The Enigmatic Autoimmune Response in Endemic Pemphigus Foliaceus

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Abstract. Endemic pemphigus foliaceus, known as Fogo Selvagem (FS) in Brazil, is a disease characterized by subcorneal blisters and IgG4 anti-dermoglein 1 (Dsg1) autoantibodies. Epidemiological studies of FS strongly support an environmental etiology. A 15-year surveillance of the Limao Verde Amerindian reservation in Brazil has uncovered information on the transition of the autoimmune response from the pre-clinical stage to disease state. This incubation time may evolve over several years. The serological markers of the pre-clinical state of FS are IgM anti-Dsg1, IgE and non-IgG4 autoantibodies against Dsg1. The disease stage of FS is characterized by the rise of pathogenic IgG4 anti-Dsg1 autoantibodies. In this review, the authors reviewed the literature on the relevance of the humoral autoimmune response of FS as well as the possible environmental triggers of anti-Dsg1 autoantibody formation. Based on epidemiological observations, the authors hypothesize that the pathogenic IgG4 response in FS may be triggered by hematophagous insect bites.

Key words: fogo selvagem, desmoglein, pemphigus foliaceus, pemphigus vulgaris, ELISA.

Non-Endemic and Endemic Pemphigus Foliaceus

The spectrum of clinical phenotypes of skin diseases known collectively as pemphigus include pemphigus vulgaris (PV), pemphigus foliaceus (PF), drug-induced pemphigus and paraneoplastic pemphigus1. These diseases are characterized by spontaneous intraepidermal blister formation on the skin and mucous membranes and pathogenic autoantibodies against desmosomal glycoproteins2-4. In PV and PF the autoantibodies recognize desmoglein 3 (Dsg3) and Dsg1 respectively2. PF is a life-threatening skin disease characterized by subcorneal blisters and IgG anti-Dsg1 autoantibodies. These autoantibodies are known to be pathogenic by passive transfer experiments4. Fogo Selvagem (FS) is an endemic form of PF which occurs among local inhabitants of rural Brazil5, Colombia6,7 and Tunisia8. FS shares similar clinical, histological and immunologic features with the non-endemic form of PF seen in the USA and around the world9. Importantly, the IgG anti-Dsg1 autoantibodies are predominantly IgG4 and recognize the ectodomain of this molecule10,11.

We have described an active focus of FS in Brazil, the Limao Verde Amerindian reservation, where the approxi-
approximately 1,300 members of the Terena tribe show a ~3 % prevalence of the disease. The following relevant observations have been made during the last 15 years of follow up of this human settlement: a) serological transition from preclinical to clinical stage of FS, including epitope mapping studies and IgG subclass switch; b) detection of IgM anti-Dsg1 autoantibodies in FS patients and healthy individuals from endemic areas; c) development of an IgG4-based classifier for FS sera, d) detection of anti-Dsg1 autoantibodies in the sera of patients with leishmaniasis, Chagas disease and onchocerciasis. Finally, we have detected a significant number of FS sera possessing IgE anti-Dsg1 autoantibodies that show correlation with IgG4 autoantibodies. These findings have prompted us to probe the immunological relationships of the IgG4 and IgE anti-Dsg1 autoantibody responses in FS patients and their possible link to a common environmental “allergen”, that in turn, may trigger the autoimmune response in FS.

**The Immune Response Against Dsg1 Begins Early in Life with Non-Pathogenic IgM Autoantibodies whereas the IgG Response Increases Gradually with Age**

