CYCLO-OXYGENASE 2 IS PRESENT IN THE MAJORITY OF LESIONAL SKIN FROM PATIENTS WITH AUTOIMMUNE BLISTERING DISEASES

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Abstract
Introduction: The in situ immune response within skin biopsies from patients affected by autoimmune skin blistering diseases (ABDs) is not well characterized.
Aim: Based on the fact that the ABD immune response is considered an adaptive immune response, both an innate immune response and inflammation would be expected in these diseases. Our investigation investigates the presence of cyclo-oxygenase-2 (COX-2), since this enzyme is commonly involved in innate immune responses.
Methods: We utilized immunohistochemistry (IHC) to evaluate the presence of COX-2 in lesional skin biopsies of patients affected by ABDs. We tested 30 patients with endemic pemphigus foliaceus (EPF), 15 controls from the endemic area, and 15 biopsies from healthy controls from the USA. We also tested archival biopsies from patients with selected ABDs, including 20 patients with bullous pemphigoid, 20 with pemphigus vulgaris, 8 with pemphigus foliaceus and 12 with dermatitis herpetiformis.
Results: Most ABD biopsies stained positive for COX-2 in the lesional blister and/or the dermal inflammatory infiltrate, accentuated in the upper neurovascular plexus. In BP and EPF, the COX-2 staining was also seen in the sweat glands. All controls were negative.
Conclusions: We document that COX-2 is expressed in lesional skin of patients with ABDs.

Key words: Cyclo-oxygenase 2; autoimmune skin diseases; endemic pemphigus foliaceus

Abbreviations and acronyms: Bullous pemphigoid (BP), immunohistochemistry (IHC), direct and indirect immunofluorescence (DIF and IIF), hematoxylin and eosin (H&E), basement membrane zone (BMZ), intercellular staining between keratinocytes (ICS), pemphigus vulgaris (PV), cicatricial pemphigoid (CP), autoimmune blistering skin diseases (ABDs), fogo selvagem (FS), endemic pemphigus foliaceus in El-Bagre, Colombia (El Bagre-EPF).

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What does the current study add?
Immune system markers such cyclo-oxygenase-2 seem to be expressed in the majority of lesional skin of patients with ABDs, and may represent a consistent feature of the inflammation commonly present in these diseases.

Introduction
Multiple therapies have been utilized for the treatment of cutaneous autoimmune blistering skin diseases (ABDs). Steroids represent a commonly utilized therapy, because many autoimmune disorders are B-lymphocyte mediated processes depositing autoantibodies and complement deposits in the skin. Well documented correlations exist between titers of autoantibodies and the clinical severity of the diseases [1-3].
Materials and Methods

Subjects of study:
We tested 30 biopsies from patients affected by EPF in El Bagre, Colombia, South America (El Bagre-EPF) and 15 normal controls from the endemic area [4-8]. We also utilized 15 control skin biopsies from cosmetic reduction plastic surgery patients in the USA, taken from the chest and/or abdomen. Biopsies were fixed in 10% buffered formalin, then embedded in paraffin and cut at 4 micron thicknesses. The tissue was then submitted for hematoxylin and eosin (H&E) and IHC staining. In addition, we also tested biopsies from the archival files of two private laboratories led by board certified dermatopathologists in the USA; these patients underwent initial diagnostic biopsies, and therefore were likely not taking immunosuppressive medications at the time of biopsy. We evaluated 20 biopsies from bullous pemphigoid (BP) patients, 20 from patients with pemphigus vulgaris (PV), 8 patient biopsies with pemphigus foliaceus (PF) and 12 from patients with dermatitis herpetiformis (DH). For all of the El Bagre area patients and controls, we obtained written consents as well as Institutional Review Board permission from the local hospital. The archival biopsies were IRB exempt due to the lack of patient identifiers. In both dermatopathology laboratories, each biopsy also was sent for direct immunofluorescence as previously described [3], for correlation with the H&E diagnoses.

Quantitative digital morphometry and IHC staining:
The staining intensity of the antibodies was also evaluated in a semiquantitative mode by an automated computer image analysis system, designed to quantify IHC staining in hematoxylin-counterstained histologic sections. Slides were scanned with a ScanScope CS system, utilizing brightfield imaging. IHC staining was performed as previously described. For IHC, we utilized a Dako monoclonal mouse anti-human COX-2 antibody, clone CX-294; staining was performed as previously described [4-7].

