Annular skin lesions in childhood: Review of the main differential diagnoses

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INTRODUCTION

Annular lesions are extremely common in daily clinical dermatological practice, but can be misleading for general practitioners unfamiliar with them.

The term “annular” is derived from the Latin word “ring”. The lesions appear as macules (changes in skin color) circular or ovoid, or plaques (solid content lesions more than 1 cm. of diameter in which predominate the extent and surface rather than deep) with an erythematous border and a clear center.

The most common cause of annular lesions found in the adult and child population is ringworm, which can be successfully diagnosed without a biopsy, in typical cases. However, conditions other than this, may present the same clinical appearance (Table 1).

Various mechanisms have been proposed to explain the annular configuration of lesions, not always satisfactory. One of the proposed mechanisms is based on irrigation, so that each round macula represent the territory irrigated by a single arteriole. Other authors consider that it could be centrifugal extension of a pathological process, whether infectious, neoplastic or allergic phenomenon.

Tinea Corporis

Ringworm is caused by fungi of the genus Microsporum, Trichophyton and Epidermophyton. The transmission thereof occurs by direct contact from person to person, from animals to humans and from the soil to animals and/or humans, depending if you try to anthropophilic, zoophilic or geophilic species [1].
It is characteristic in the skin of the trunk and extremities, restricted to the stratum corneum and most commonly occurs on exposed skin, although it can develop anywhere on the body.

It is most common in tropical regions like our country. Pets are an important factor in transmission, especially those zoophilic species. The incubation period is 1 to 3 weeks. The infection spreads centrifugally from the point of skin infection with a clear center, resulting in typical erythematous annular lesions of different sizes, with well circumscribed scaly, papular, vesicular or pustular edge. As they increase in size, the plates may have concentric rings (Fig. 1) [2,3].

Topical and oral imidazoles are the treatment of choice for ringworm. General measures are indispensable, such as avoiding moisture or maceration of the skin, weight loss and the use of cotton clothes [4,5].

**Pityriasis Rosea**

Common skin disease of children and young adults characterized by been a self-limited disease. It is usually preceded by upper tract respiratory symptoms [6].

Initially a heraldic plaque or mother plaque, typically located in the trunk, between 2 and 10 cm of diameter appears. The plaque is oval, erythematous, with a collarette desquamation on the periphery. After days, others plaques appear, similar in appearance to heraldic plaque but smaller in size, between 5 and 10 mm in diameter distributed throughout the trunk and less commonly in extremities, along the lines of division of the skin. If they are located on the back take on the appearance of a Christmas tree.

Skin biopsy usually is not indicated in these patients, although in borderline cases is performed. It shows a spongiform- subacute pattern with chronic non-specific inflammatory infiltrate, with apoptotic keratinocytes in the epidermis (especially in heraldic plaque). The picture is resolved between five to eight weeks from the beginning [4].

Rarely needs treatment, but if patients have accompanying pruritus, emollients such as zinc oxide, calamine lotions, and topical low power corticosteroids can be used [7]. Systemic antihistamines can also be used. In more severe cases treatments with ultraviolet light and natural sun exposure are described [4,8].

<table>
<thead>
<tr>
<th>Diagnosis</th>
<th>Clinical features</th>
<th>Histopathology</th>
<th>Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pityriasis roseae</td>
<td>Small salmon plates, with scales on the edges, along the lines of division of the skin. One is higher (herald).</td>
<td>Irregular epidermal acanthosis, spongiosis, diskeratosis (in heraldic plate), dermal hematic extravasation.</td>
<td>Topical or systemic corticosteroids; UVA; UVB.</td>
</tr>
<tr>
<td>Granuloma annulare</td>
<td>No scaly skin color patches, with raised indurated edges, extremities.</td>
<td>Necrobiosis area of dermal collagen surrounded by a palisade of inflammatory cells.</td>
<td>Spontaneously resolution or after biopsy.</td>
</tr>
<tr>
<td>Hansen’s disease</td>
<td>Hypochromic macules or erythematous plaques without scales. The edges will be more or less defined depending on the type of Hansen.</td>
<td>Lymphocytic (H1), granulomatous (HT, HBT) or macrophage (HBB, HBL, HL), periadnexal, perivascular and perineural infiltrate. ZN+for BAAR.</td>
<td>Dapsone, rifampicin, clofazimine.</td>
</tr>
<tr>
<td>Centrifugal annular erythema</td>
<td>Annular plate with scale leaves the erythematous border.</td>
<td>Superficial and deep perivascular dermal lymphocytic infiltrate. Variable skin alterations.</td>
<td>Topical or systemic corticosteroids. Oral antihistamines. Treating the underlying cause.</td>
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</tbody>
</table>

**Table 1: Main annular lesions that occur in childhood**

**Figure 1:** a) Tinea faciei, b) Tinea corporis. Erythematous annular lesions of different sizes, with well circumscribed scaly in the face and arms.
**Granuloma Annulare**

Self-limiting inflammatory entity of unknown etiology, characterized by annular skin color or slightly erythematous plaques, with elevated periphery and depressed center (Fig 2a and 2b), possibly result of non-specific reactions from various agents [9-11].

Located more often in the hands, feet, wrists and ankles, they can be located anywhere in the body. These lesions are usually asymptomatic, although some patients report mild itching. It is more frequent in females, predominantly in children and young adults. Four clinical forms are described: classical or localized, generalized, perforating and subcutaneous [12]. It has been linked to certain triggers such as: vaccination, insect bites, contact dermatitis to nickel, solar exposure, intestinal parasites and Ebstein-Barr virus, HIV, Herpes zoster virus infections among others. Is thought to be the result of vasculitis, necrobiosis secondary to trauma, activation of monocytes, Langerhans cell or a T cell phenomenon of type IV hypersensitivity. Some even believe there is a Koebner phenomenon [13].

