Case Report: Pemphigoid Nodularis—Five Patients With Many Years of Follow-Up and Review of the Literature

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Pemphigoid nodularis is a rare form of pemphigoid that joins the clinical picture of prurigo nodularis and the immunological features of bullous pemphigoid, which is therapeutically challenging. Here, we analyze five female patients with a long-lasting course of nodular pemphigoid in terms of clinical and immunological characteristics and therapy. All the patients fulfilled clinical and immunological criteria of nodular pemphigoid. We applied numerous techniques allowing the proper diagnosis: direct and indirect immunofluorescence, salt split skin, ELISA, BIOCHIP, and fluorescence overlay antigen mapping using laser scanning confocal microscopy. Our study showed that 4 of 5 patients fulfilled the clinical and immunological criteria of nodular bullous pemphigoid. Two out of 4 patients presented exclusively nodular lesions; in the other two patients, blisters and erythematosus lesions preceded prurigo-like lesions by a few years. The remaining patient had clinical and immunological criteria of nodular mucous membrane pemphigoid, presenting oral erosions, scarring conjunctivitis, and numerous disseminated nodules on the skin. All the patients were treated with multiple medicines; however, it was observed that the use of clobetasol propionate on the entire body plus antidepressants best controlled the disease. Pemphigoid nodularis mainly occurs in elderly women. In cases with coexisting psychological problems, antidepressants should be considered as an important complementary therapy to the basic one with clobetasol propionate.

Keywords: pemphigoid nodularis, nodular mucous membrane pemphigoid, clobetasol propionate, antidepressants, treatment
INTRODUCTION

Bullous pemphigoid (BP) is the most common autoimmune subepidermal blistering disease equally affecting men and women over 60 years of age (1). The disease is characterized by the development of autoantibodies against the NC16a domain of BP180, an antigen of the basement membrane zone (BMZ) (1). This leads to the characteristic clinical picture consisting of erythematous and edematous lesions as well as tense blisters localized on the trunk and extremities (2). However, it is widely accepted that BP may present unusual clinical features, like dyshidrotic, localized erythoderma (2). Pemphigoid nodularis (PN) is another rare form of pemphigoid that occurs more frequently in older women (3–5). This variant of BP is extremely challenging in terms of diagnostics and therapy since it may be clinically indistinguishable from prurigo nodularis. It is noteworthy that the etiology of prurigo nodularis is not clear, but neuroimmune alterations play a role. It is also considered that prurigo nodularis is associated with other dermatological conditions (atopic dermatitis, xerosis), systemic disorders (renal failure), as well as medications. Associated with other dermatological conditions (atopic dermatitis, xerosis), systemic disorders (renal failure), as well as medications. It is also noted that psychiatric disorders accompanied by pruritus significantly predispose to prurigo nodularis (6). However, the main difference between PN and prurigo nodularis is the lack of characteristic immunological findings in patient’s skin and serum. Here, we present five female patients with PN presenting different immunological characteristics, course, and therapy.

MATERIALS AND METHODS

Patients

The study included 5 patients, all women (mean age, 74 years; range 62–85) (detailed characteristics are depicted in Table 1), who presented clinical and immunological criteria of PN. The clinical criterion was the presence of long-lasting disseminated nodules accompanied by pruritus. The immunological criterion was positive direct immunofluorescence (DIF) with the presence of linear IgG and/or C3 along the BMZ and indirect immunofluorescence (IIF) revealing circulating anti-BMZ antibodies (Table 2).

The study was approved by the Bioethical Committee of the Medical University of Warsaw. Informed and written consent form was obtained from the participants.

Methods

Direct Immunofluorescence

Direct immunofluorescence was performed on patient’s normal-appearing tissue according to a previously described method (7).

Indirect Immunofluorescence and on Salt Split Skin

Patients’ sera were studied by routine indirect immunofluorescence method (IIF) and on the salt split skin substrate (SS-IIF) according to the method described previously (8).

Enzyme-Linked Immunosorbent Assay for BP180 and BP230

Anti-BP180 and anti-BP230 antibodies were studied with commercially available ELISA kits (Euroimmun, Lubeck, Germany) that employ the extracellular domain of recombinant proteins BP180 NC16a and BP230. Assays were carried out in accordance with the manufacturer’s instructions. The results were measured by an ELISA plate reader (KHB-ST 360). The standard recommended cutoff values were 20 RU/ml for both BP-180 NC16a and BP-230 (9).

