Review: dermatitis herpetiformis*
Revisão: dermatite herpetiforme

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DOI: http://dx.doi.org/10.1590/abd1806-4841.20131775

Abstract: Dermatitis herpetiformis (DH) or Duhring-Brocq disease is a chronic bullous disease characterized by intense itching and burning sensation in the erythematous papules and urticarial plaques, grouped vesicles with centrifuge growth, and tense blisters. There is an association with the genotypes HLA DR3, HLA DQw2, found in 80-90% of cases. It is an IgA-mediated cutaneous disease, with immunoglobulin A deposits appearing in a granular pattern at the top of the dermal papilla in the sublamina densa area of the basement membrane, which is present both in affected skin and healthy skin. The same protein IgA1 with J chain is found in the small intestinal mucosa in patients with adult celiac disease, suggesting a strong association with DH. Specific antibodies such as antiendomysium, antireticulina, antigliadin and, recently identified, the epidermal and tissue transglutaminase subtypes, as well as increased zonulin production, are common to both conditions, along with gluten-sensitive enteropathy and DH. Autoimmune diseases present higher levels of prevalence, such as thyroid (5-11%), pernicious anemia (1-3%), type 1 diabetes (1-2%) and collagen tissue disease. The chosen treatment is dapsone and a gluten-free diet.

Keywords: Celiac disease; Dermatitis herpetiformis; Skin diseases, Vesiculobullous

Resumo: Dermatite herpetiforme é uma doença bolhosa crônica caracterizada por intenso prurido e sensação de queimação em pápulas eritematosas e placas urticariformes, vesículas agrupadas com crescimento centrífugo e bolhas tensas. Apresenta associação com genótipos de HLA DR3, HLA DQW2 encontrados em 80 a 90% dos casos. É uma doença cutânea mediada por IgA com depósito de imunoglobulina A em padrão granular no topo da papila dérmica na área da sublâmina densa na zona da membrana basal, presente tanto na pele lesada com em área de pele sã. A mesma cadeia J da proteína IgA1 é encontrada na mucosa do intestino delgado em pacientes com doença celiaca do adulto, sugerindo forte associação com a dermatite herpetiforme. Anticorpos específicos com anti-endomísio, anti-reculina, anti-gliadina, e recentemente identificado, o subtipo transglutaminase epidérmica e tecidual, assim como a produção aumentada da zonulina, são descritas em ambas as afeções enteropatia sensível ao glúten e a dermatite herpetiforme. Exibe depósitos de IgA em padrão granular na papila dérmica. Doenças auto-imunes exibem maior prevalência como tireoidopatia em 5 a 11%, anemia perniciosa em 1 a 3%, diabetes tipo 1 em 1 a 2% e doença do colágeno. O tratamento de escolha é a dapsona e dieta isenta de glúten.

Palavras-chave: Dermatite herpetiforme; Dermatopatias vesiculobolhosas; Doença celiaca
INTRODUCTION
Bullous diseases constitute one of the most extraordinary chapters of dermatology. Pathophysiological mechanisms diversity subordinated to its varied etiology, extensive range of clinical manifestations with often systemic disease involvement, require a well-conducted medical evaluation method; therefore, translate into surprising difficulties that require specialized treatment and habilitation in overcoming the diagnostic and therapeutic challenge.

The bubbles are efflorescence filling with liquid composed of plasma and inflammatory cells, resulting from the change of cell structures and intercellular junctions structures responsible for the adhesion of epithelial tissue. Vesicles are known as the diameter of the cavity less than 0.5 cm, and bubble is greater than 0.5 cm, intraepidermal if the lesion is present in the basal layer to the stratum corneum. Through knowledge of the pathophysiology of the cleavage plane, the characteristics of the inflammatory infiltrate and especially the mechanism of blistering, it is possible to distinguish bullous dermatoses.

