HPE): the specimen HPE showed fistulous courses in the subcutaneous tissue, suggesting hidradenitis suppurativa (Figure 5).

Second surgery: the approach to the fistula channeled by seton in the first surgery was performed 40 days later, with excellent surgical result.

Patient control: the patient was examined two and four months after the second surgery, and at these two moments, the final aspect of the surgery was documented (Figures 6 and 7).

LITERATURE

Other names
Hidradenitis Suppurativa, Hidrosadenitis Suppurativa, Acne Inversa, Acne Conglobata, Verneuil’s Disease, Velpeau’s Disease1,3,12,13.

History
HS was first described by Velpeau (1839), who reported unusual processes related to the peculiar locations of axillary, mammary and perianal abscesses. For this reason, it was named “Velpeau’s disease”. After some time, Velpeau was not included in the references, despite his classical study of 183913.

However, that was Verneuil, through his several publications on the subject in the 1850’s, who most contributed to the knowledge of this disease, until then known as “Velpeau’s disease”. For this reason, “Verneuil’s disease” was also established, recognizing the importance of his publications12.
But the relation of HS with sweat glands was made only about 30 years after that, by Dubreuilh (1893), in his classical study of almost 120 years ago. Pollitzer, also in 1893, repeated the studies made by Dubreuilh, and the first case of HS was published on around 40 years after Brunsting (1939).

**General**

HS is a chronic, recurrent and debilitating disease, affecting mainly women, especially in their second and third decades of life. Its most common incidence is in the axillary, inguinal, perianal and inframammary regions. Its cause has been attributed to apocrine duct occlusion caused by keratin plugs, involving some triggering factors – friction of adipose tissue, sudoresis, heat, stress, tight clothes – and some facilitating factors – genetic and hormonal factors.

**Diagnosis**

The diagnosis is eminently clinical, based on symptoms reported by the patient and signs observed by the physician. The initial symptoms include discomfort, pruritus, erythema and hyperhidrosis in the affected area. With the disease progress, the symptoms are more evident. The physical examination detects lesions in the form of multiple diffused abscesses, with chronic drainage in the form of multiple fistulas through fistulous orifices of varied aspects. The skin and subcutaneous tissue in the affected area become inflamed and tender.

![Figure 3. Internal view (A) and external view (B) of the resected surgical specimen.](image)

![Figure 4. Panoramic view of the immediate result of the surgery, after using cutaneous flaps.](image)

![Figure 5. Panoramic view of a plate at the histopathology examination showing cryptic formations in the epithelium and dense conjunctive tissue with perifollicular inflammatory infiltrate.](image)
hardened, fixed and fibrotic, welding the dermis and epidermis layers together, keeping them from sliding apart.\textsuperscript{4,7,8,11} The clinical condition is characterized by long-term symptoms and signs that may reach 30 years of progress, with recurrent abscesses and fistulas, which, after the inflammatory phase, leave sequelae such as areas of fibrosis, fistulous orifices and scarce purulent secretion.\textsuperscript{4,7,8,11} When the disease affects the perianal area, it rarely involves the anal sphincter.\textsuperscript{8,10} The diagnosis, in the chronic form of the disease, is clinical and easily performed, depending on the coloproctologist’s experience. In certain situations, a biopsy is required to confirm the diagnosis, such as in the atypical cases of perianal Crohn’s disease, tuberculous ulcer and carcinoma. The association with spinocellular carcinoma, in case of long-term progress, is very rare.\textsuperscript{1,7,15,16}

The most important acute complication is characterized by an inflammatory and later infectious process, affecting superficial and deep tissues with cellulitis, abscesses and suppuration. Its chronic complications result from fistulas and the involvement of important structures, such as the sacrum and coccyx, the anal sphincter, urethra and great-caliber vessels, such as the groin vessels.\textsuperscript{1,4}

Complementary exams are required in the presence of extensive suppurrative areas, deep fistulas and involvement of noble structures. Bacterioscopy and culture of secretions, radiography of the sacrum and coccyx, fistulography and computed tomography of the pelvis can be performed.

The histopathology exam shows a cellular reaction into the lumen of apocrine sweat glands, with distention by leukocytes and cellular infiltration of adjacent conjunctive tissue. In the macroscopic perspective, the subcutaneous tissues have higher density, skin purple discoloration and fistulous orifices with little purulent secretion.\textsuperscript{4}

The secretion culture can isolate \textit{Streptococcus milleri}, \textit{Staphylococcus aureus}, anaerobic \textit{Streptococcus} and bacteroids.\textsuperscript{7,8}

**Differential diagnoses**

The diseases that should be considered and ruled out as differential diagnoses include (Figure 8): Crohn’s disease (Figure 8A), anorectal fistulas (Figure 8B), perianal fistulas (Figure 8C), cutaneous tuberculosis (Figure 8D), lymphogranuloma venereum (Figure 8E), pilonidal cyst, as well as other rarer diseases, such as anthrax, epidermoid cyst (infected dermoid cyst), erysipelas, furuncle, granuloma inguinale, steatocystoma multiplex and actinomycosis.\textsuperscript{5,7,8,15}

**Associated conditions**

The conditions that can be associated with hidradenitis suppurativa include: acanthosis nigricans, certain forms of arthritis, Crohn’s disease, Down’s syndrome, Graves’ disease, Hashimoto’s thyroiditis, herpes simplex, hyperandrogenism, irritable bowel syndrome and Sjögren’s syndrome.\textsuperscript{15}
Figure 8. Some morbid entities that should be considered as differential diagnoses of hidradenitis suppurativa: Crohn’s disease (A), multiple non-specific anorectal fistulas (B), non-specific perianal fistulas (C), cutaneous tuberculosis (D) and lymphogranuloma venereum (E).

Anogenital hidradenitis suppurativa

It affects more frequently the groin, involving the inguinal region, pubic region, internal face of the thigh and lateral scrotum. The perineum, buttocks and perianal folds are frequently included; the fistulas may deeply dissect in the tissue, involving the musculature, fascia and bowel. In the perianal form, biopsies are indicated to remove coexisting cancer and perianal
Crohn’s disease. It rarely affects the anal canal, but if involved, it is never above the pectineal line\textsuperscript{7,8,15}.

**Treatment**

Multiple treatment options are available, including antibioticotherapy, systemic retinoids, intralesional corticosteroids, hormonal therapy, immunosuppressors, radiotherapy, cryotherapy, local care of the lesion, laser therapy and surgical treatment. No isolated treatment was effective to the patients\textsuperscript{1-4,6-9,17}.

The surgical approach still seems to be the ideal option, and it may vary from simple incision and acute abscess drainage to radical excision of the whole tissue with apocrine glands. The radical excision of the whole affected tissue is the definitive and gold-standard treatment, as the recurrence rate is inversely proportional to the surgical radicality: the recurrence rate ranges from 100\% in three months (isolated drainage) to 25\% (cases of surgical radicality in 20 months after the radical excision)\textsuperscript{15,8,11}.

**CONCLUSION**

The approach used in the treatment of hidradenitis suppurativa remains challenging to physicians and frustrating to patients. As the number of randomized studies is insufficient, due to the lack of long series of cases, comparing the several types of treatment, the best approach is based on the patient’s clinical condition, results from prior nonsurgical treatments and the physician’s experience. The radical excision is considered the gold standard, and it should be the treatment of choice, since it is well indicated. Leaving the surgical wound resulting from enlarged resection of skin and subcutaneous tissue to heal at a second moment should not taken as the best alternative, although this is an option in certain patients; with flap rotations, when well indicated and performed, as an important factor in the immediate quality of life of the patient.

**REFERENCES**


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