BIOCHIP Mosaics Technique

In our study, we used the BIOCHIP Dermatology Mosaic 7 (Euroimmun, Lubeck, Germany), which consists of (i) monkey esophagus (BIOCHIP-ME), (ii) salt split skin (BICHIP-SSS), (iii) recombinant BP180 NC16a, (iv) HEK293 cells transfected with BP 230, (v) HEK293 cells transfected with Dsg1, and (vi) HEK293 cells transfected with Dsg3. The assay was carried out in accordance with the manufacturer’s instructions (9).

Fluorescence Overlay Antigen Mapping Using Laser Scanning Confocal Microscopy

The FOAM-LSCM technique was performed according to the method described previously (10). The tissue sections were incubated with mouse monoclonal antibodies directed against BMZ markers: antibody to laminin-332 (Chemicon, MAB 1949) directed against epitope of laminin-332 located in the lower lamina lucida was used as a marker of the lamina lucida–lamina densa border, whereas antibody to type IV collagen (Sigma, clone col-94) served as a marker of the lamina densa. In the second stage, the sections were treated with a conjugate mix: Cy5-labeled anti-mouse IgG antibody giving a red stain (Chemicon, AP160S) and anti-human IgG antibodies conjugated with FITC giving a green stain (Cappel, alfa chain). A confocal microscope Radiance 2000 with an appropriate means of laser lines for Cy5 and FITC (637 and 488 nm) and appropriate filters (a long-pass filter of 660 nm for Cy5 and a bandpass filter of 500–560 nm for FITC) was used to avoid cross-reactivity of reagents (11).

CASE REPORTS AND RESULTS

Characteristics of the patients are depicted in Tables 1, 2.

Case 1

The first case involves an 85-year-old woman with hypertension and with a 12-year history of BP. The onset of the disease was characterized by disseminated nodules and severe pruritus; thus, she was considered as prurigo nodularis. The diagnosis of BP was established 1 year later. During the entire course, the patient presented exclusively numerous nodules, excoriations, and subsequently postinflammatory hyperpigmentation with any
### Clinical and therapeutic characteristics of patients included in the study.

| No | Sex | Age | History of nodular pemphigoid | Comorbidities | Type of skin lesions and location | Treatment |
|----|-----|-----|-------------------------------|---------------|----------------------------------|-----------|
| P1 | F/85 | 12-year duration with numerous relapses | HT | Exclusively nodules on the trunk and limbs | Prednisone, tetracycline, methotrexate | Clobetasol propionate, mirtazapine + duloxetine |
| P2 | F/62 | 9-year duration | The longest remission lasted several months | HT, COPD, anxiety | Initially disseminated blisters and urticarial lesions, then only nodules + severe pruritus | Prednisone, methyloprednisone, tetracycline, methotrexate, oral antihistaminics |
| P3 | F/80 | 1.5 years before therapy | HT | Exclusively nodules on the trunk and limbs | Exclusively clobetasol lesionally, tacrolimus, quetiapine |
| P4 | F/67 | 9-year duration | Conjunctiva and oral mucosa in stable state for several years Nodules on the skin occasionally appear | T2DM, HT, asthma, obesity | Initially disseminated urticarial lesions and few blisters, then only numerous nodules + severe pruritus |
| P5 | F/77 | 10-year duration | With remissions lasting a few months | CHD, coronary heart disease; obesity, asthma, sleep apnea, depression | Extensive oral erosions, scarring and BP180 was negative. Combined therapy of topical clobetasol 25 g/day and mirtazapine 15–30 mg/day resulted in very good clinical response along with itching reduction. Unfortunately, the patient was not cooperating and unsystematic with the treatment, causing recurrences of nodules. Recently, she was diagnosed with depression and started duloxetine. Adding prednisone 40 mg/day and clobetasol propionate topicaly led to the significant reduction of itch and disappearance of nodules. The dose of prednisone has been gradually tapered to 5 mg used every other day. The patient has been in remission for 6 months. |

**Case 2**

A 62-year-old woman with hypertension, chronic obstructive pulmonary disease, and anxiety disorder was referred to our clinic in 2009 year with a 3-year history of BP presenting erythemas and large blisters on the trunk and extremities with severe pruritus. For the next 6 years, we observed alternating remissions and relapses. During the last 2 years of observation in our clinic, the patient presented only nodules without any blister formation (Figure 1B).