Immunization of naïve individuals to a particular antigen is known to trigger a primary immune response characterized by low-affinity IgM antibodies. Re-exposure to the same antigen subsequently leads to a sustained secondary response characterized by high-affinity IgG antibodies. In humans, detection of antigen-specific IgM antibodies in sera may be an indicator of a recent infection or re-infection with viral diseases such as rubella, infectious mononucleosis or hepatitis B, bacterial infections such as Lyme disease, or parasitosis such as toxoplasmosis. We have found a relatively high prevalence of IgM anti-Dsg1 autoantibodies in sera from both FS patients and clinically normal donors residing in rural settings in or near an endemic area. Patients with PF, other autoimmune bullous diseases, as well as healthy individuals from more urban settings, did not demonstrate a significant IgM anti-Dsg1 response. A modest number of hospitalized FS patients that were temporarily isolated from their native outdoor environmental exposure showed positive IgM response. Importantly, over 50% of sera from healthy donors from three cohorts between the ages of 5 and 20 from Limao Verde possessed IgM anti-Dsg1 autoantibodies. There was no significant trend in IgM anti-Dsg1 autoantibody response with age. A recent study showed that neonates from mothers with FS do not possess IgM anti-Dsg1 antibodies. Finally, these studies showed that the prevalence of IgG anti-Dsg1 autoantibodies gradually increased with age of the donors. These novel findings support the hypothesis that an environmental antigen(s) sensitizes individuals living in these rural endemic areas of FS. This process began after the neonatal period of life because over 50% of children after the age of 5 already show IgM anti-Dsg1 autoantibodies in their sera. Further, they reinforce our findings that B cell repertoire in FS in the preclinical stage is antigen-driven.

While the observed seroepidemiology of the IgM anti-Dsg1 autoantibody response appears to suggest exposure to an environmental antigen(s), it is also known that unimmunized individuals may possess IgM antibodies to toxins, bacteria and erythrocytes that comprise a population of polyreactive low-affinity natural antibodies. These natural polyreactive IgM antibodies represent the first barrier against infection, eliminating bacteria by complement activation, thus bridging innate to adaptive immunity. Many of these IgM natural antibodies recognize single or multiple self-antigens and have been detected from early childhood throughout life. Natural IgM autoantibodies against self-antigens have been reported in autoimmune diseases such as lupus erythematosus, autoimmune hemolytic anemia, and autoimmune thrombocytopenia. It is hypothesized that polyreactive IgM or the IgG/IgM ratio of anti-dsDNA antibodies in systemic lupus erythematosus may modulate the disease and prognosis, especially in patients with nephritis.

In summary, these seroepidemiological observations suggest that the IgM response in FS patients from Limao Verde likely arises from recurrent and persistent antigenic exposure to an environmental cross-reactive antigen(s) harbored in this and other Amerindian reservations. Sensitization begins in early childhood and continues throughout life in these individuals, resulting in the production of non-pathogenic IgM and IgG anti-Dsg1 autoantibodies. Low affinity IgM anti-Dsg1 autoantibodies may not be demonstrated by routine indirect IF assays. Pathogenic IgG anti-Dsg1 autoantibodies and clinical disease may occur in only a small fraction of genetically predisposed individuals. Importantly, FS is more common among poor laborers of all races and both sexes sharing the HLA DRB1*0404, DRB1*1402 or DRB1*1406 alleles (RR: 14). It is likely that serum concentrations of IgM anti-Dsg1 autoantibodies from FS patients referred to metropolitan hospitals, away from their native environment, decrease due to elimination of the environmental antigenic stimuli.

**The Pathogenic IgG Response in FS is IgG4 Restricted and Predicts the Disease. The Incubation Time of FS Last Years**

It has been known for several years that the humoral immune response against Dsg3 in PV is IgG4 restricted.
This IgG4 restriction is also observed in non-endemic PF and FS. These studies have been confirmed using indirect IF, immunoblotting, and Dsg1 ELISA. Moreover, we demonstrated that not only the total IgG4 was pathogenic but also its F(ab’)2 and Fab’ fragments when tested in the mouse model of FS. Additional studies in FS have demonstrated that the autoantibody response in FS exhibits a limited heterogeneity, consisting of oligoclonal IgG1 and IgG4 banding, when tested with epidermal antigens by affinity immunoblotting. These studies suggested that the autoantibody response in FS exhibits an early IgG1 response followed by a sustained IgG4 response.

