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**Epidermolytic acanthoma: A case report**

Ginsberg AS *et al*. Epidermolytic acanthoma

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**Abstract**

BACKGROUND

Epidermolytic acanthoma (EA) is a rare benign skin lesion, usually found in the genital area of men and women, with epidermolytic hyperkeratosis as its distinguishing histologic characteristic. It is commonly misdiagnosed as condyloma accuminatum, verruca, and seborrheic keratosis. Since this lesion is benign, treatment is not necessary. However, it is often misdiagnosed, and patients are likely to undergo incorrect counseling and unnecessary treatment, causing undue burden to the patient. This study seeks to increase awareness of this rare condition to prevent future misdiagnoses.

CASE SUMMARY

A 55-year-old man living with human immunodeficiency virus presented for anal cancer screening. His physical examination revealed a flesh colored papule at the anal margin. The initial differential diagnosis included molluscum contagiosum, anal condyloma, and basal cell carcinoma. The lesion was excised to obtain a definitive diagnosis and was discovered to be EA.

CONCLUSION

EA is often misdiagnosed due to its similarity to other dermatologic conditions. Careful examination and pathologic evaluation should be obtained to ensure proper diagnosis.

**Key words:** Epidermolytic acanthoma; Epidermolytic hyperkeratosis; Condyloma accuminatum; Anal cancer; Verruca; Case report

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**Core tip:** Epidermolytic acanthomas (EAs) are papular skin lesions whose hallmark histologic feature is EHK. Herein, we describe a patient who presented with a solitary perianal EA and summarize the existing literature. EAs often present as pale papules in the genital region, trunk and extremities. EA is benign and cannot be spread to others, however they are commonly misdiagnosed as condyloma accuminatum, verruca, and seborrheic keratosis. Careful histopathological examination is necessary to obtain an accurate diagnosis and prevent unnecessary treatments. Although treatment of EA is not medically necessary, many patients prefer treatment, which can be accomplished with topical therapy or with excision.

**INTRODUCTION**

Epidermolytic acanthoma (EA) is a benign lesion, typically skin colored, brown, or white, small in size with a smooth surface, located in the groin and genital area, with a duration from one week to several years[1-4].The median age of diagnosis is 39 to 65 years[1,4-6]. The incidence is 4.5 cases per 100000 and most studies show a similar rate among men and women[4]. One study, however, reported that genital lesions were more common in males[4].  The lesions are not transmitted by skin-to-skin or sexual contact[7,8]. EA may present as a single lesion or in clusters[2-4,8]. Additionally, EA may be asymptomatic or pruritic[2,9].

The typical histologic features include well demarcated papular lesions with perinuclear vacuolization, reticular degeneration in the granular and upper spinous layers, and epidermolytic hyperkeratosis (EHK)[2,8].

In this case report, we describe one patient’s presentation, histologic findings, and prognosis. We also conducted a literature review of the condition. We performed a PubMed search of relevant literature on EA, using key words “Epidermolytic Acanthoma” resulting in 53 related publications. The most relevant articles based on title and abstract were chosen and reviewed. The purpose of this report is to contribute to a better understanding of EA and to encourage clinicians to consider this diagnosis when examining patients with small papular lesions.

**CASE PRESENTATION**

***Chief complaints***

The patient was a 55-year-old and was referred to our practice for anal cancer screening.

***History of present illness***

The patient had no known history of present illness.

***History of past illness***

The patient has been living with human immunodeficiency virus (HIV) for 17 years.

***Physical examination***

He had no gastrointestinal or anorectal symptoms and his physical examination was positive only for a three millimeter, smooth, flesh colored papule at the anal margin (Figure 1).

***Laboratory examinations***

His cluster of differentiation 4 (CD4) count was 2200 cells/mm3, with a nadir of 1300 cells/mm3. His HIV viral load was undetectable. Anal cytology collected at this visit was benign.

***Imaging examinations***

Digital anorectal examination and high resolution anoscopy (HRA) were otherwise normal.

***Preoperative diagnosis***

The differential diagnosis included molluscum contagiosum, anal condyloma, and basal cell carcinoma. To obtain a definitive diagnosis, the anal margin lesion was excised in the office, using local anesthetic.

***Histopathology and immunohistochemical findings***

Upon examination of the specimen, the lab determined the patient had EA.

**FINAL DIAGNOSIS**

The patient was diagnosed with EA.

