Retrospective study of Hallopeau-type pemphigus vegetans throughout a 20-year period. Review of the literature

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ABSTRACT

Hallopeau-type pemphigus vegetans is a rare form of pemphigus characterized by the development of vegetating lesions located mainly on flexural areas. A retrospective study was carried out of patients with diagnosis of pemphigus admitted at our Department (1988-2007). The three cases found to be Hallopeau-type pemphigus vegetans are described and a review of the literature is performed (Dermatol. Argent., 2011, 17(4): 294-300).

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Introduction

Pemphigus vegetans (PV) is considered by many authors to be a rare variant of pemphigus vulgaris and is characterized by vegetating lesions, acantholysis in histopathology and IgG deposition and/or complement with honeycomb pattern in immunofluorescence. It is an infrequent disease whose exact prevalence is unknown.

PV was referred to as a variant of pemphigus vulgaris by Neumann in 1876, and 13 years later, Hallopeau described at the First International Congress of Dermatology and Syphilography, in Paris, a different variant of pemphigus vegetans characterized by the presence of pustules, which bears his name.

These two forms of PV differ in their clinical presentation, course and prognosis. By virtue of the therapeutic implications that this entails, we highlight the relevance of differentiation between both variants.
Objectives

- To estimate the prevalence of Hallopeau-type pemphigus vegetans in the overall number of pemphigus cases and the relative frequency of each clinical manifestation of the disease.
- To characterize its presentation, response to treatment and course.

Material and methods

A review was carried out of the patients who were hospitalized at our Department between January 1998 and December 2007 who were diagnosed with pemphigus through clinical, histopathological evaluation and immunofluorescence (the latter in most of the cases).

Results

A total of 203 patients were analyzed (Graph 1). Pemphigus vulgaris is the most frequent variant - 70.5% of cases. The second most frequent variant is seborrheic pemphigus (15%), pemphigus foliaceous (6%), pemphigus herpetiformis (6%), IgA pemphigus (0.5%) and paraneoplastic pemphigus (0.5%). Hallopeau-type pemphigus vegetans constitutes 1.5% of the patients population studied and corresponds to the three cases described below.

Clinical cases

Case 1

45-year old female, without relevant pathological antecedents, who consulted at our Department because of lesions on the scalp, lip commisures and axillae after a one-year course.

**Physical examination:** on scalp, we observed a large oval raised lesion, 5 x 7 cm in diameter, pink-grayish in color and with irregular surface, with isolated erosions covered by small yellowish crusts (Photo 1). It was soft, with a non-infiltrated base and painless. On the axillary area, there were many papillomatous agminated formations with a purulent secretion and fetid odor (Photo 2). Such lesions were also present on lip commisures (Photo 3). On labial semi-mucosa, there were erosions mostly covered by hematic and “honey-colored” yellow crusts. The rest of the skin was unaffected, Nikolsky sign was negative and her general condition was good.

**Complementary tests:** laboratory: WBC count 6,000 cell/mm³ (Eo 10%, 600 cell/mm³), the rest was unremar-
Case 2

40-year-old male, with asthma and epilepsy antecedents, under treatment with diphenylhydantoin, who was first seen at our Department in 1994 due to exophytic, pruriginous lesions on axillae and inguinoscrotal and intergluteal folds, after a course of several months.

**Physical examination:** on the axillae, papules and reddish-purple plaques which converged forming vegetating-like lesions. These lesions were also visible on the inguinoscrotal region and intergluteal fold, where they presented an erosive, oozing, smelly surface (Photo 5). There were erosions on labial semi-mucosa. He did not evidence any other cutaneous manifestation, Nikolsky sign was negative and, like in the previous case, the patient’s general condition was good.

**Complementary examinations:** laboratory: WBC count 10500 cell/mm³ (Eo 5%, 525 cell/mm³, the rest was unremarkable). Cytodiagnosis of vegetating lesions and erosions on oral semi-mucosa: flaps of acantholytic cells. **Histopathological skin examination:** epidermal acanthopapillomatosis and collections inside it formed by eosinophils and acantholytic cells. Indirect immunofluorescence: IgG with an interkeratinocyte pattern, titer 1/40.

**Diagnosis:** Hallopeau-type pemphigus vegetans.

**Treatment:** methylprednisone 0.5 mg/kg/day orally, dapsone 100 mg/kg/day orally, ciprofloxacin 500 mg every 12 hours and metronidazole 500 mg every 8 hours (for 14 days).

**Course:** complete remission of dermatosis for 5 years after which he relapsed with the same places previously affected. These lesions also evolved favorably when treatment was resumed.

Case 3

84-year-old male, without known pathological antecedents, who, 14 months before consultation (2006), had recurrent outbreaks of pustules on large folds area which progressed into vegetating lesions.