In a recent study we conducted the serological evaluation of IgG and IgG subclass anti-Dsg1 autoantibodies in 214 FS cases (45%) and 261 normal individuals (55%) and generated a highly sensitive and specific “IgG4 classifier/predictor.” FS patients were seen in 5 Brazilian hospitals (Campo Grande, Sao Paulo, Goiania, Brasilia and Belo Horizonte) from 1980 to 2006. Healthy controls were obtained from the blood banks of the same Brazilian hospitals and from UNC Hospital. The sera were tested by Dsg1 ELISA optimized for the detection of human IgG subclasses using murine monoclonal anti-human IgG subclass-specific antibodies and the results were expressed as index values. A logistic regression model was used to develop a “classifier” that predicts case-control status based on the four IgG subclass index values. For the purpose of developing and evaluating a classification rule, the data set (n = 475 samples) was divided at random into three parts; a training set (n = 239), a validation set (n = 118) and a test set (n = 118) containing 50%, 25% and 25% of the cases and controls, respectively. A rigorous statistical procedure was applied in order to choose the best model for prediction. Each of 15 possible models were estimated from the training set. Sensitivity, specificity and area under the curve (AUC) were estimated from the validation set. The AUC is a summary of the whole ROC curve for a given model and is not affected by choice of the cut point. Thus, AUC was used as the criterion for model selection. An observation with fitted probability above 0.45 was classified as a case. The model with only IgG4 had an AUC of 0.961 (in the validation set). Using additional predictors had a negligible effect on the AUC. Thus, the model with only IgG4 was chosen as the final model because of its parsimony and high AUC value.

The classification rule based on IgG4 classifies a given subject as a case if the IgG4 index value exceeds 6.43; otherwise it classifies as a non-case. The estimated AUC is 0.971 (95% CI: 94-100%), sensitivity is 92% (95% CI: 82-98%), and specificity is 97% (95% CI: 89-100%) of the “classifier” that was determined on the test set. It is worth mentioning that the “classifier” was developed entirely in the training and validation sets, yet it performed extremely well in the test set.

The positive predictive value (PPV) and negative predictive value (NPV) of the classifier, when applied in a population such as the Limao Verde reservation where the prevalence of FS is 3%, were calculated. It was estimated that an individual from Limao Verde, classified as positive by the classifier, has a 49% chance of having FS while a subject classified as negative has a 99.7% probability of being disease-free. The PPV and NPV for IgG and other IgG subclasses were lower than for IgG4. Since the prevalence of IgG anti-Dsg1 autoantibodies in normal inhabitants of endemic areas of FS, i.e. Limao Verde is high, the use of IgG as a classifier in these human settlements would be very limited. On the contrary, IgG4 anti-Dsg1 autoantibodies are detected in the sera of individuals developing FS or during recurrences. For similar reasons the IgM anti-Dsg1 autoantibodies did not perform well.

The IgG4-based classifier was further validated by analyzing other groups of patients and people at risk to develop FS:

1. Eleven FS patients during the preclinical stage of the disease. In the first group, the classifier predicted FS in 5/11 individuals (45%) during the preclinical stage and in all samples during the clinical stage of FS (100%). It must be emphasized that this classifier identifies subjects with serological features of FS regardless of the presence of active skin disease. We propose that this IgG4-based classifier is a serological marker of FS during the preclinical and clinical stages of the disease. During the preclinical stage this classifier may show variations over time due to fluctuations in environmental antigenic stimulation.

2. Sixty Japanese patients. Twenty patients with mucosal PV, possessing only anti-Dsg3 autoantibodies, were classified as normal donors by the classifier (because of the absence of anti-Dsg1 autoantibodies). The IgG4-based classifier performed well in a group of 20 PF patients, of whom 18 were identified as cases. In a group of 20 mucocutaneous PV patients, possessing only anti-Dsg1 and anti-Dsg3 autoantibodies, the classifier predicted the disease in 17 cases. Hence the IgG4-based classifier performed well in the Japanese group of patients with PF and mucocutaneous PV since both groups of patients possess anti-Dsg1 autoantibodies.