Immunological tests were consistent with the diagnosis of BP, i.e., DIF showed linear deposits of IgG and C3 at the BMZ. IIF disclosed circulating IgG anti-BMZ antibodies. Both BIOCHIP and ELISA disclosed reaction of circulating IgG antibodies with the NC16a domain of BP180. The patient was treated with oral prednisone, methyloprednison, tetracycline, methotrexate, and oral antihistaminic without long-term remission achievement. Finally, topical clobetasol propionate in cream applied on the entire body led to very quick (in 3 weeks) disappearance of nodules. The patient was observed for the next several months without relapse and then she failed to continue the follow-up.

**Case 3**

An 80-year-old patient with hypertension was referred to our clinic in May 2015 with a 1.5-year history of pruritic papules that subsequently transformed into nodules involving her trunk and limbs (Figure 1C). There has never been any evidence of blister formation or erythema. DIF revealed linear deposits of IgG and C3 along the BMZ. On IIF, circulating IgG anti-BMZ antibodies were detected labeling the roof of the blister on SS-IIF. The BIOCHIP and ELISA were negative in terms of target antigens. The patient was treated with topical clobetasol with an initial dose of 25 g of cream per day on the entire body resulting in complete remission of skin lesions and itching in the first 2 weeks. Then, the cream application was continued with progressive dose reduction according to the original description for a few months without recurrence of skin lesions. She has been in clinical remission for 6 years.
Case 4
A 67-year-old woman with type 2 diabetes, hypertension, asthma, and obesity was referred to our outpatient clinic 7 years ago with a 2-year history of extensive oral erosions on gingiva, palates, and buccal mucosa and single erosions on the trunk (Figures 2A–C). The DIF performed on oral mucosa and skin revealed IgG and IgA deposits along the BMZ. Overlay antigen mapping using laser scanning confocal microscopy disclosed IgG deposits below laminin 332 and above collagen type IV, typically for MMP (Figures 2D, E). Topical treatment with clobetasol propionate and tacrolimus for a few weeks was not effective; therefore, prednisone 40 mg/day and dapsone 100 mg/day were launched, leading to the remission lasting 2 years. After self-discontinuation of therapy, she relapsed oral erosions and developed scarring conjunctivitis in the left eye. DIF performed on patient’s conjunctiva showed in vivo bound IgG along the BMZ confirming MMP. Dapsone in combination with prednisone 0.2 mg/kg b.m. resulted in complete remission of MMP. However, during therapy, skin itching, initially periodic, then later became permanent as well as nodules on the trunk and extremities were reported by the patient. In repeated DIF from skin biopsy, linear
deposits of IgG and IgA at the BMZ were detected, confirming the diagnosis of nodular MMP. ELISA revealed the presence of IgG antibodies against BP230 antigen only. The dapsone therapy was enriched in clobetasol propionate in cream lesionally as well as antidepressants such as escitalopram and paroxetine, which caused slight reduction of itch. Eventually, launching clonipramine with quetiapine led to the dramatic reduction of itch and prevented the formation of new nodules.

**Case 5**

A 77-year-old woman with numerous comorbidities (obesity, hypertension, ischemic heart disease, heart failure, type 2 diabetes, primary biliary cirrhosis, sleep apnea, and depression) was diagnosed with BP 10 years ago. The diagnosis was based on DIF (IgG and C3 deposits along the BMZ), IIF on salt split skin (circulating anti-BMZ IgG antibodies reacted with the roof of an artificial blister), and ELISA (positive reaction of circulating IgG antibodies with the NC16a domain of BP180 antigen). Initially, she presented classical features of BP (blisters and erythematous lesions accompanied by pruritus). At that time, she was treated with numerous medicines (prednisone, dapsone, and methotrexate) that had to be discontinued due to side effects. Then, the patient was treated with topical clobetasol propionate with an initial dose of 40 g of cream per day, which led to the consolidation phase and clearing of most of the lesions; however, with time, the patient could not keep up with her therapy regime. During the last 3 years, the patient complained of persisting severe itching and presented exclusively disseminated nodules (**Figure 1D**). Oral tetracycline in combination with vitamin PP was partially helpful in controlling the disease. The addition of mirtazapine and escitalopram significantly decreased the intensity of pruritus, but did not provide complete remission. The patient has still experienced several alternating periods of remission and relapse.

