Granulomatous vaginal ulceration due to metastatic cutaneous Crohn’s disease

HUGH J FREEMAN MD, RAYMOND KWONG MD, STEPHEN L SACKS MD


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Ulcération vaginale granulomateuse attribuable à une maladie de Crohn cutanée métastatique


Crohn’s disease may involve the vulva due to continuous extension of the anal, rectal and/or colonic inflammatory process with indolent, undermined perineal ulceration showing granulomatous inflammation (1-3). In addition, direct enteric fistulous communication with the genitourinary tract may occur. Finally, vaginal or other cutaneous lesions (ie, erosions, ulcers) separated from the gastrointestinal tract by normal skin (ie, ‘skip lesions’) have very rarely been reported in Crohn’s disease; in the case of genital disease, these are generally characterized by the absence of perineal and/or perianal disease and histologically by the presence of giant cells, noncaseating granulomas or both (4-8). This phenomenon, called ‘metastatic’ cutaneous Crohn’s disease (MCCD) (9-13), appears to have a variable clinical course, but requires careful differentiation from other, especially infectious, causes.

The present patient was initially referred to a sexually transmitted diseases clinic for evaluation of painful vaginal ulcers thought to be due to herpes simplex virus. Clinical assessment of the vaginal lesions, however, suggested a different cause. Studies to exclude...
other (especially infectious) etiologies, combined with features of previously diagnosed Crohn’s disease, led to a diagnosis of MCCD of the vagina.

CASE PRESENTATION

A 28-year-old Caucasian female was initially seen at a community hospital in April 1984 with intermittent fever, malaise, weight loss, abdominal pain and diarrhea for one year. Examination revealed multiple oral aphthous ulcers and right lower quadrant tenderness. Investigations showed hemoglobin 116 g/L, white blood cell count 11,500/mm$^3$ and platelet count 567,000/mm$^3$. Erythrocyte sedimentation rate was 40 mm/h (normal up to 20). Urinalysis and sigmoidoscopy were normal. Barium radiographs revealed narrowing of the ileal lumen for approximately 15 cm with mucosal swelling and ulceration, consistent with Crohn’s disease (Figure 1). An $^{111}$In autologous whole body white blood cell scan showed abnormal activity in the region of the ileum, cecum, and ascending and transverse colon consistent with Crohn’s disease. Treatment included sulfasalazine and prednisone, but nausea and vomiting developed with increased abdominal pain. Because of possible drug intolerance, medications were discontinued and gastrointestinal symptoms resolved.

In August 1984, the patient developed bilateral vaginal labial pain. Oral and pharyngeal aphthous ulcers were present involving the right tonsillar pillar, floor of the mouth, tongue and hard palate. Indirect laryngoscopy was normal. There were no gastrointestinal symptoms. Rectal and pelvic examinations were normal. Over the next three weeks, however, vaginal pain persisted and progressive swelling with ulceration of the labia minora developed.

She was referred to a sexually transmitted diseases clinic at University Hospital in Vancouver, British Columbia in October 1984. There was no history of known sexually transmitted disease nor sexual activity during the previous two years. Examination showed multiple aphthous ulcers of the palate and buccal mucosa, bilateral episcleritis and swollen labia minora with multiple superficial erosions and ulcerations to, but not extending through, the introitus (Figures 2, 3). The cervix was normal. There was no cervical or inguinal adenopathy. Abdominal and rectal examination, flexible sigmoidoscopy and rectal biopsy were normal.

Investigations revealed hemoglobin 123 g/L, normal red blood cell indexes, white blood cell count 7500/mm$^3$ with a normal differential and platelet count 414,000/mm$^3$. Venereal Disease Research Laboratory test was negative. Blood urea nitrogen, serum creatinine, calcium, glucose, folic acid, vitamin B₁₂, iron, iron binding capacity and protein electrophoresis were normal.

Fecal studies for ova and parasites, salmonella, shigella, campylobacter, aeromonas, yersinia (14), Clostridium difficile and C difficile cytotoxin were negative. Vaginal and cervical cultures for viruses, including herpes simplex, chlamydia, trichomonas, candida and Haemophilus ducreyi, were negative. Vaginal labial biopsy showed focal ulceration with an acute and chronic inflammatory infiltrate, as well as granulomatous inflammation with giant cells (Figure 4). Special stains and cultures for acid-fast organisms and fungi were negative.

The patient’s symptoms and vaginal lesions improved over two to three weeks; pain symptoms were initially controlled with codeine and acetaminophen along with local cleansing.
with sitz baths. By December 1984, the patient was completely asymptomatic with no persisting vaginal lesions; she has remained well since that time with no further gastrointestinal or gynecological symptoms.

**DISCUSSION**

Crohn's disease may involve all parts of the gastrointestinal tract as well as extraintestinal sites, including the genital tract. The term 'metastatic cutaneous Crohn's disease' has been used to describe involvement remote from the gastrointestinal tract, ie, 'skip lesions', separated from the gastrointestinal tract by normal intact skin.

In the extraintestinal tract manifestations of Crohn's disease, involvement of the skin is common; indeed, 20 to 40% of Crohn's disease patients may have dermatological features. MCCD, however, is rare, with fewer than 30 reported cases in the literature (4-13, 15-20). The lesion of MCCD may manifest as an ulcer, nodule or plaque and has been identified principally in the retroauricular region and inframammary area, as well as the face, upper and lower extremities, umbilicus, lower abdominal wall, penis and vulva. It appears that ulceration occurs primarily in sites of apposition of moist skin surfaces (3).

The clinical appearance of MCCD is relatively distinctive, and it should be possible, on clinical grounds alone, to differentiate this condition from other dermatological entities associated with Crohn's disease. For example, pyoderma gangrenosum typically begins with a crop of small, discrete pustules that ulcerate, while erythema nodosum presents as tender, erythematous non-necrotic nodules, characteristically located on the anterior tibial surfaces. Neither of these resembles the MCCD lesion. Histologically, granulomatous inflammation with giant cells and/or discrete noncaseating granulomas is present. The MCCD lesion correlates poorly with disease activity. Indeed, rare patients have been recorded with perineal involvement alone and no gastrointestinal disease (21).

A number of specific criteria are required to diagnose MCCD: first, cutaneous lesions, typically erosions or ulcerations, must be separated from intestinal tract by normal skin, ie, 'skip lesion'; second, pathological demonstration of an inflammatory reaction characterized by giant cells and noncaseating granulomas is required; and third, other causes of granulomatous vulvovaginitis, particularly infectious, require exclusion. MCCD may also resemble other, primarily granulomatous, cutaneous conditions, ie, fungal and mycobacterial infections, sarcoidosis, necrobiotic granulomas and foreign body reactions.

MCCD is a rare cause of genital ulcers; other possibilities should be considered (22, 23). Stevens-Johnson syndrome may also have other ulcers (ie, mouth, eyes), sometimes with target 'iris' lesions on the skin. In addition to genital ulcers, Reiter's syndrome includes an iritis with balanitis and arthritis, while Behçet's syndrome.

**REFERENCES**