Statistical analysis:
For statistical analysis, the non-parametric Mann–Whitney U-test was used to calculate significant levels for all measurements. Values of p<0.05 were considered statistically significant.

Result
We noted that 26/30 patients with EPF were positive in the epidermis in spot areas of the corneal layers, around the neurovascular areas of eccrine and hair follicles. Only 2 controls from the endemic area showed some corneal reactivity (p<0.05). Further, 17/20 biopsies from BP patients were positive for COX-2 in the sweat glands, under the blisters and along the bases of the blisters. Some reactivity was seen in the corneal layers (p<0.05). Reactivity was seen in the upper neurovascular plexus of the dermis, and in some type of junction between endothelial cells and the extracellular dermal matrix (Fig. 1). In patients with PV, 16/20 biopsies were positive in the upper dermal inflammatory infiltrate, and around the epidermal blisters (p<0.05). In patient biopsies with PF, 5/8 were positive in the epidermis in spotty areas of the corneal layers, and around the neurovascular supplies of eccrine glands and hair follicles. In 9/12 patients with DH, positive staining was noted, mostly under the BMZ (p<0.05). In Figure 1, we highlight the most common patterns of positivity found in these patients.

Discussion
Because adaptive immunity has been demonstrated to be play a pathogenic role in ABDs, it is important to note that innate immunity is the first step in an adaptive immune response. Unfortunately, few studies have specifically studied molecules involved in the adaptive immune response in ABDs. Hallmark pathologic events in ABDs include vasodilatation of the microcirculation, resulting in increased blood flow to the affected area. The vasodilatation is responsible for the heat and redness that occurs at sites of inflammation. In ABDs, we also often see increases in the permeability of upper dermal blood vessels, promoting the movement of fluid and plasma proteins into the interstitial areas [8]. In ABDs, chemotactic neutrophils, monocytes and other white blood cells manifest as an inflammatory infiltrate. Activated complement, another innate immune marker found in pemphigus blister fluids, suggests a pathogenetic role for complement in this disorder [9].

In our study, we investigate the immune response induction marker COX-2, a key enzyme of arachidonic acid metabolism. Cyclooxygenase exists as two distinct isoforms. COX-1 is constitutively expressed in most tissues, whereas COX-2 is inducibly expressed at sites of inflammation. An additional recently documented isoform, COX-3, is produced via alternative splicing of COX-1 and has also been described [10]. COX-2 is also inducibly expressed in neoplastic tissues. Prostanoids are produced by many cell types, and act on target cells through specific G protein-coupled receptors. Although prostanoids have traditionally been considered acute inflammatory mediators, studies using knockout mice show that prostanoids may regulate selected aspects of both innate and adaptive immunity; such a dual role may be the case in ABDs. Each prostanoid, depending on which receptor it acts on, exercises specific effects on immune system cells such as macrophages, dendritic cells, and B and T lymphocytes, often in concert to microbial ligands and cytokines. These cellular actions affect the strength, quality, and duration of immune responses. Prostanoids play a critical role in immunopathology, via inflammation, autoimmunity and cancer pathophysiology [8]. We found limited specific information regarding autoimmune diseases and the expression of COX-2. In oral lichen planus, increased levels of COX-2 have been reported [11]. In experimental autoimmune encephalomyelitis, celecoxib (a new generation COX-2 inhibitor) retards inflammation [12]. Since our study utilized archival biopsies, the requisition forms did not contain specific information regarding administration of immunosuppressive agents. The precise role of this enzyme in ABDs requires further investigation since this was a pilot study. We thus recommend larger studies, utilizing only patients known to have no immunosuppressive therapy to study the role of this molecule in ABDs.

We found no specific studies regarding the presence of COX-2 in ABDs. For this reason and because our study has a small sample size, we recommend larger studies to further define the role of COX-2 in these disorders.

Conclusion
We suggest that molecules such as complement and COX-2 are present in the majority of skin lesional biopsies from patients with ABDs; further, it is possible that COX-2 may consistently contribute to the inflammation seen in these diseases. Recent data has also demonstrated that prostanoids regulate selected aspects of both innate and adaptive immunity, and this may also be the case in ABDs.

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