**DISCUSSION**

There are several publications on PN in the literature, mainly single case reports (4, 5, 10, 13). All authors have agreed that the correct diagnosis and treatment of the patients with PN remains a challenge. If the patients present from the beginning of the disease disseminated nodules accompanied by pruritus, they are recognized as having and are treated for prurigo nodularis for a long time. If diagnostic procedures are not performed at that time, the correct diagnosis and therapy are delayed.

Here, we report a large case series on PN, involving 5 patients with many years of follow-up. All of them were women, over 60 years of age (from 62 to 85), and this is consistent with most of the patients described in the literature; however, there are a few articles on PN in children (14, 15).

The patients presented in our case series were divergent in terms of clinical and immunological features as well as long-lasting course and therapy.

Clinically, among our patients, two presented only nodules disseminated on the trunk and extremities throughout their disease course, and two others initially developed classic BP with disseminated blisters and erythematous lesions, which, after years, transformed into exclusively PN (4, 14–16). That was consistent with previously reported cases. Thus, one should be aware that PN may be a secondary presentation of a classic bullous pemphigoid. PN may be generalized or limited to one area of the body (16–18). The remaining patient in our group fulfilled the clinical criteria of MMP at the beginning, presenting oral erosions and scarring conjunctivitis, subsequently developing numerous nodules on the trunk and upper and lower extremities. As far as we know, this is the first description of nodular MMP. This patient for several years presented as classic MMP and then developed disseminated nodular lesions. Since she complained of anxiety problems, she was considered as having prurigo nodularis, but repeated DIF of the skin disclosed IgG and IgA deposits at the BMZ, characteristic of MMP. It is important to emphasize that all our patients reported very severe pruritus for the duration of the disease, even during the clinical remission periods. It is widely accepted that pruritus is a strong trait of BP with an intensity much stronger than psoriasis and comparable with that observed in atopic dermatitis as we proved in our previous study (19). It is accepted that pruritus in BP is associated with elevated IgE serum levels, but its nature in other blistering disorders like MMP, EBA, or epidermolysis bullosa is not clear. On the other hand, itching may be a result of many factors such as dry skin, old age, drugs taken, comorbidities, i.e., type II diabetes mellitus, kidney diseases, and psychological status. All these conditions lead to scratching and releasing of pruritic cytokines with sustained itching. Thus, it is likely that overlapping of these conditions may be responsible for itching and the subsequent development of prurigo nodularis-like lesions.

The etiology of PN is completely unknown, though the disease may be caused by drugs, like nifedipine, etanercept, dipeptidyl-peptidase 4 inhibitor, psoralens, and PUVA (20–24). Interestingly, there are a few reports on PN coexisting with rheumatoid arthritis, lichen planus, psoriasis, or diabetes—diseases that are pruritic (20, 22, 24–30). It is highly likely that chronic, severe itching in these cases led to the damage of BMZ and exposition of its antigens and subsequently to the PN development. In our patients, drugs were excluded as provocative factors; however, some of them had emotional problems of various intensities, which are known to aggravate the itching.