DERMATITIS HERPETIFORMIS
Dermatitis herpetiformis (DH) was described in 1884 by dermatologist Louis Duhring, who placed it in the same clinical category as pemphigus and pemphigoid, thus composing the class of bullous diseases. In 1888 Brocq described similar skin lesions diagnosed as "polymorphic pruritic dermatitis" and after examining Duhring’s report, admitted that it was the same pathology. Therefore, Duhring-Brocq’s disease is now used as a synonym for DH.

In 1943, through distinction of vesiculation mechanism, Civatte differentiated pemphigus (intraepidermal bullae), pemphigoid and DH (blistering of the basement membrane zone). The association with celiac disease, a gluten-sensitive enteropathy, and DH was observed in the sixties by Mards et al., Fry et al. and Shuster et al.

Epidemiologically, DH is a rare disease. It affects mainly young adults, although it had been diagnosed in infants aged eight months as well as in elderly people aged ninety years. Males are more affected, with a ratio of 2:1, but in patients under 20, the ratio is 12 females for every 8 males. Prevalence of DH varies across different countries, with 1:1,000,000 new cases / year in Germany, 11 per 100,000 in Scotland; 20-39 per 100,000 in Sweden and 58.8 per 100,000 in Ireland.

There are reports of disease in other members of the same family, either DH or adult celiac disease, in 2.3 to 10.5% of cases. Ethiopathogenesis has an immunological cause but is not fully understood. It is known that there is a higher incidence of genotypes HLA DR3, HLA DQw2 in 80-90% of patients, HLA B8 and HLA DQ8 in 10-20% of cases, as well as adult celiac disease.

Cutaneous lesions begin with itching or a burning sensation in erythematous papules and urticarial plaques. There are grouped vesicles and tense blisters with centrifugal growth, whose contents may be serous or hemorrhagic, with symmetrical distribution (Figures 1, 2 and 3). Bullous elements rupture, culminating in denuded areas of exulcerated skin and crust. Subsequently, there is residual hypopigmentation or hyperpigmentation.

The topography usually affected are the extensor areas: lower limbs (anterior thigh and knee), elbows, buttocks and sacral region, although the shoulder, scapular region and scalp may also be affected. Lesions in the oral mucosa are unusual.

Gastrointestinal clinical manifestation of gluten-sensitive enteropathy may occur at any age, either in childhood when cereals are introduced into the oral diet, or in adulthood without any prior food intolerance reaction. Symptoms include diarrhea, steatorrhea, malabsorption with resulting anemia, metabolic bone disease, weight loss and malnutrition. However some patients have no gastrointestinal signs or symptoms at all, since the majority of DH patients are asymptomatic, as only 20% of them develop intestinal symptoms.

Gluten is an amorphous protein composed of gliadin and glutenin amino acids, which are found in cereal seeds from the Grameneas family, such as wheat, barley, oat, malt and rye. They are cereals that
This can only be eradicated through adopting a gluten-free diet for several years, because even drug therapy does not alter this pattern.23

Patients present gluten-sensitive enteropathy diagnosed by villi atrophy in a small intestine biopsy, in addition to specific antibodies in the serum, although most of them are asymptomatic.8,9,12,20,24

The same protein IgA1 with J chain and secretory component found in the small intestinal mucosa in adult celiac disease, is also present in the skin of patients with DH, suggesting a strong association between both diseases.15 Other antibodies such as specific endomysium, antireticulina, gliadin and, recently identified, the epidermal and tissue transglutaminase subtypes, as well as overproduction of zonulin, are found in both conditions, along with enteropathic sensitivity to gluten and DH.25-29 IgM, IgG and C3 may also be present.