3. Three cohorts (n = 96) of normal individual (ages 5-20) from Limao Verde. The IgG4 classifier identified 21 individuals (22%) with serological features of FS: 6/34 individuals (17.6%) in cohort 1 (age 5-10) were positive, 6/39 (15.3%) in cohort 2 (age 11-15) were also positive as well as 9/24 (37.5%) in cohort 3 (age 16-20). The same donors show the following percentages of total IgG anti-Dsg1 autoantibodies: cohort 1, 1/34 (2.9%), cohort 2, 3/41 (7.3%) and cohort 3, 7/24 (29%). Inter-
estingly, one member of the third cohort (JDM), classified as a case, has developed FS in the course of the study. According to the PPV of the IgG4 classifier, it is estimated that about 50% of these positive subjects from Limao Verde have FS in the preclinical stage and are at risk to develop clinical disease if the conditions are appropriate. Similarly, each of the 75 subjects identified as normal by the classifier have a 99% chance of being disease-free. Forecasting active clinical disease in individuals of both groups (positive and negative) using the classifier is the subject of current investigation in our laboratory. An ongoing prospective study of these cohorts will further validate this immunological instrument not only as an identifier of current FS serology but also as a predictor of future disease. These cohorts are evaluated clinically every 4 months and serologically every 2 years.

It is concluded from these studies that the bulk of pathogenetic anti-Dsg1 autoantibodies in FS are predominantly IgG4. In fact, a recent study showed that progression from preclinical to clinical stage of the disease is associated with a dramatic rise in IgG4 anti-Dsg1 autoantibodies as determined by ELISA assays. Further, we believe that the IgG4 anti-Dsg1 based classifier would be extremely useful in identifying individuals during the preclinical stage of FS. HLA typing and the IgG4-based classifier would become powerful tools for the selection of individuals to undergo close clinical and serological surveillance. Moreover, as the environmental risk factor(s) can also be assessed among potential FS patients, these immunological markers may enhance our ability to identify the factor(s) involved in triggering the autoimmune disease in FS.

**Antigen Selection of Anti-Desmoglein 1 Autoantibody Occurs not only During but Also Prior to the Onset of Fogo Selvagem Disease**

The diversity and clonality of the anti-Dsg1 response, as well as the likelihood that anti-Dsg1 B cells are antigen selected, were studied with peripheral blood samples from 8 FS patients and one individual with prior-to-clinical FS. Human hybridomas were generated by fusion of EBV-transformed blood lymphocytes with either mouse myeloma cells (P3X63Ag8.653) or MFP-2s myeloma cells as fusion partners. Dsg1 was used to detect anti-Dsg1 antibodies by ELISA during the screening and cloning of hybridomas. The V genes of both H and L chains of the Dsg-specific autoantibodies from these hybridomas were sequenced and analyzed for the H and L chain pairing of the autoantibodies, the diversity of H and L chain V gene usage, and the extent of somatic mutation.

Seventy-eight monoclonal anti-Dsg1 hybridomas were isolated; 38 secreted IgG, and 40 secreted IgM. Multiple lines of evidence suggest that the anti-Dsg1 autoimmune response in FS is antigen selected. First, clonally related sets of anti-Dsg1 hybridomas were identified from individual FS patients. Second, H and L chain V gene use appears to be biased, particularly among IgG hybridomas, as some H chain V gene families were over-represented and others under-represented. Third, both IgM and IgG hybridomas exhibited a high frequency of VH mutation. The replacement versus silent mutation (R/S) ratios exhibit a bias in favor of CDR (complementary determining region) amino acid replacement mutation and in favor of silent mutation of FWRs (framework regions). Interestingly, similar selection was evident among anti-Dsg1 hybridomas from an individual with prior-to-clinical disease. The V genes of hybridomas from this individual were also extensively mutated, and showed similar R/S mutation bias in CDRs and FWRs of their H and L chain V genes to that of FS patients. Thus, selection of anti-Dsg1 B cells begins well before the onset of the disease. This is consistent with the hypothesis that an environmental antigen(s) may induce the cross-reactive anti-Dsg1 response in these genetically predisposed individuals to initiate FS.