**Physical examination:** on axillary region, voluminous vegetating lesions, of irregular surface, with deep grooves in which purulent, fetid material collected and alternated with pustule-strewn plaques (Photo 6). Some less significant lesions could be observed on left submammary fold. On the inguinoscrotal region, the lesions were of a considerable size and adopted a cerebriform aspect (Photo 7). Like in the previous cases, there were no other cutaneous lesions, the mucosal membranes were unaffected, Nikolsky sign was negative and the dermatosis did not alter the general condition.

**Complementary examinations:** laboratory: Hb 9.4 g/dl, Hto 27% (ferrokinetic profile: chronic disorder anaemia); WBC count 6,700 (eosinophils 15%), VSG 71 mm/1st
hour, albumin 3.2 g/dl. The rest was unremarkable. Cytodiagnosis: flaps of acantholytic cells. Histopathological skin examination 40 X: epidermal parakeratosis, acanthosis, irregular lengthening of rete ridges and an intraepidermal pustule. 100X: it was observed that this pustule contained some neutrophils, numerous eosinophils and isolated acantholytic cells. On the epithelium, eosinophilic spongiosis was observed and the underlying dermis evidenced moderate inflammatory infiltrate consisting of lymphocytes and eosinophils. IFD: IgG deposition and interkeratinocyte C3.

**Diagnosis:** Hallopeau-type pemphigus vegetans.

**Treatment:** methylprednisone 1 mg/kg/day, ciprofloxacin 500 mg every 12 hours, clindamycin 600 mg every 6 hours EV.

**Course:** improvement of dermatosis, but the patient presented sepsis from the cutaneous lesions. The initiation of immunosuppressive treatment was suggested but the patients did not return for controls.

**Discussion and literature review**

Pemphigus vegetans is predominant in adults between 40 and 50 years old but it may occur in other age groups, including children and it might be a little more frequent in females. It is an autoimmune disease involving antibodies directed against desmosomal proteins present in stratified squamous epitheliums. Such antibodies exhibit pathogenic activity. Anti-desmoglein 3 antibodies (130 kD) are usually produced in both variants. However, in the Neumann type, an 85 kD protein not present in the Hallopeau type, is also detected by immunoprecipitation. In some patients, anti-desmoglein 1 and anti-periplakin and desmocollin 1 and 2 antibodies have also been detected. Even if the antigen of pemphigus vegetans and classic pemphigus vulgaris is the same (130 kD desmoglein 3), it is posited that in the former, different subclasses of IgG would be produced (IgG 2 and 4, whereas IgG 1 and 4 are produced in pemphigus vulgaris), capable of strongly activating the complement system. This could explain the greater affluence of neutrophils and mainly of eosinophils which occurs in vegetating lesions. Notwithstanding this, there are isolated PV cases with IgG 1 and 4. A further case of pemphigus vegetans (confirmed by histology and immunofluorescence) has been described over a skin graft. The assumed physiopathogenic mechanism is that the injury produced by surgery would expose antigens normally hidden from the immune system thus triggering a secondary immune response with the development of antibodies capable of inducing the lesions. This phenomenon, also called epitope spreading, has already been described for other dermatoses such as bullous pemphigoid and mucosal pemphigoid.

There are drug-induced cases, among which we can mention the angiotensin converting enzyme (captopril, enalapril) and penicillamine. It has even been suggested that a patient might have developed pemphigus vegetans as a consequence of intranasal heroin abuse. The chemical structures of both captopril and penicillamine are made up of highly reactive sulphydryl groups which could bind to pemphigus antigens and generate acantholysis. Heroin could act as another ligand of these antigens. In the case of enalapril, its relationship with pemphigus induction is not very clear-cut.

Clinical manifestations

**Hallopeau-type variant:** it begins with outbreaks of pustules which become vegetating plaques located mainly on flexural areas. Oral cavity affection is frequent for this clinical form; two of the three cases described evidenced this manifestation. Oral involvement is referred to in the literature in 92% of pemphigus vegetans patients. It may cause papillomatous lesions on commissures, vermillion lip and cerebriform tongue or scrotum. There are cases exclusively circumscribed to the lips and oral mucosa, scalp, and others which involve only the foot. Some atypical forms have also been reported to affect fingers and toes mimicking acrodermatitis continua, or to present verrucous acral lesions (including 20-nail verrucous paronychia with nail dystrophy). The cases which mimic acrodermatitis continua can be really severe and...
lead to alterations in the phalanges and loss of nails. Besides, this clinical form may be the first manifestation of the disease after which folds and oral mucosa involvement may develop some months later. Furthermore, it is interesting to note that these patients may present HLA related in turn with pustular psoriasis. The clinical condition may turn to be very florid and the vegetating lesions may develop a considerably large size. Nikolsky sign tends to be negative, as was observed in the cases reported. There is no alteration of the general condition or inner body parts.