Studies have shown that cross-reaction occurs between some substances because they present common epitopes, and are therefore recognized as self-antigens triggering the disease.24

High association autoimmune diseases include thyroid (5-11%), pernicious anemia (1-3%), type 1 diabetes (1-2%), and collagen tissue disease.8,9,14,16,30 Thus, Caproni et al. suggest screening for autoimmune diseases in patients with DH, as determination of antiperoxidase antibodies (present in 20% of cases), TSH, T4 and T3, anti-gastric parietal cells (10-25% positive), ANF, anti -Ro/SSA and glucose.16

In 1970, Gjone & Nordoy were the first to report increased incidence of lymphoma in DH patients, which was confirmed for decades, especially T-cell lymphoma.14,16,31-33 However, Lewis et al, in a study of 846 patients published in 2008, contradicted this relationship, such as found no increased risk of mortality in these patients.30

Although 100% of patients with DH present sensitivity to gluten enteropathy, only a minority develop symptoms of colic or intestinal malabsorption, it is described the ratio of 1:5.7,10,20 There is evidence that a gluten-free diet alone brings about improvement or even complete remission of intestinal symptoms, and improvement of skin lesions in DH.7,11,18

The skin biopsy must come from a new, intact bulla.14 Histopathological studies have revealed the Piérard microabscess on top of the dermal papilla in a granular pattern at the top of the dermal papilla in the area of the sublamina densa of the basement membrane, which is present both in affected skin areas and in healthy skin.8,9,19
The direct immunofluorescence of perilesional skin affected is the gold standard to confirm the diagnosis with the deposition of IgA1 in granular pattern in the lamina lucida of the basement membrane zone (Figure 4).2,9,34,35 Less than 5% of cases present deposits of IgA in a linear pattern, which must be distinguished from linear IgA dermatosis.7 A fibrillar IgA standard deposit is reportedly found in 50% of the Japanese population with HD, but several authors have questioned whether it corresponds to a variant of DH, a different disease, or only to an alternating pattern consequent longitudinal and transverse orientation of microfibrils, visualized through an electron microscopy, as suggested by Ko CJ et al.36 Sometimes, it is necessary to carry out a second test for the diagnosis because, during the early stages of the disease, this typical feature may not be found. The deposit of immunoglobulin A does not change with dapsone, but approximately two years of gluten-free diet abolishes this finding.1

Indirect immunofluorescence may be useful to detect the presence of autoantibodies and circulating anti-endomysial, anti-gliadin, and anti-reticulin IgA, and anti-epidermal transglutaminase antibodies.1,2 Tissue transglutaminase antibodies (anti-tTG) can be measured by ELISA, showing greater than 90% specificity and sensitivity of 47-95%. It is used to diagnose DH and to assess patients adhesion to gluten-free diets and intestinal damage. Anti-tTG is 64% homologous to anti-epidermal transglutaminase (anti-eTG), which acts against the specific antigen in DH.16,37 Jaskowski et al. suggest that 20% of patients have anti-tTG negative, but these patients are anti-eTG positive. The same authors found a higher sensitivity of anti-eTG, which may help in cases of difficult diagnosis. Prevalence of anti-eTG decreases in children compared with adults.37

IgA1 antibodies against smooth muscle have 100% specificity and 52 to 100% sensitivity in the diagnosis of HD. However anti-tTG may be absent in patients with gluten free diets.16

Studies show that 100% of patients with DH exhibit histopathological changes of celiac sprue, ie, villous atrophy characterizing an aspect flat surface cuboidal epithelial cells that lose the orientation of the basal nucleus, increased proliferation of crypt cell causing hyperplasia and loss of their structures with increased lymphocytes and plasmocytes cells in the lamina propria. These findings are not pathognomonic of the sprue, but the return of histological pattern after gluten-free diet confirms diagnosis.37

Villous atrophy of small bowel mucosa biopsy found in most patients with DH is less severe than in celiac disease or no tropical sprue.29 The absorption test D-xylose is altered in 10-33% of cases. Finding of iron deficiency anemia or megaloblastic anemia for folate deficiency and steatorrhea is not uncommon.2 Although the clinical manifestations of D differ strongly in EC, both improve significantly with gluten-free diets. This measure provides relief in the itching and burning sensation of the vesicle-erythematous-papule, disseminated or localized on skin. Sometimes, it causes total regression of the cutaneous manifestations. This behavior seems to be great for the prognosis of the disease. However it is arduous and difficult in our country since gluten is a ubiquitous substance, and not all food products in the market contain explicit information regarding the presence or absence of gluten. Furthermore, the high cost of alternative products offered to patients with gluten sensitivity represents a financial obstacle for most Brazilian citizens. Thus, it is very difficult to adhere to and depends on determination, guidance and investment. 9,38,39 Iodine strict restriction is also suggested by some authors.13

