A primer on extramammary Paget’s disease for the urologist

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Abstract: Extramammary Paget's disease (EMPD) is a rare and lethal intraepithelial malignancy that remains poorly understood. No standardized guidelines or consensus statements exist with regards to the diagnostic evaluation, therapeutic approaches and follow-up management. Complete surgical excision with negative margins has been accepted as the mainstay of treatment for EMPD to decrease the risk of local recurrence and to maximize durable cure. Non-invasive therapies, such as laser therapy, photodynamic therapy, radiotherapy and topical chemotherapy have been utilized, but are best reserved for non-surgical candidates. While the debate on the surgical approach between wide local excision (WLE) and Mohs micrographic surgery (MMS) continues, several studies have demonstrated the ability of WLE to be performed safely and effectively and to yield equally satisfactory outcomes with similar rates of recurrence to MMS. Patients undergoing surgical excision often require complex closures with skin grafting or local flaps to close genital defects. We aim to provide an up-to-date review of the current knowledge of EMPD. In addition to discussing the clinical presentation and prognostic outcomes, we focus and elaborate on the diagnostic approaches and treatment alternatives available. This information may serve as a primer for the urologist who may be called upon to treat this malignancy primarily or to provide wound coverage secondarily.

Keywords: Extramammary Paget's disease (EMPD); Mohs micrographic surgery (MMS); surgical margins; wide local excision (WLE)

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Introduction

Extramammary Paget's disease (EMPD) is a rare yet lethal cutaneous malignancy with an overall survival rate of 60.2% at 10 years postdiagnosis (1). This disease is distinct from Paget's disease of the breast and predominantly affects the apocrine gland-bearing skin such as the perianal, genital and axillary regions of the body. Due to the rarity of this disease, controversies exist within the literature regarding its true prevalence, its association with concurrent internal malignancies and the diagnostic evaluation, therapeutic approaches and follow-up management. Currently, treatment recommendations for EMPD are based off single institution series and small case reports only, with many different treatment options, both in clinical and experimental use, yielding variable outcomes and results (2-4). We aim to provide an up-to-date review of the current knowledge of EMPD. In addition to discussing the clinical presentation and prognostic outcomes of this disease, we also focus and elaborate on the diagnostic approaches and treatment alternatives available to physicians when faced with patients presenting with this malignancy.

Epidemiology & pathophysiology

The incidence of EMPD has been reported to be as
low as 0.12 per 100,000 people and represents 21% of primary scrotal cancers and 2% of primary vulvar cancers, respectively (5-7). With regards to gender and racial preponderance, Asian men seem to have a fourfold increased risk of being diagnosed with EMPD when compared to their Caucasian counterparts (1,8). Previous studies have also reported a female predominance in Caucasian populations (M:F ratio of 1:2–1:7) with the opposite being true among the Asian population (M:F ratio of 2:1–4:1) (5). EMPD represents 6.5% of all cutaneous Paget’s disease and it predominantly affects patients between ages 50 to 80 years, with a peak age of 66 years old (9-11). The most common sites affected by EMPD is the vulva (65%), followed by the perianal region (20%) and subsequently the penoscrotal and groin areas (14%) (12).

Two prognostically distinct pathogenesis of EMPD have been described (13). In the primary or intraepidermal form, carcinoma develops in situ from the apocrine gland ducts and is less commonly associated with an underlying malignancy; however, it is still capable of invading the dermal layers and metastasizing over time (14). In contrast, the secondary form arises from an epidermotropic spread of malignant cells from a primary tumor within a contiguous epithelium, such as the genitourinary or gastrointestinal tract, or from a dermal adnexal gland (15,16).

**Clinical presentation**

EMPD most commonly presents as a well delineated or poorly defined erythematous and scaly plaque that may encrust, ulcerate or develop pigmentation (Figure 1) (17). Due to its similar clinical presentation to many benign conditions, the diagnosis of EMPD is often delayed for years, with only 17% of patients being correctly

![Figure 1](https://example.com/image1.png)

**Figure 1** Genital EMPD can present as a well delineated or poorly defined erythematous lesion that may affect the (A) scrotal, (B) suprapubic, (C) inguinal or (D) perineal regions of the body. EMPD, extramammary Paget’s disease.
diagnosed at first presentation (18). The most common presenting symptoms include pruritus (up to 60–72%), rash or erythema, which usually raises little suspicion for a malignant etiology (19,20). The initial differential diagnoses are commonly contact dermatitis, seborrheic eczema and fungal infections (12,18). Thus, conservative management such as topical emollients or corticosteroids, antifungal creams or other oral therapies are often prescribed first for these benign conditions. Patients typically experience a delay in definitive diagnosis for an average of 21 to 43 months after multiple rounds of failed treatment and persistent symptoms (10,11). Therefore, providers should biopsy any recalcitrant lesions that fail to respond to expectant treatments or acquire a specialist referral to decrease the risk of delayed diagnosis.