**Neumann variant:** It occurs within the context of a classic pemphigus vulgaris. It starts with flaccid bullae which slough giving way to erosions some of which adopt a vegetating aspect. The bullae may be found mainly on folds, where friction and maceration favor their development. However, they may occur in other places such as scalp, trunk, acral areas mimicking common warts and causing paronychia with onychodystrophy (onycholysis, onychomadesis, subungual hyperkeratosis and trachyonychia, among others). The classic involvement of folds may be absent in these atypical cases. Exceptionally, there might be cases in which the esophagus is affected in a similar way to pemphigus vulgaris and there is significant dysphagia. In these cases, endoscopy reveals erosions and whitish plaques surrounded by a reddish halo, and histology shows rounded epithelial cells and infiltrate rich in eosinophils. There might occasionally be acantholytic cells. Immunofluorescence evidences intercellular IgG deposition and Nikolsky sign tends to be positive.

To sum up, it is obvious that there is an overlapping of clinical manifestations between both forms. The vegetating lesions, then, originate from bullae which become eroded (Neumann) or from pustules which gradually turn into hypertrophic and exophytic lesions. In the former, however, the lesions alternate with flaccid bullae and erosions and the general condition of the patient and their inner body parts tend to be altered. In both variants, the lesions may present superinfection by bacteria and/or candida (as is the case for the three patients reported). The presence of eosinophilia in the blood count is usual in both variants.

**Diagnosis:** it is performed on the basis of clinical, histopathological and immunofluorescence findings (Table 1).

**PV and neoplasias:** the incidence of neoplasias in patients with pemphigus is much higher than in the general population and constitutes between 5 to 12% of the total number of pemphigus cases. Neoplasias may precede, be concomitant with or appear after pemphigus. In most cases, neoplasia is usually diagnosed before the appearance of dermatosis. If pemphigus vegetans is itself a rare disease, its coexistence with neoplasia is even rarer. There are 5 reported cases of PV and neoplasia in the literature (2 lung neoplasias, a lymphosarcoma, a gastric carcinoma and a case of colon cancer) and in one of them there is remission of the lesions when the tumor is removed. It should be taken into account that paraneoplastic pemphigus may present vegetating-type lesions within their clinical polymorphism.

**PV and HIV:** there are various cases of patients with HIV and autoimmune bullous diseases (pemphigus vulgaris, herpetiform pemphigus, pemphigus vegetans and bullous pemphigoid). Whether there is a real association or they are fortuitous cases is unknown. However, antibo-
dies with pathogenic capability could develop within the usual polyclonal hypergammaglobulinemia in prone patients. This kind of response would occur at the early stages of HIV infection due to the clonal expansion of B cells related to cellular immunity depression. In spite of this, as immunodepression progresses, both responses decrease dermatoses improve.

**Differential diagnosis:** condyloma accuminata, pemphigoid vegetans, benign familial pemphigus (Hailey-Hailey disease). In atypical location cases, squamous cell carcinoma, chromomycosis, hypertrophic lichen and pyoderma gangrenosum vegetans, among others.

**Course:** outbreaks and remissions. Hallopeau-type has a benign course (the second case reported corresponds to a patient who is still being controlled after 13-year follow-up). It has a good therapeutic response to low doses of steroids and prolonged remissions. In contrast, the Neumann variant is associated to greater skin and general condition involvement. Thus, it requires a more aggressive treatment to control the disease. However, its prognosis would be slightly more favorable than classic pemphigus vulgaris.

**Treatment**

This being an infrequent disease, it is difficult to carry out controlled, randomized trials. Publications consist in series or isolated cases. Systemic steroids are the treatment of choice for most authors. The patients who have the Hallopeau-type variant require low doses, whereas the ones with the Neumann-type one need higher doses. Corticoids usually improve the lesions and are capable of inducing remission, but the disease tends to recur when their doses are lowered or they are discontinued. In other cases, the vegetating lesions are refractory to such treatment. The drugs which have been used associated to steroids as adjuvants with a good response include dapsone, immunosuppressive drugs (azathioprine, cyclophosphamide, cyclosporine, mycophenolate mofetil) and retinoids (etretinate, acitretin). Dapsone inhibits the chemotaxis of polymorphonuclears. Retinoids reduce cell proliferation and have immunomodulatory and antiinflammatory effects, though they are not very well clear-cut. Other treatments to consider include intralesional corticoids and metotrexato. Sawai et al describe a case of pemphigus vegetans with esophageal involvement which was successfully treated with minocycline and nicotinamide without association with steroids.

**Conclusion**

Hallopeau-type pemphigus vegetans represents only 1.5% of all pemphigus cases at Hospital F. J. Muñiz, which coincides with international literature publications (between 1 to 5% of all pemphigus cases). It presents benign biological behavior, it does not require high doses of steroids and it is controlled with dapsone as adjuvant therapy. Even if pemphigus vegetans is very rare, very important to differentiate both variants, Hallopeau-type and Neumann-type due to the prognostic and therapeutic implications that such differentiation entails.

**Bibliography**


