Palmoplantar Pustular Lesions during Ovulation Inducement Therapy: New Insight into the Pathomechanism of Palmoplantar Pustulosis?

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Key Words
Buserelin acetate • Estradiol

Palmoplantar pustulosis (PPP) is a recurrent pustular dermatosis localized on palms and soles [1]. Although focal infections such as chronic tonsillitis, sinusitis or periodontitis have been suggested to be modifying factors in a portion of the cases, for the most part, the essential pathomechanism presently remains unknown. Female predominance has also been recognized as a factor, which suggests an association with sex hormones. We had 2 cases of PPP following ovulation inducement therapy using buserelin acetate, a GnRH agonist. These cases suggest the relation of PPP to the alteration of hormonal conditions.

Case Reports
Case 1 is that of a 33-year-old Japanese woman who presented with pustules on her palms (fig. 1A, B). The patient did not have any personal or familial history of PPP or psoriasis and laboratory findings did not indicate any abnormality in blood counts, biochemistry or thyroid function. The patient was treated for sterility and noticed skin lesions appearing after starting ovulation inducement therapy using buserelin acetate nasal spray and estrogen patches. She had, however, been smoking for several years. Buserelin acetate nasal spray was administered every day from the beginning of the high body temperature phase and the patient used estrogen patches starting at the beginning of menstruation. Skin lesions appeared 1 or 2 days after every beginning of buserelin acetate treatment but before application of the estrogen patch. Parts of the skin on which estrogen patches or buserelin acetate were applied did not show any contact dermatitis or pustules. Pustules and erythemas on the palms recurred after every usage of buserelin acetate (fig. 1C) but they did not appear on the soles. The peak number of pustules gradually decreased with repeated ovulation inducement therapy treatments (fig. 1D). These skin lesions were clinically indistinguishable from PPP. Although these lesions improved and worsened cyclically along with buserelin acetate administration, the symptoms were relatively easily controlled with a topical steroid and no arthritis associated with SAPHO syndrome was apparent during all courses.

Case 2 is that of a 34-year-old Japanese woman who presented with pustules on her palms and soles (fig. 1E). She noticed persistent palmar pustules and vesicles about a month after the administration of buserelin acetate. Although she had also been smoking for several years, she had never experienced palmoplantar lesions until this time. The administration protocol of buserelin acetate and an estrogen patch was the same as that in case 1. Her lesions were recalcitrant to topical steroid, topical maxacalcitol and topical PUVA treatments. The condition improved slightly after the cessation of buserelin acetate (fig. 1F). The parts of the skin on which estrogen patches or buserelin acetate were applied did not show any contact dermatitis or pustules and the patient did not have any personal or familial history of PPP or psoriasis just as in case 1. Signs of arthritis suggesting SAPHO syndrome did not appear during all courses.

Discussion
PPP, which is considered a palmoplantary localized pustular dermatosis, is relatively common especially in middle-aged smoking women [1, 2]. Although its relation to focal bacterial infection is known, the treatment of focal infection occasionally improves skin lesions in limited cases. Recently, metal allergy has also been suggested as a cause of PPP [3]. In addition, genetics has been identified as a factor. In both cases reported here, there was a history of smoking, yet there were no other past skin diseases.

Although both patients received ovulation inducement therapy using buserelin acetate and estradiol in our cases, skin lesions appeared only several days after usage of buserelin acetate but before application of the estrogen patch. In addition, there were no signs of contact dermatitis on the parts of the skin where both agents were administered. These facts indicate that the skin lesion could be mainly associated with buserelin acetate via hormonal or immunological mechanisms other than an allergic response. In psoriasis, the association between sex steroid and skin lesions has been reported [4, 5]. Previous reports have indicated a positive correlation between the estrogen/progesterone ratio and the improvement of psoriasis during pregnancy [6]. On the other hand, it is also known that tamoxifen, an anti-estrogen agent, improves psoriatic arthritis [5]. Buserelin acetate, a GnRH analogue, inhibits release of estrogen and progesterone via downregulation of GnRH receptor in long-term administration. Nevertheless, in the early phase, it leads to transient estrogen release known as a flare-up phenomenon following the ovulation inducement therapy [7]. TNF-α, one of the inflammatory cytokines, is known to be involved in the pathogenesis of inflammatory skin disorders in-
cluding PPP [8, 9] and psoriasis. The level of serum TNF-α fluctuates during the menstrual cycle and it is at the peak around days 7–9 of the cycle [10]. Also, the association between GnRH and TNF-α has been identified [11]. In addition, Babayof et al. [12] reported that the serum level of TNF-α is upregulated transiently during GnRH administration. Furthermore, psoriasiform eruption after administration of an anti-TNF-α agent, so-called paradoxical adverse reaction, has been reported [13–15]. The appearance of these skin lesions usually cycles along with the administration of anti-TNF-α agent. However, in a few cases, these eruptions persist even after cessation of anti-TNF-α agent [13–15]. The differences in the clinical features between case 1 and case 2 may be linked to this phenomenon. Although the pathomechanism of PPP is still unclear, our 2 cases suggest hormonal factors may be significantly involved in PPP. Hormonal circumstances might explain the female predominance of PPP and the periodically worsening of PPP symptoms, which may lead new insight into this disease’s pathogenesis and treatment.

References


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