**TREATMENT**

The anal lesion was fully excised during his initial office visit.

**OUTCOME AND FOLLOW-UP**

Histologically, the specimen was described as a hyperkeratotic papillomatous lesion, consistent with a diagnosis of epidermolytic acanthoma (Figure 2).  The excision site healed without complication.

In accordance with the New York State Department of Health guidelines, the patient will undergo annual anal cancer screening[10].This screening will consist of a digital anorectal examination, anal cytology, and anoscopy. If the cytology is abnormal, the patient will then undergo HRA with biopsy of any lesion that appears to be malignant or premalignant.

**DISCUSSION**

EAs are small lesions, often located on the trunk or genital area, and are commonly misdiagnosed[2].Therefore, close histopathological examination is important to achieve an accurate diagnosis[8]. The principal histologic feature of EA is EHK on 50% or more of the EA surface[2,4].EHK is defined by keratinocytes in the granular and spinous layers that have reticular degeneration, perinuclear vacuolation, coarse keratohyalin granules, and perinuclear eosinophilic inclusions[2,11,12].

While EHK is the hallmark histological finding in EA, it is not specific to this condition[13,14].  A correct EA diagnosis requires the characteristic histologic features in the proper clinical setting. EHK is also seen in epidermolytic ichthyosis (EI), epidermolytic epidermal nevus, or as an incidental finding[13,14]. These conditions are histologically indistinguishable. However, EI has distinguishing clinical features from EA, as it presents at or shortly after birth, and the skin of affected individuals appears red and blistering[15,16].  While EI is known to be caused by germline mutations in *KRT1*and *KRT10*genes, no such mutation has been found in EA[2,17,18]. However, reduced expression of cytokeratin (CK)1and CK10in EA has been observed on immunohistochemistry[17,19].

Many reports have confirmed the absence of human papillomavirus (HPV) in EA[2,20]. In one case report, however, a lesion demonstrated the presence of HPV type 16[[21]](https://word-edit.officeapps.live.com/we/wordeditorframe.aspx?new=1&ui=en%2DUS&rs=en%2DUS&wdorigin=OFFICECOM-WEB.START.NEW-INSTANT&wdenableroaming=1&mscc=1&wdodb=1&hid=C9A74B9F-607F-B000-0C6A-7D781DB01832&wopisrc=https%3A%2F%2Flasersurgerycare-my.sharepoint.com%2Fpersonal%2Ffrontdesk_lasersurgerycare_com%2F_vti_bin%2Fwopi.ashx%2Ffiles%2Fccf503163fbc4cc49bff199b5ed0af61&wdhostclicktime=1587666730670&jsapi=1&newsession=1&corrid=95ba930d-2488-405e-ba2e-9d405e5b0e3c&usid=95ba930d-2488-405e-ba2e-9d405e5b0e3c&wdredirectionreason=Unified_SingleFlush#_edn18). It is thought that in this case, HPV was an incidental finding rather than the causative agent for the EA[16].

While the patient described here was diagnosed with HIV 17 years before presenting with EA, his virus was well suppressed since diagnosis, and he always maintained high CD4 cell counts. To our knowledge, there are no published studies showing an association between HIV and EA. However, there have been case reports describing immunosuppressed patients who presented with EA. Two relevant case reports included two separate kidney transplant recipients, both developing EA with the commencement of immunosuppressive medications[7,22]. Further investigation is necessary to understand if EA is associated with immunosuppression.

EA is often misdiagnosed, and the most common misdiagnoses reported are condyloma acuminatum, verruca, and seborrheic keratosis[[2,4,23].](https://word-edit.officeapps.live.com/we/wordeditorframe.aspx?new=1&ui=en%2DUS&rs=en%2DUS&wdorigin=OFFICECOM-WEB.START.NEW-INSTANT&wdenableroaming=1&mscc=1&wdodb=1&hid=C9A74B9F-607F-B000-0C6A-7D781DB01832&wopisrc=https%3A%2F%2Flasersurgerycare-my.sharepoint.com%2Fpersonal%2Ffrontdesk_lasersurgerycare_com%2F_vti_bin%2Fwopi.ashx%2Ffiles%2Fccf503163fbc4cc49bff199b5ed0af61&wdhostclicktime=1587666730670&jsapi=1&newsession=1&corrid=95ba930d-2488-405e-ba2e-9d405e5b0e3c&usid=95ba930d-2488-405e-ba2e-9d405e5b0e3c&wdredirectionreason=Unified_SingleFlush#_edn19) Condyloma acuminate are more commonly known as genital warts and verruca are common warts. There are several defining features that differentiate EA and condyloma acuminatum. In one study by Lee and Wu, the average age for those with EAs was older than for condyloma (51 years and 17-33 years, respectively)[2]. Furthermore, EAs were lighter in color, smaller, and sporadically distributed compared to condyloma[2]. One of the reasons for the frequent misdiagnosis of condyloma is that both types of lesions are often found in the genital region[2]. EA has also been reported on the trunk and the extremities[4,11,24].