Immunologically, classic BP is well characterized by the development of circulating IgG antibodies against the NC16a domain of BP180 and the presence of IgG and C3 deposits along the BMZ by DIF (1). Among our patients, all had positive DIF and all but one had circulating IgG antibodies labeling the roof of the artificial blister in SS-IIF. Additional studies (ELISA and BIOCHIP) disclosed, in two patients, reaction of circulating IgG antibodies with the NC16a domain of BP180 and/or BP230, like in several cases with PN in the literature. In other words, two of our patients failed to detect any target antigens despite using numerous techniques. This was observed in other papers. It may be associated with low titer of circulating antibodies in atypical forms of pemphigoid, i.e., nodular or localized BP as disclosed by Kawahara et al. (31). It was also shown that patients with PN may have circulating antibodies directed to other BP180 antigens of BMZ, like laminin γ1, which is characteristic of a separate entity:
anti-laminin γ1 pemphigoid (32). In the current study, we showed for the first time that nodules may also develop in a course of MMP. The diagnosis in that case was established using not only serological tests but also FOAM-LSCM on the patient’s tissue (33). Therefore, we postulate that PN seems to be not a subtype of bullous pemphigoid, but a symptom of different blistering disorders (anti-laminin γ1 pemphigoid, MMP). That is why it is mandatory to perform detailed tests to establish the proper diagnosis as it determines proper management. Moreover, in case of MMP, it is necessary to periodically examine the patient in terms of progress of skin and mucosal lesions, i.e., development of scarring conjunctivitis or esophagitis, as it was with our patient.

On the basis of the literature, patients with PN were extremely challenging since they were treated using various methods like oral steroids, azathioprine, cyclophosphamide, IVIG, dapsone, minocycline, methotrexate, and others with different results (27, 30, 34–37). According to the European guideline of BP, cream with clobetasol propionate on the entire body is recommended as a first-line therapy since majority of BP patients are elderly with numerous internal disorders (1). Moreover, it is worth mentioning that many patients with pemphigoid have neurological and/or psychiatric disorders (not infrequently associated with pruritus), which undoubtedly affect treatment compliance and may explain such enormous therapeutic difficulties. On the other hand, different medications may trigger BP, mainly diuretics, ACEI, and some antibiotics. In the literature, there are only a few papers on the provocative role of antidepressants in BP (38–42). In general, patients with BP should always be analyzed in terms of drugs they take; however, in a BP patient who experienced extensive pruritus (no response to common antipruritic drugs), antidepressants are worth consideration, because benefits from such therapy may be superior to the risk of BP exacerbation as we observed in presented patients. Four of our 5 patients had depression or anxiety disorder; therefore, in three of them, antidepressants were added to the dermatological treatment leading to the fairly good control of pruritus and skin lesions (see Table 1). The patient with nodular MMP (P4) has been treated all the time with dapsone with good control of oral lesions and conjunctivitis. However, dapsone did not control pruritus and nodules at all. In this patient, different combinations of antidepressants were tried, finally attaining success after clomipramine with quetiapine. This particular case suggests that, in some cases, the efforts should be made to find the right medications to control the pruritus.

In terms of course of the disease, in one of our patients (P3), it was relatively mild, similarly to the course of classical BP, i.e., the 7-month therapy led to the complete long-term remission. The remaining four patients presented chronic and recurrent course of the disease, resistant to many therapies. This could have been caused by chronic itching of the skin leading to the scratching and the formation of new nodules. Finally, adding antidepressants allowed for reduction of pruritus and enabled the healing of nodules.

In conclusion, this paper disclosed that:

1. Patients with PN are immunologically heterogeneous; therefore, there is a need for precise analysis of target antigens recognized by circulating antibodies since this determines their management.
2. Patients with PN should be verified in terms of coexisting emotional problems similarly to those with prurigo nodularis.
3. In most of the cases with PN, clobetasol propionate in cream as the first-line treatment should be considered.
4. In case of PN with very severe itching, adding antidepressants to clobetasol propionate might be considered.

DATA AVAILABILITY STATEMENT

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

ETHICS STATEMENT

The studies involving human participants were reviewed and approved by the Bioethical Committee of the Medical University of Warsaw, Warsaw, Poland. The patients/participants provided their written informed consent to participate in this study.

AUTHOR CONTRIBUTIONS

All had full access to all of the data in the study and responsibility for the integrity of the data. Accuracy of the data analysis: KS, AA, CK, and KW. Study concept and design: KW, KS, and CK. Acquisition of data: KS, AA, BJ, EP and KW. Analysis and interpretation of data: KS, AA, KW, BJ, EP and CK. Drafting of the manuscript: KS and KW. Critical revision of the manuscript for important intellectual content: CK and KW. All authors contributed to the article and approved the submitted version.

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