Hidradenitis Suppurativa: The Third Cause of Vulva Carcinoma

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Abstract

The development of Squamous Cell Carcinoma (SCC) is a severe complication of chronic Hidradenitis Suppurativa (HS). Male patients with prolonged HS in extra-axillary areas are especially at risk of this complication. The case in our paper of HS associated with vulvar SCC emphasizes that women can also develop this complication. In addition to lichen sclerosus vulvae (via dVIN) and high risk HPV (via uVIN) there is a third disease that can lead to vulvar cancer, namely chronic HS. The clinician should look out for a malignant transformation in the presence of severe, chronic HS, and have a low threshold for biopsy. Staging, therapy and follow-up should be performed by gynecologic oncologists.

Keywords: Chronic hidradenitis suppurativa; Sclerosis vulvae; Vulvar cancer

Case Presentation

A 56-year-old woman, with a 30-year history of Hidradenitis Suppurativa (HS), presented herself to the emergency department with a painful growth on her right labium majus. A physical examination uncovered a 2 cm diameter verrucous tumor on an erythematous, enlarged, diffuse infiltrated labium majus on a background of HS Hurley stage III with scarring and sinus tract formation (Figure 1). An abscess was suspected in the context of a HS whereupon a general surgeon performed an incision and drainage. However, no pus oozed out and a skin biopsy was taken. Histopathology showed a Squamous Cell Carcinoma (SCC), but was high risk (hr) human papillomavirus (HPV) negative (Figure 2). The sentinel lymph nodes were free of disease. SCC is a rare but severe complication of HS. Malignant transformation in long-standing HS results from chronic inflammation, a phenomenon known as Marjolin’s Ulcer [1].

There is a lack of reliable incidence rate estimates of SCC arising in HS. Only 7 cases, including our patient, of HS associated vulvar SCC have been reported [2]. Although HS is three times more common in women than in men, SCC development in HS is observed more frequently in males (4:1 ratio) [3]. This can be explained by the fact that perineal/gluteal and perianal HS lesions have the highest predisposition to becoming malignant and these are the typical HS locations in men [1]. HS rarely occurs in these locations in women, but rather in the groin or the chest [4].

SCCs develop after 20-30 years of chronic HS [3]. Early recognition is important where the prognosis of vulvar cancer is associated with lymph node metastases in the groin and tumor size [5]. In HS, SCCs can be recognized by a persisting tumor or ulcer and the lack of typical HS features like sinus tracts and inflammation. 'High risk' (hr) human papillomavirus (HPV) can be identified in one-third of vulvar SCCs and is supposed to be a contributory factor of HS associated perianal/perineal SCC [6]. Studies showed that HPV types, especially HPV-16, are present in these carcinomas [3]. The presence of hrHPV was not investigated in the previously reported cases of HS associated vulvar SCCs. Our case suggests that hrHPV does not play a role in HS associated vulvar SCC.

Our patient was referred to the gynecologic oncologist who performed a sentinel node (SN) procedure and wide local excision. Histopathology showed negative SN and clear margins. No signs of recurrence were seen six months after the diagnosis. Staging, therapy and follow-up of vulvar SCC should be performed by gynecologic oncologists, according to the guidelines for vulvar cancer [7].
Conclusion

This case shows that, in addition to lichen sclerosus (via differentiated-type VIN) and high-risk HPV (via usual-type VIN), a third disease can progress to vulva carcinoma, namely chronic HS [8]. Therefore, clinicians should be vigilant for malignant transformation in HS and have a low threshold for biopsy.

References