The treatment is successful in patients who tolerate dapsone.33,6 The initial dose is generally between 100-200 mg per day and the response occurs within three hours to two days, with no new lesions appearing.29 The patient should take a minimum dose, sufficient to suppress the disease. Some patients take 25 mg of dapsone per week, but others need 400 mg daily.2

The side effects of drugs are dose-dependent. 50mg of dapsone can cause some hemolysis. Doses of 150 mg may decrease 2 grams of hemoglobin, which can be asymptomatic in healthy patients or trigger several symptoms and signs in patients with cardiac or lung disease, or even in elderly people.2 Methemoglobinemia is ferrous iron (Fe +2) oxidation from hemoglobin to ferric iron, (Fe +3) caused by chemical substance. It progresses with erythrocytes

**Figure 4:** Direct immunofluorescence. Fluorescence in granular pattern in the basement membrane zone
precipitation and hemolysis characterized by Heinz bodies and “bitten cells.” Methemoglobin causes the deviation of the oxygen dissociation curve to the left, resulting in insufficient oxygen to the tissues.\textsuperscript{19}

Methemoglobinemia is generally less than 5% and does not exceed 12% in DH. The signs and symptoms occur at methemoglobin 3%, characterized by cyanosis, a greyish colour, weakness, headaches, tachycardia, nausea and abdominal pain.\textsuperscript{2}

Other adverse effects are peripheral neuropathy (reversible with dose reduction); morbiliform eruption, erythema nodosum, erythema multiforme, exfoliative dermatitis, toxic epidermal necrolysis, severe hypoalbuminemia with anasarca, leukopenia, agranulocytosis, which may be fatal during the first three months, cholestasis and hepatitis.\textsuperscript{3,40} The sulfone syndrome usually occurs within 6 weeks of treatment, regardless of dosage, and is characterized by exfoliative dermatitis, hepatitis, fever, lymphadenopathy, leukocytosis, headaches, vomiting and haemolysis.\textsuperscript{41,42}

Due to both the side effects and gravity of some cases, patients taking dapsone must be closely and continuously monitored. To avoid disastrous developments before starting treatment, deficiency in glucose-6-phosphate dehydrogenase enzyme must be investigated, as it evolves with severe hemolysis because enzyme methemoglobin reductase - (NADPH - Methemoglobin reductase) - depend on NADPH to act.\textsuperscript{2,19} A complete hemogram should be performed weekly during the first month, every two weeks within two months and often during the treatment. Renal and hepatic tests should also be required before starting the drug, with regular monitoring.\textsuperscript{2}

Patients who cannot tolerate the use of dapsone may benefit from sulfapyridine (1 to 1.5 g / day, tetracycline 2 g / day, together with nicotinamide 1.5 g / day or cyclosporine for resistant cases).\textsuperscript{2,11,42} Sacchidanand S. et al revealed satisfactory results with dexamethasone-cyclophosphamide pulse therapy.\textsuperscript{43}

An important observation is that anti-inflammatory drugs usually worsen DH.\textsuperscript{43}

Prognosis courses with periods of remission and exacerbation.\textsuperscript{3,44} An emotional event or infection may trigger a new worsening of the disease.\textsuperscript{3}

**FINAL CONSIDERATIONS**

It is extremely important for bullous diseases to be recognized by clinicians to prevent worsening of the symptoms with contraindicated or ineffective drugs, since the peculiar characteristics of bullous disease pose a high risk of serious and systemic consequences arising from electrolyte imbalance.\textsuperscript{43,44}
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