Atopic dermatitis with mononuclear phagocytic activity deficiency

W.C.N. Forte, M.C. Santos de Menezes, S.M. Cipolli Guerra de Oliveira, and S. Bruno

Immunology Section of Santa Casa Medical School and Hospital. São Paulo. Brazil.

SUMMARY

Five patients with atopic dermatitis, three males and two females, aged 2 to 17 years, had positive reactions to air allergens (Dermatophagoides pteronyssinus and/or farinae). All the patients suffered from severe recurrent dermatophytosis that responded poorly to antifungal treatment. The results of immunologic evaluation by laboratory tests were normal, except for a decrease in the ingestion phase by mononuclear phagocytes.

After diagnosis of immunodeficiency, ketoconazole shampoo was used prophylactically and at the very first signs of recurrence of dermatophytosis, systemic antifungal treatment was started, without concurrent use of macrolides and with monitoring of hepatic function. The fungal infections responded well to this treatment and the patients’ quality of life markedly improved.

Key words: Atopic dermatitis. Dermatophytosis. Phagocytic deficiency. Monocytes.

INTRODUCTION

Atopic dermatitis or atopic eczema is a chronic and recurrent inflammatory disease with exacerbations triggered by different stimuli (1). Clinically, it is characterized by pruritic lesions with scaling leading to lichenification, the latter considered to be the cutaneous component of the atopic complex (2, 3).

Several atopic dermatitis patients present recurrent or persistent infections and should be submitted to immunologic evaluation (4). Studies have demonstrated some deficiencies in the immunologic response of atopic dermatitis patients; the type, degree and prevalence of these defects however are as yet unclear (5).

The diagnosis of deficient phagocytic activity by mononuclear phagocytes is less often made as compared with the diagnosis of deficient polymorphonuclear neutrophil phagocytosis, as seen in chronic granulomatous disease, in malnutrition and other nutritional disturbances (6-8). In the presence of altered mononuclear phagocytic activity, serious infections are encountered especially by fungi, but also by viruses and intracellular bacteria. Infections are caused by Candida sp, M. gypseum, T. tonsurans, Mycobacterium tuberculosis and any type of virus.

The prognosis of atopic dermatitis with immunologic involvement is directly related to early diagnosis and specific treatment, as seen in practically all immune deficiencies.

Few cases have been described in the literature of patients with atopic dermatitis who had deficient mononuclear phagocytic activity, even though this is so important for adequate treatment and evolution of the patients’ disease.

CASE REPORTS

First case

G.P.M., female, 17 years old. At the age of eleven, she started to present intense pruritus over the whole body when the diagnosis of atopic dermatitis was made. At 13, there was aggravation of the lesions with diagnosis of Tinea corporis. This was so serious that she often could not go to school. At the age of 17, she presented papulo-vesicular lesions with eryt-
hematous base and serous exudate associated with whitish round lesions with definite contour, disseminated all over her body, in which state she was first seen at our Immunology Section. There was also an intense vaginal discharge.

Second case

T.A.S., male, 2 years old. Diagnosis of atopic dermatitis was made at the age of one. The clinical presentation aggravated when the atopic dermatitis developed fungal infection which responded poorly to topical treatment as well as frequent recurrences. At the age of 2, he was brought to the Immunology Section with intense pruritis, papulo-vesiculo-erythematous confluent lesions predominantly on the face, extensor regions and lower members associated with crusty lesions.

Third case

F.C.N., male, 12 years old. Diagnosis of atopic dermatitis was made at the age of three. At the age of four, he developed *Cryptococcus neoformans* meningitis confirmed by positive cerebrospinal fluid culture, associated with whitish oral mucosa lesions. At the age of ten, he developed crusty erythematous lesions on the face, trunk and members.

Fourth case

H.G.M., female, 12 years old. At the age of three, she presented with disseminated maculo-papular lesions, mainly on the extensor surface of the members, which progressed to lichenification. There was aggravation and at the age of four, there was alopecia with tonsured hair, areas of scaling on the face, trunk, upper members, nail thickening, labial fissures and repeated viral infections.

Fifth case

J.A.S., female, 17 years old. At the age of two, she had erythematous exudative lesions on the face. At the age of three, these lesions spread to the whole body, except hand palms and foot soles, with intense pruritis. At the age of seven, there was aggravation of her condition, when she arrived at the Immunology Section with severe atopic dermatitis with fungal infection.