A case study published by Dupont *et al*[24] outlined the original misdiagnosis of condyloma acuminatum before the correct diagnosis of EA in a 51-year-old male patient[24]. The misdiagnosis of condyloma led to a prescribed treatment of cryotherapy. The lesion did not resolve after initial treatment and was biopsied after treatment. Pathologic examination demonstrated acanthosis, hyperkeratosis, cytoplasmic vacuolization, coarse keratohyalin granules, and eosinophilic inclusions. The lesion was then correctly diagnosed as EA.After arriving at the proper diagnosis, the patient was spared further treatment.

As EAs are benign, treatment is not mandatory[2].However, if patients desire treatment, there are several methods that may be employed. These methods include excision, cryotherapy, curettage, and ablative laser therapy[2,8]. There are also various topical therapies such as imiquimod, retinoids, lactic acid, glycolic acid, salicylic acid, and calcipotriol, which regulate the proliferation and differentiation of keratinocytes[2,8,9,20,25].One patient also reported complete elimination of EA after four weeks imiquimod 5% applied topically, while another reported softening of the lesions with lactic acid cream[8,25].Another patient found 0.1% tacrolimus ointment effective for relief of pruritus associated with EA after only a few weeks of use[26].

If EA is suspected, we recommend that the provider considers biopsy or excising the lesion to ensure an accurate diagnosis. Once EA is definitively diagnosed, we recommend explaining that treatment is not medically necessary. If a patient still desires removal of the benign papules for aesthetic, psychological, or other reasons, medical providers should explain possible treatment options as well as their associated risks and benefits to be sure an informed decision is made. If the patient has only one or two EA lesions, excision is often preferred. Topical therapy is most useful with a higher disease burden that would be associated with a painful recovery after surgical treatment of widespread disease.

Correctly diagnosing EA is extremely beneficial for patients and medical practitioners. If EA is misdiagnosed, patients are likely to undergo unnecessary treatments, as outlined above. These treatments are often painful and uncomfortable for patients and may also place an unnecessary financial burden on them. It is relevant to mention the psychological burden of misdiagnosis. In their case report, Fletcher *et al*[8]noted that their patient “was relieved to learn that her lesions did not represent a sexually transmitted disease and were consistent with agminated epidermolytic acanthoma[8,12,16].” We believe this case is of interest because EA is relatively uncommon and often confused with other dermatological conditions. Our primary goal in presenting this review is to provide clinicians with a knowledge to properly diagnose and treat EA.

**CONCLUSION**

In this manuscript, we describe a patient who presented with a solitary perianal EA and summarize the available literature. EAs are benign lesions, often presenting as pale papules in the genital region, trunk and extremities that are commonly misdiagnosed as condyloma accuminatum, verruca, and seborrheic keratosis. Careful histopathological examination demonstrating more than 50% EHK is necessary to reach an accurate diagnosis. Although treatment of EA is not medically necessary, many patients desire treatment, which includes topical or surgical therapy.

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**Footnotes**

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**Figure Legends**



**Figure 1 External inspection of the anus demonstrates a pale colored papule three millimeters in size located one centimeter from the anal verge at the right anterior position (left panel).** Magnified view of the lesion with high high resolution anoscopy shows a smooth, well-demarcated perianal lesion (right panel). The right panel shows the same lesion as the left panel at a higher magnification. The magnified photo was taken through a clear anoscope that was used to flatten out the perianal skin folds. The shine is a result of the lubricating gel.



**Figure 2 Hematoxylin and eosin photomicrographs of the leison.** The left panel demonstrates compact hyperkeratosis (yellow star) and acanthosis (orange arrow), with the formation of keratin crypts. Higher magnification (right panel) shows intracellular vacuolar degeneration of the epithelial cells of the spinous cell layer, as indicated by the orange arrows.