On histology, lesions usually reveal an epidermal infiltration of Paget cells, which appear as large, round cells with an abundant, pale-pink cytoplasm, surrounding a hypochromatic nuclei, occasionally with a prominent nucleolus (21). Pathognomonic cells contain intracytoplasmic sialomucin, which is capable of staining periodic acid-Schiff (PAS), mucicarmine, colloidal iron and alcin blue stains, which can aid in the diagnosis of EMPD (9). Cells may also express cytokeratin (CK), which are easily identifiable on immunohistochemical staining. While CK7 has been reported to have good sensitivity for EMPD, ranging from 86–100%, CK20 appears to be more specific for this disease (22-24). The expression of hormonal receptors has also been examined. A lack of both the estrogen and progesterone receptors together with the presence of androgen receptors and overexpression of HER-2 protein is often suggestive of EMPD (25,26). Furthermore, the presence of tumor suppressor protein p53 as well as the expression of tumor proliferation markers, such as Ki-67 and cyclin D1, has been linked to the secondary form of EMPD, which also predicts the invasiveness of EMPD lesions (27,28). While there are no grading systems for EMPD histology, the unified perception is that all EMPD lesions are considered high grade (29).

**Course and prognosis**

Patients with primary or intraepithelial EMPD have a favorable prognosis, with studies showing that the mortality rates of patients treated adequately for non-invasive disease do not differ significantly from that of the general matched population (30). Primary disease, though seemingly treatable, has the potential to disseminate and become invasive (31). Overall mortality rates for the secondary form of EMPD have been reported to be 26–66%, either from metastatic EMPD or from their associated internal malignancies (19). Moreover, mortality rates were found to be significantly higher in patients with an underlying adenocarcinoma when compared to those who did not (46% vs. 18%, P<0.05) (12).

Several factors such as dermal invasion, elevated CEA levels and lymph node metastasis portend a greater risk of adverse prognosis, with the former being the most significant. Hatta found that the 5-year survival rate for patients with deep invasion beneath the reticular dermis was significantly lower than patients with no invasion or with microinvasion to the papillary dermis (32). Histological assessment is therefore needed for appropriate risk stratification (33-35). Serum CEA levels have also been associated with predicting systemic metastasis in EMPD patients with a sensitivity of 70% and specificity of 94% (36). CEA may be used as an indicator to monitor treatment effects and outcomes for EMPD (37,38). Regional lymph node metastasis has also been shown to significantly affect the prognosis of EMPD. Unfortunately, limited studies have evaluated the utility of lymphadenectomy as a treatment strategy for EMPD (32,34). Further assessment of lymphadenectomy should be performed before it can be accepted as a standard treatment. Studies have also examined the efficacy of sentinel lymph node biopsy (SLNB) and reported a significant association between SLN positivity and increased dermal invasion rates, lower overall 5-year survival rates, and higher lymphovascular metastatic rates (39-44). Due to the limited existing data regarding lymphadenectomy, chemotherapy, immunotherapy and radiation therapy in EMPD disease, patients with metastatic disease should be evaluated via a multidisciplinary approach to develop a treatment plan that may maximize outcomes for individual patients.

**Association with internal malignancy**

In contrast to Paget’s disease of the breast, whereby 100% of patients have an underlying ductal breast carcinoma, the rates of EMPD with an associated malignancy is reported to be 21–29% (12,45). Controversy exists within the literature regarding this association with concurrent internal malignancies. Chanda demonstrated that topographic locations of EMPD appear to be closely related to the anatomic sites of the underlying malignancy. For example, neoplasms of the male genitourinary tract (e.g., prostate)
were associated with penoscrotal EMPD, neoplasms of the gastrointestinal tract (e.g., rectal) were associated with perianal EMPD, and neoplasms of the female genitourinary tract (e.g., squamous cell and adenocarcinoma of the cervix, Bartholin glands) were associated with vulvar EMPD (12). Therefore, directed screening tests for underlying malignancies based on clinical presentations have been recommended.