**Fogo Selvagem is an Environmentally Triggered Autoimmune Disease of the Skin**

FS shows several unique and remarkable features such as the geographic and temporal clustering of cases, the increased frequency of cases among young adults and children, the increased frequency of familial cases, and an association with certain distinct HLA-DR alleles. FS occurs in Brazilian states located between 45º to 60º west longitude and 5º to 25º south latitude in regions with an altitude between 500-800 meters (1,600-2,600 feet). FS is rare at altitudes below 400 meters (1,300 feet) or above 1,000 meters (3,300 feet). The weather in endemic regions of FS is subtropical and in northern regions supports coffee, sugarcane and cacao, while in southern regions corn, soybeans and cotton are the predominant crops. The FS patient is usually a farmer (or a member of the family) who farms in these fields or works outdoors. They are usually young adults or children. In general, poor families move to a farm and live in rustic houses located near rivers or creeks. It is also common among them to raise chickens, hogs, cows, horses and have a variety of pets such as dogs, cats, and birds. There is no reported sex or race predisposition for the development of FS. The daily activities of a family include agriculture, care of livestock and home chores. Wives and children remain at home to perform routine chores, i.e. cooking, caring for small ani-
mals or washing laundry in nearby rivers or streams. However, a great number of wives help their husbands in farming activities. The farmer and his family usually have a common bedroom and wear light clothes appropriate for the weather. The houses are usually built of reed walls and thatched roofs with open doors and open windows. Commonly, these houses harbor rodents, other small wild animals and are usually infested with blood-feeding arthropods such as bedbugs and Reduvid bugs. Because they contain shallow waters and rock beds, some rivers and streams are infested with a variety of insects, including Simulids (black fly, also known as borrachudo in Portuguese) and sandflies. The number of new cases of FS is greatest at the end of the rainy season (September to March) and least during the dry summer (April to August), suggesting that insect multiplication and the increase in the number of FS patients are related phenomena. Deforestation and the arrival of humans to settle in these regions have been salient features of FS as described by Brazilian investigators.

Interestingly, the same ecological systems found in the “pemphigus country” overlap with those described in Chagas disease, leishmaniasis and FS in the endemic states, as compared with those seen in the middle part of the 20th century. The diseases were common among workers laboring in the construction of railroads and highways in the interior of the endemic states, i.e., Sao Paulo, Minas Gerais. It is known that these diseases tend to disappear once the working and living conditions of the settlers improve. In recent years the frequency of cases of Chagas disease, leishmaniasis and FS in the endemic states, as compared with those seen in the middle part of the 20th century has dramatically decreased and this decrease has been associated with modernization in agricultural techniques. It is clear that the ecological systems of FS share similarities with these insect vector borne diseases.

Finally, the Amerindian reservation of Limao Verde, located in the state of Mato Grosso do Sul, Brazil, is the home of 1,351 members of the Terena tribe of Amerindians (as of September 10, 2006), and is an active focus of these diseases. As described above, and based on the information generated from the Limao Verde endemic focus of FS, we hypothesize that exposure of this population to an environmental risk factor triggers the cross-reactive autoimmune response that leads to FS. During early childhood a substantial number of individuals start producing IgM anti-Dsg1 autoantibodies, a process that may continue for years if the person remains in Limao Verde. It is feasible, although unproven, that in individuals moving out of an endemic area such as Limao Verde, this IgM anti-Dsg1 response may decrease and disappear. The IgG response in the Limao Verde population appears after the age of 5 years and continues through adulthood. The IgG anti-Dsg1 autoantibodies may be found in up to 1/3 of the normal population and have no pathogenic consequences.