RESULTS

Our patients were positive for air allergens (*Dermatophagoides pteronyssinus* and/or *farinae*). Cultures of skin scales showed *Candida sp*, *M. gypseum* and dermatophyte *T. tonsurans*. In the first case, *Candida sp* was found in the vaginal discharge. In all five patients, the values of immunoglobulins, lymphocytes, T and B cells, CD4+ and CD8+ cells, C3 and C4 components of complement as well as chemotactic and phagocytic activity for neutrophils were normal while phagocytosis was persistently decreased in monocytes.

All five patients showed decreased phagocytic ingestion phase for monocytes (table I) when compared to normal values: 27 ± 9; 78 ± 7; 75 ± 10 to 27 ± 5; 65 ± 7; 68 ± 7, according to age (9).

After the diagnosis of atopic dermatitis with dermatophytosis, associated with deficient monocyte phagocytic activity was made, all patients received systemic antifungal treatment during the fungal infection. No concurrent macrolides were given and there was control of hepatic transaminases and coagulation tests. Prophylactic use of cetoconazol was also initiated. Immediately at first signs of fungal infection, systemic antifungal therapy was started once more, always with hepatic function control. The patients started to have satisfactory response to the treatment.

DISCUSSION

Zimossan (Zy) particle phagocytosis used in our methodology assesses the monocyte ingestion phase (9). Monocytes present receptors for C3b and C5b implicated in the ingestion phase of phagocytosis, where C5b is considered the more important one. Mononuclear cell incubation with serum promotes complement activation, resulting in the formation of C3b and C5b components which unite to Zimossan particles. The union of C3b and C5b with their respective receptors on the surface of monocytes cul-

### Table I

<table>
<thead>
<tr>
<th>Monocytes</th>
<th>Chemotaxis (micra)</th>
<th>Phagocytosis (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Case</td>
<td>Case</td>
</tr>
<tr>
<td>Control</td>
<td>1º 2º 3º 4º 5º</td>
<td>1º 2º 3º 4º 5º</td>
</tr>
<tr>
<td>With homologous serum</td>
<td>36 35 42 81 66</td>
<td>35 32 47 62 30</td>
</tr>
<tr>
<td>With autologous serum</td>
<td>36 35 46 83 69</td>
<td>35 37 52 68 41</td>
</tr>
</tbody>
</table>

Allergol et Immunopathol 2002;30(5):263-6
minates with the ingestion of the Zimsoan particles by these phagocytes. The control test evaluates the viability of the method with spontaneous particle ingestion. The second and third tests evaluate zimsoan ingestion in the presence of C3b and C5b components of heterologous and homologous serum respectively.

The results demonstrated no significant difference between the second (mononuclear leucocytes incubated with Zy and human serum “pool”) and the third test (mononuclear leucocytes incubated with Zy and the patient’s serum), with normal complement values of these patients (10), indicating that the decrease in phagocytosis was due to an intrinsic monocytic defect. These data are coherent with the greater frequency of fungal infections in these patients. The children studied were of normal height and weight, which makes it impossible that the phagocytic deficiency in monocytes might be due to malnutrition (9).

Studies confirm that the skin of atopic dermatitis patients is more frequently colonized by *Staphylococcus aureus* than non-atopic individuals and that this colonization is more intense in the presence of more serious degrees of dermatitis, which contributes to the chronicity of the disease (11, 12).

Patients with atopic dermatitis may show exacerbation of their eczema which is triggered by various inflammatory stimuli, through IgE mediated mechanism, including to dermatophytes; these patients thus become sensitized to these agents and are more susceptible to cutaneous dermatophyte infections (1, 13). The positive tricofitin reactivity observed in atopic dermatitis does not necessarily signify sensitization to the dermatophytes but is probably a sign of cross-reaction to fungus (14).

*Candida albicans* which belongs to the normal human microflora, can induce synthesis of specific IgE in patients with atopic dermatitis and asthma. The *Candida albicans* sensibilization manifested in children with serious atopic dermatitis is frequently associated with immunodeficiency (15) and has been suggested as a component of the atopic dermatitis pathogenesis (16). On the other hand, eczema is one of the cutaneous manifestations of primary immunodeficiency which leaves a doubt as to the initial disease factor (17). Neutropenic patients have greater risk to develop systemic infections by *Candida albicans*, while the fungus also has the ability to activate Th2 response as evasive strategy (18, 19).

An increasing number of children with atopic dermatitis are recognized to possess immune defects, while the exact incidence is still unknown (5). Monocytes and T helper lymphocytes exert an important role in the immunologic dysfunction of atopic dermatitis. Studies have evaluated the citocin pattern in the immune response of atopic dermatitis, but results have been conflicting, while the main studies realized have used stimulation by different mitogens, which makes comparison difficult (20). Fisher et al have observed significant decrease in chemotaxis by monocytes in patients with serious mucocutaneous candidiasis and also in some patients with atopic dermatitis (21).