**Investigations and diagnostic evaluation**

A thorough history and physical examination should be performed with special attention to the location, distribution, size, color and morphology of the EMPD lesion. Palpation for enlarged lymph nodes and hepatosplenomegaly should be performed with a breast, pelvic and digital rectal exam as indicated. A biopsy of the lesion should be performed if not done so already. Pathology should be reviewed for factors such as dermal invasion which portends a higher risk of adverse prognosis (18,32,45).

A comprehensive laboratory workup, including serum carcinoembryonic antigen (CEA) levels should be obtained. Cross-sectional imaging of the abdomen and pelvis may be performed to evaluate for nodal or metastatic disease. Screening for associated internal malignancies, such as cervical, colorectal, bladder and prostate may be done as clinically indicated with Pap smear, colonoscopy, cystoscopy and serum prostate-specific antigen, respectively. If patients have lapsed the recommended screening intervals, respective specialty referral may be warranted (46).

**Table 1** Response and recurrence rates of non-invasive therapies for extramammary Paget’s disease (EMPD)

<table>
<thead>
<tr>
<th>Treatment modality</th>
<th>Response rate (%)</th>
<th>Recurrence rate (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Laser therapy (48-53)</td>
<td>86–100</td>
<td>29–100</td>
</tr>
<tr>
<td>Photodynamic therapy (54-59)</td>
<td>14–50</td>
<td>38–56</td>
</tr>
<tr>
<td>Radiotherapy (60-64)</td>
<td>62–100</td>
<td>0–35</td>
</tr>
<tr>
<td>5-fluorouracil &amp; Bleomycin (65-70)</td>
<td>57–100</td>
<td>25</td>
</tr>
<tr>
<td>Topical Imiquimod (71-77)</td>
<td>52–80</td>
<td>19</td>
</tr>
</tbody>
</table>

**Non-surgical treatment**

**Laser therapy**

Laser treatment for EMPD has garnered much interest as a conservative approach that may preserve anatomy and sexual function. Several reports have demonstrated the effectiveness of both the CO$_2$ and Nd:YAG lasers to treat EMPD with the advantage of shorter operative and hospitalization times; however, the lack of histologic evidence for analysis, post-operative pain and anesthesia requirements have prompted both physicians and patients to seek alternative treatments (48-50). Several reports have also noted high recurrence rates even up to 67–100% with this treatment modality (51,52). This is likely due to the multifocal and extensive nature of EMPD lesions and the overly superficial ablative techniques provided by laser therapy which may not adequately treat microinvasive or invasive disease (53).

**Photodynamic therapy (PDT)**

PDT has been used to treat several neoplasms such as non-melanoma skin cancer, esophageal carcinoma and even...
non-small cell lung carcinoma (54,55). It relies on the interaction between oxygen and a photosensitizer, either topical 5-aminolevulinic acid (ALA) or systemic porfimer sodium (PS), to generate reactive oxygen species that selectively destroy neoplastic tissue (56). While topical ALA offers excellent cosmetic outcomes, its ability to treat invasive and multifocal extensions of the disease remains questionable (57,58). While PS may alleviate this concern, systemic administration of a photosensitizer may generate a more severe local reaction which requires longer healing times. The safety and efficacy of PDT have yet to be clearly elucidated and therefore should be limited to patients who are unable to undergo surgery or with lesions in difficult anatomic locations (59).

**Radiotherapy**

Use of radiation therapy in the treatment of EMPD has been reported in several case reports (60-64). The majority of these patients presented with primary lesions on functionally delicate areas or were non-surgical candidates. Acute and chronic radiation toxicity is the major adverse effect. Initial investigations have described many different treatment techniques with a diverse range of radiation beam types, energy and dosages, all of which have yielded varying results and outcomes. Further studies evaluating the safety and efficacy of radiotherapy should be conducted before conclusions can be drawn.