Possible Etiological Factors Of Fogo Selvagem-Hypothesis

The diagram shown in figure 1 depicts the central hypothesis investigated in our laboratory about the immunological events triggered in genetically predisposed individuals that lead to FS. As described above, and based on the information generated from the Limao Verde endemic focus of FS, we hypothesize that exposure of this population to an environmental risk factor triggers the cross-reactive autoimmune response that leads to FS. During early childhood a substantial number of individuals start producing IgM anti-Dsg1 autoantibodies, a process that may continue for years if the person remains in Limao Verde. It is feasible, although unproven, that in individuals moving out of an endemic area such as Limao Verde, this IgM anti-Dsg1 response may decrease and disappear. The IgG response in the Limao Verde population appears after the age of 5 years and continues through adulthood. The IgG anti-Dsg1 autoantibodies may be found in up to 1/3 of the normal population and have no pathogenic consequences.
in the individual. Our studies demonstrate that, when the IgG anti-Dsg1 response in normal individuals shows predominant IgG4 autoantibodies, this individual will be at high risk to develop FS. The IgM and IgG response against Dsg1 remains as a serological marker of the pre-clinical stage of FS. After an incubation time that may last from 1 to more than 10 years, some genetically predisposed subjects may increase their IgG4 anti-Dsg1 autoantibodies titers and develop clinical FS.

The biochemical interactions between a vector and the host’s skin during the process of blood feeding by hematophagous insects are complex. In order to steal a blood meal from the host, the hematophagous vector must move quickly using mechanical tools (mouthparts) to perforate the skin and inject a cocktail of salivary substances to counteract host defense mechanisms (i.e. coagulation, vasoconstriction, platelet aggregation, etc.)77-80. The host defense ranges from the mechanical squashing of the offending insect to the in situ tissue and humoral response at the site of the bite, i.e. vasoconstriction, inflammation and antibody response. Medically relevant insects (blood feeding and others) trigger several responses in the host: a) an acute or chronic process in which the host may exhibit a local or systemic reaction modulated by IgE and IgG4 antibodies against an insect product (e.g. bee venom)81; b) they are carriers of parasites causing disease (e.g. hematophagous bugs in leishmaniasis, Chagas disease, filariasis, malaria, etc.); c) sensitization and production of anti-saliva antibodies that block parasite invasion as occurs in malaria or leishmaniasis82 where salivary proteins may promote parasite invasion77-80. Additionally, in filariasis there is a predominantly IgG4 response83. Interestingly, a small number of FS patients have antibodies to maxadilan, a potent vasodilator isolated from sand fly saliva84. Although some of the salivary components from blood-feeding insects have been extensively characterized at the molecular level85-89, others including cross-reactive carbohydrate determinants90,91 remain unstudied.

We are testing the intriguing hypothesis that a component(s) of the saliva of these hematophagous insects (reduvii, simuliiid and lutzomia) contain cross-reactive antigens with human epidermal Dsg1 and thus trigger the anti-Dsg1 autoantibody response (fig. 2). Interestingly, the genome of a well-characterized insect like the fruit fly Drosophila melanogaster contains 19 genes encoding cadherin-like molecules92; whether these molecules are present in the saliva of these insects is unknown. Considering the persistent IgM anti-Dsg1 response in normal individuals of Limao Verde we entertain the idea that the sensitizing antigen(s) may be a T-independent (TI) type 2 antigen, perhaps a carbohydrate, derived from the environment (blood feeding bugs, parasites, etc.)93,94. Examples of TI-2 antigens include, among others, multivalent polysaccharides (or other antigens with repetitive structures) from

![Figure 2. Hematophagous insects and Fogo Selvagem (FS).](image)

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### Conflict of interest

Authors have no conflict of interest to declare.

### References