These observations as well as the literature propose the hypothesis of deficient mononuclear leukocyte phagocytic activity in patients with atopic dermatitis associated with fungal infections and alert that special therapeutic measures must be taken in order to improve these patients’ quality of life.

**RESUMEN**

Cinco pacientes con dermatitis atópica, tres varones y dos mujeres, de 2 a 17 años de edad, tuvieron reacciones positivas a neuroalergenos (*Dermatophagoides pteronyssinus* o *farinae*). Todos padecían una dermatofitosis grave y recurrente que respondían con dificultad al tratamiento antifúngico. La evaluación inmunológica mediante pruebas analíticas fue normal, excepto por la presencia de disminución de la fase de ingestión por fagocitos mononucleares.

Después de diagnosticar la inmunodeficiencia, se utilizó de manera profiláctica champú de ketoconazol y ante los primeros signos de recidiva de la dermatofitosis se inició tratamiento antifúngico por vía general, siempre sin el uso concurrente de macrólidos y con control de la función hepática. Las infecciones fúngicas ahora respondieron bien y la calidad de vida de los pacientes mejoró considerablemente.

**Palabras clave:** Dermatitis atópica. Dermatofitosis. Deficiencia fagocítica. Monocitos.
REFERENCES

1. Nissen D, Petersen LJ, Esch R, Svejgaard E, Skov PS, Poulsen
   K, et al. IgE sensitization to cellular and culture filtrates of
   fungal extracts in patients with atopic dermatitis. Ann Allergy

2. Castro, APBM. Dermatite atópica In Grumach, AS – Alergia e
   imunologia na infância e na adolescência, Atheneu, 1ª ed. São
   Paulo 2001;185-201.


4. Weston WL, Huff JC. Atopic dermatitis: etiology and patho-

5. Rogge JL, Hanifin JM. Immunodeficiencies in severe atopic
   dermatitis. Depressed chemotaxis and lymphocyte transfor-

6. Forte WCN, Carvalho Jr, FF. Immunodeficiências secundárias às
   alterações nutricionais. In Grumach, AS- Alergia e imunologia
   na infância e na adolescência, Atheneu, 1ª ed. São Paulo
   2001;571-577.

7. Forte WCN, Gonzales CCL, Carignani S, Mímica I. Avaliação
   de neutrófilos na desnutrição moderada. Revista da Asso-
   ciação M. Brasileira 1999;45:147-51.

8. Forte WCN, Noyoya AM, Carvalho Jr, Bruno S. Reapited fun-
   culosis in adult male with abnormal neutrophil activity. Aller-

9. Forte WCN, Campos JVM, Leão RC. Non-specific immunolo-
   gical response in moderate malnutrition. Allergologia et im-

10. Forte WCN, Forte AC, Leão RC. Complement system in mal-

11. Goh CL, Wong JS, Giam YC. Skin colonization of Staphylococ-
    cus aureus in atopic dermatitis patients seen at the National

    Staphylococcal toxic shock syndrome toxin1 inhibits monocy-

    Recent investigation on the relationship between fungal skin
    diseases and atopic dermatitis. Acta Derm Venereol Suppl

14. Rajka G, Barinn C. On the significance of the trichophytn re-
    activity in atopic dermatitis.Acta Derm Venereol 1979;59:
    45-7.

15. Samulova TL, Morinosova MA, Kranoprosina Li, Sdokhova
    SA, Sewrgeeva AS. Candida albicans sensitization in patients
    with atopic bronchial asthma and atopic dermatitis. Ter Arkh

16. Henseler T, Tausch I. Mycoses in patients with psoriasis ou

17. Saurat JH. Eczema in primary immune-deficiencies. Clues to
    the pathogenesis of atopic dermatitis with special reference
    to the Wiskott-Aldrich syndrome. Acta Derm Venereol Suppl

18. Romani L, Bistoni F, Puccetti P. Initiation of T-helper cell im-
    munity to Candida albicans by IL-12: the role of neutrophils.

    HW. Functional characterization of skin-infiltrating lymphocy-

20. Lee HJ, Lee HP, Ha SJ, Byun DG, Kim JW. Spontaneous ex-
    pression of mRNA for IL-10, GM-CSF, TGF-beta, TGF-alpha,
    and IL-6 in peripheral blood mononuclear cells from atopic der-

21. Fischer TJ, Gard SE, Rachelefsky GS, Klein RB, Borut TC,
    Stehlm ER. Monocyte chemotaxis under agarose: defects in
    atopic disease, aspirin therapy, and mucocutaneus candidia-