**Topical chemotherapy**

Topical chemotherapeutic agents have been utilized for EMPD. While these agents have reported response rates as high as 57–100% in localized disease, the side effects related to these agents including severe pain and dermatitis, moist desquamation and allergic reactions have mostly

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**Figure 2** Advantages and disadvantages of Mohs micrographic surgery, wide local excision, and non-surgical therapy for EMPD. EMPD, extramammary Paget's disease.
caused them to fall out of favor (65-67). 5-fluorouracil (5-FU), mostly used as an adjunct to surgery, has been reported to be useful in clearing residual lesions which may not be excisable during surgery (67). It may also have a role in highlighting subclinical areas of EMPD, allowing better visibility during surgery or be useful postoperatively to detect recurrence of disease (68). Most investigators, however, recommend against the use of 5-FU as a monotherapy as it has only been shown to clear clinical but not pathological disease.

Topical bleomycin has also been considered for the treatment of EMPD. Watring et al. reported varying responses and outcomes in a series of seven patients treated for recurrent vulvar EMPD. Of these, four patients experienced complete therapeutic response with bleomycin, one of which required retreatment 30 months later and subsequently showed no evidence of recurrence (65).

Topical 5% imiquimod cream, more commonly known for treating genital warts, has been increasingly prescribed for off-label use in localized EMPD. EMPD response rates have been reported to be as high as 52–80% with a 19% recurrence rate (71). Imiquimod is an immune response modifier that enhances both innate and acquired immunity via the stimulation of cytokines, such as interferon-α and tumor necrosis factor-α. These cytokines in turn augment the anti-tumoral immune system to increase neoplastic cell death and destruction. Although the safety and efficacy of imiquimod for EMPD have yet to be completely exemplified, its side effect profile is better tolerated than much of the earlier chemotherapeutic agents, with mild dermatitis being the most commonly reported symptom (2,72-74). Depigmentation is another adverse effect that has been previously described and may obscure visible margins on gross examination at the time of excision or post-operative follow-up (75,76). Further studies thoroughly evaluating imiquimod are required before it may be considered as an adjunct to surgery, a potential alternative in non-surgical candidates or as part of a therapeutic combination with other non-invasive treatment modalities (77).

### Surgical treatment

Surgical excision with negative margins is the mainstay treatment for EMPD. Lesions may skip, be multifocal, or develop asymmetrically, which makes achieving negative surgical margins challenging (20). Furthermore, margins cannot be clearly identified with visual inspection alone as malignant cells are capable of extending microscopically beyond the clinically evident lesion. Achieving negative margins is paramount for long-term survival as it has shown to prevent further invasion, metastasis and recurrence (Table 2) (17,78-81). Choi et al. identified that marginal status and lymphovascular invasion were the most valid

<table>
<thead>
<tr>
<th>Author</th>
<th>Surgery type</th>
<th>R+/SM+</th>
<th>R+/SM−</th>
</tr>
</thead>
<tbody>
<tr>
<td>Jung (11)</td>
<td>WLE</td>
<td>3/10 (30%)</td>
<td>1/9 (11%)</td>
</tr>
<tr>
<td>Hegarty (17)</td>
<td>WLE, MMS</td>
<td>2/3 (67%)</td>
<td>3/12 (25%)</td>
</tr>
<tr>
<td>Curtin (78)</td>
<td>WLE</td>
<td>2/8 (25%)</td>
<td>3/14 (21%)</td>
</tr>
<tr>
<td>Baehrendtz (79)</td>
<td>WLE</td>
<td>8/12 (67%)</td>
<td>5/16 (31%)</td>
</tr>
<tr>
<td>Bergen (80)</td>
<td>WLE</td>
<td>2/6 (33%)</td>
<td>1/8 (13%)</td>
</tr>
<tr>
<td>Black (81)</td>
<td>WLE</td>
<td>14/20 (70%)</td>
<td>3/5 (60%)</td>
</tr>
<tr>
<td>Choi (82)</td>
<td>WLE</td>
<td>6/15 (40%)</td>
<td>2/16 (0%)</td>
</tr>
<tr>
<td>Long (83)</td>
<td>WLE, MMS</td>
<td>14/35 (40%)</td>
<td>17/98 (17%)</td>
</tr>
<tr>
<td>Yang (84)</td>
<td>WLE</td>
<td>9/18 (50%)</td>
<td>0/18 (0%)</td>
</tr>
<tr>
<td>Wang (85)</td>
<td>WLE</td>
<td>5/9 (56%)</td>
<td>3/72 (4%)</td>
</tr>
<tr>
<td>Fishman (86)</td>
<td>WLE</td>
<td>2/5 (40%)</td>
<td>3/9 (33%)</td>
</tr>
<tr>
<td>Composite</td>
<td></td>
<td>67/141 (48%)</td>
<td>41/277 (15%)</td>
</tr>
</tbody>
</table>

R, recurrence; SM, surgical margin; WLE, wide local excision; MMS, Mohs micrographic surgery.

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Table 2 Correlation of surgical margin status to recurrence rate for both wide local excision and Mohs micrographic surgery
prognosticating factors for EMPD recurrence, while Long et al. demonstrated that patients with positive margins had a four-fold increased risk of suffering local recurrence when compared to those with negative margins (82,83).

Several techniques have been adopted to help achieve negative margins, including the use of preoperative mapping biopsies, wide margins, intraoperative frozen sections and immunohistochemical and fluorescein staining (24,84,85,87-90). Yang et al. reported a 92% rate of obtaining negative margins with intraoperative frozen sections versus 26% without frozen sections. Misas et al. reported a 97.4% positive predictive value and 99.9% negative predictive value when assessing disease extent with fluorescein when compared to direct visualization (84,88). Despite these efforts, obtaining negative margins remains challenging (91). A study in 2005 found that 10 out of 19 patients (53%) were found to have positive margins despite undergoing intraoperative frozen sections during WLE (11). This may occur when only limited amounts of pathology frozen sections are assessed intraoperatively and complicated by the presence of multicentric, multifocal disease (83,86,92). Given the importance of obtaining negative margin status, we recommended surgical treatment to be performed at higher volume centers with the resources available to effectively perform adjunctive approaches.

The decision of surgical approach has been a long-standing debate among physicians. While MMS involves examining all the histological margins of the tumor during surgery, intraoperative frozen sections during WLE may sample a more limited area. Conceptually, MMS may seem like the preferred option over WLE for the treatment of EMPD. As such, certain investigations have shown that MMS may be superior to WLE in achieving negative margins (8,90,93). Unfortunately, this disease develops multifocal and skip lesions, which may account for some of the treatment failures with MMS. Effective use of MMS requires specialized training and may not be appropriate for larger lesions or lesions located in sensitive genital locations.

Although composite rates of achieving negative margins with MMS (97%) was higher than WLE (65%), other studies have also demonstrated comparable outcomes in obtaining adequate cancer control with both methods (6,11,17,83,94). Lesion size may play a role in selecting the surgical approach. A review of 38 cases of EMPD treated with MMS found that 76% of lesions were ≤ 10 cm while 97% of cases had lesion sizes ≤ 15 cm (93,95,96). Conversely, 95% of lesions treated with WLE were found to be ≥ 10 cm with 70% of lesions treated by Chung et al. being ≥ 15 cm, with the largest lesion measuring 30 cm in diameter (6,97). Several studies have demonstrated the ability of WLE to be performed safely and adequately and to yield equally satisfactory and durable outcomes to MMS with recurrence rates ranging from 21–60% in WLE and 0–26% in MMS (Table 3) (78,80,98-103). This suggests that while MMS may be effective in obtaining negative margins, it may be less amendable to treating larger EMPD lesions, in which case WLE may be the more favorable option.

Morbidity and complication rates differ between these two surgical approaches. MMS allows for maximal tissue sparing and decreased complication rates when compared to WLE which is associated with higher risk of neurovascular injuries, lymphedema and skin defects (94). However, MMS is limited to smaller lesions, is performed under local anesthesia and may require closure of the defect at a separate encounter by another surgeon (93). Conversely, WLE may be performed for larger lesions or lesions in sensitive areas. Patients undergoing WLE may also benefit from simultaneous primary closure or admission for inpatient wound care prior to delayed closure as final pathology is being reviewed (97). Both MMS and WLE have their respective advantages and an individualized, shared-decision making approach is recommended.

EMPD defects may require reconstruction with the use of complex primary closures, skin grafts and flaps (11). Skin grafts are a versatile reconstructive method with reported use as high as 44–80% in the management of EMPD. Advancement flaps, rotational flaps and myocutaneous flaps of the gluteal and thigh muscles may also be used for coverage and may require the assistance of a plastic surgeon (104-108). In general, penile shaft lesions may best be covered with skin grafts to preserve cosmesis and reduce the risk of contracture or chordee formation. Suprapubic, scrotal, inguinal, and perineal lesions may be amenable to complex primary closure by aggressive mobilization of neighboring tissues, flaps, or skin grafts. Due to the skills needed to both excise and reconstruct the genitalia, a reconstructive urologist may be best suited to manage localized genital EMPD.

At our institution, WLE is the preferred method for excision of genital EMPD lesions (Figure 3) (97). Pre-operative mapping biopsies and intra-operative frozen sections are used in conjunction with a 2-cm margin to help overcome the insidious nature of EMPD. Murata et al. found that the distance between the resected edge of the EMPD lesion to the last lesional cell on histopathology
Table 3 Correlation of surgical approach to recurrence rate

<table>
<thead>
<tr>
<th>Author</th>
<th>R+/WLE (%)</th>
<th>R+/MMS (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fanning (6)</td>
<td>14/32 [44]</td>
<td></td>
</tr>
<tr>
<td>Lee (8)</td>
<td>19/57 [33]</td>
<td>2/16 [13]</td>
</tr>
<tr>
<td>Zollo (20)</td>
<td>6/12 [50]</td>
<td></td>
</tr>
<tr>
<td>Louis-Sylvestre (51)</td>
<td>7/31 [23]</td>
<td></td>
</tr>
<tr>
<td>Curtin (78)</td>
<td>2/5 [40]</td>
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<td>Bergen (80)</td>
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<td>Long (83)</td>
<td>40/119 [34]</td>
<td>4/30 [13]</td>
</tr>
<tr>
<td>O’Connor (90)</td>
<td>18/83 [22]</td>
<td>1/12 [8]</td>
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<td>Fishman (86)</td>
<td>2/8 [25]</td>
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<tr>
<td>Yugueros (92)</td>
<td>3/7 [43]</td>
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<td>Hendi (93)</td>
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<td>7/27 [26]</td>
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<td>37/112 [33]</td>
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<td>8/22 [36]</td>
<td>2/11 [18]</td>
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<td>Marchesa (99)</td>
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<td>McCarter (100)</td>
<td>6/20 [30]</td>
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<td>Sarmiento (101)</td>
<td>6/13 [46]</td>
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<td>Thomas (102)</td>
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<td>Tebes (103)</td>
<td>8/23 [35]</td>
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<tr>
<td>Composite</td>
<td>185/568 [33]</td>
<td>27/154 [18]</td>
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</tbody>
</table>

R, recurrence; WLE, wide local excision; MMS, Mohs micrographic surgery.

Figure 3 Wide local excision with pre-operative mapping biopsies and intra-operative frozen sections is the preferred method for treating genital EMPD lesions at our institution. (A) A 2-cm margin is delineated around the clinically suspicious lesion as indicated by the dotted line; (B) Wide local excision was performed and (C) the defect was closed primarily after frozen sections were confirmed negative. Final pathological margins were confirmed to be negative and the patient did not require repeat excision. EMPD, extramammary Paget’s disease.
measured 1.02 cm on average (109). This, coupled with surgeon preference, justified our rationale for obtaining 2-cm margins when managing EMPD patients. Our protocol is to delay complex wound closure (skin grafting or flaps) until negative margins have been confirmed on final pathology. Patients requiring complex wound closure are admitted for inpatient wound care with wet-to-dry dressings or xenografts while permanent specimens undergo expeditious pathology review. If positive margins are identified, further excision of the corresponding region is performed. Once negative margins are confirmed, complex wound closure with or without split thickness skin grafting or local flaps is performed during the same admission (97).

Although there are no strict guidelines regarding the need and frequency of post-operative management, continued surveillance and follow-up is absolutely warranted due to the high incidence of recurrence with EMPD. We follow patients every 3 months in the first year, 6 months in the second and annually thereafter. A routine physical examination is required during each clinic visit, while cross sectional imaging of the abdomen and pelvis can be performed with either CT or MRI to rule out systemic metastasis. Mapping biopsies may be performed according to clinical suspicion for disease recurrence while serum CEA levels are obtained to monitor treatment response.

Conclusions

Surgical excision to achieve negative margins remains the mainstay of treatment to decrease local recurrence rates and maximize durable cure in noninvasive disease. Further studies are still necessary to examine the implications of dermal invasion and lymph node involvement in EMPD and to elucidate the most efficacious treatment modality with the least morbidity. EMPD patients should be referred to centers of excellence with the experience and resources of a multidisciplinary panel that can ensure homogeneity of care, eventually allowing for the development of treatment protocols and consensus guidelines.

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None.

Footnote

Conflicts of Interest: The authors have no conflicts of interest to declare.

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Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved.


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