Three histological varieties are distinguished: granulomatous (palisade), interstitial (incomplete) and deep (nodular) forms (Fig. 2c). They differ only in the location of granulomas. The granulomatous form is characterized by a palisade of histiocytes around degenerated collagen area where it can be mucin shown with special stains (Alcian blue or colloidal iron). It can be seen eosinophils and lymphocytes in the inflammatory infiltrate. In interstitial variant, macrophages are disposed between the collagen bands without forming palisade granulomas. The nodular form is characterized by deep subcutaneous granulomas with necrobiosis, but clinical presentation is not annular, it’s nodular [14].

The diagnosis is made by clinical and histopathologic correlation. In the absence of fever or other symptoms it is not necessary to request additional tests. Some of the patients with granuloma annulare can present hyperlipidemia, hiperggammaglobulineamia and circulating antinuclear antibodies. It can be ordered fasting and postprandial and determinations of glycated hemoglobin in patients with a family history of diabetes [4].

There is no successful treatment of the disease. Up to 50% of patients with localized forms may have spontaneous resolution in two years with a likely recurrence in 80% of cases, unlike the generalized form, where it is rarer and if there is not a spontaneous resolution it last at least 3 to 4 years.

In the limited forms it is not recommended any treatment. It is advisable to explain to parents the mildness of the disease, although if the therapeutic demand is strong, you can try the local treatment with topical corticosteroids, medium or high power, to avoid intralesional injection. In the deep or subcutaneous forms excision is indicated. In generalized forms dapsone (100 mg/day) can be used [4].

**Hansen’s Disease**

Leprosy is a chronic infectious disease caused by Mycobacterium leprae; as intracellular parasite triggers an immune response in the host, with the participation of cell-mediated immunity. The development of the disease is determined by two variables: a) opportunity for exposure to mycobacteria b) the ability of immune response that the host can offer.

The household contacts are the main source of infection. It has an incubation period of between 3 and 10 years.

The diversity in clinical manifestations will be conditioned by the play of interactions between M. leprae and the immune response offered by the host. And they may be: indeterminate, tuberculoid, dimorphic or borderline and lepromatous. Of these, the dimorphic form, which is a rare type even in childhood, is characterized by annular lesions tend to enclose areas of healthy skin and result in cut wafer images (Fig. 3). No mucous membranes or internal organs are affected. The smear is generally negative, because in the pediatric population pauci bacillary forms are the most frequent.
The histopathology evidence lymphohistiocytic perivascular and perineural dermal infiltrates (Fig. 4). With Zeehl - Neelsen staining AAR may be displayed, but not in the indeterminate or tuberculoid forms [4]. The dimorphic leprosy should be treated with 3 drugs (multibacillary).

**Urticaria**

Acute urticaria, annular urticarial or acute hypersensitivity syndrome represents an allergic hypersensitivity reaction mediated by histamine after viral or bacterial infections or after consumption of food or drugs. It is frequent in infants and children aged between 4 months and 4 years.

Most authors believe that this is not a single entity. In early childhood is often misdiagnosed as erythema multiforme or as serum sickness disease. Clinically, it begins as a pruritic maculopapules growing rapidly to form erythematous annular, polycyclic or arcuate evanescent character (disappear in less than 24 hours).

It may have a central clearing or acquire a ecchymotic tone, which simulates the target lesions of erythema multiforme, but there are not epidermal necrosis, blisters or mucosal involvement (Fig. 5a).

Histologically is characterized by dermal edema, vascular dilation, presence of perivascular cellular infiltrate composed of lymphocytes, mast cells, eosinophils and neutrophils and as diagnostic key there are many intravascular neutrophils. There are no fibrin extravasation or leukocytoclasia, findings in vasculitis, despite the large number of neutrophils in biopsies. The findings are most striking in superficial dermis (Fig. 5b).

The existence of dermografism, with erythema and edema in areas of trauma, is a regular feature of urticaria. Another typical manifestations of that entity is the appearance of angioedema in the face, hands and feet. It can be confused with serum sickness disease, however in this case the individual lesions are fixed and there is no dermografism associated [15,16].

It is a self-limiting rash with resolution of episodes in about 8-10 days, with good response to oral antihistamines, reserving the use of systemic corticosteroids for more severe cases [15].

**Centrifugal Anular Erythema**

It is considered a hypersensitivity reaction to different antigens. Its appearance is preferably in adults, although there are also cases reported in infants and children. In the pediatric population centrifugal annular erythema is more frequently associated with Candida albicans, dermatophytes, VEB and poxvirus infections. It can also occur in relation to most common cancers in children, such as leukemia and Hodgkin lymphoma [17].

Usually asymptomatic or poorly pruritic rash remits spontaneously in about two or three weeks, reappearing in the same locations or in other different at variable
Lesions, single or multiple, appear anywhere on the body, but preferably on the trunk and the root of the limbs, in the form of erythematous papules that migrate slowly (2-3 mm/day), flattening as it grows, fading at its center and leading to complete annular lesions or arc segments. There are two variants, one with peripheral scaly edge and intensely itchy, and the other with deeply infiltrated sharp edge without scale and asymptomatic [18].

As we already indicate, depending on the location of the perivascular infiltrate we found two variants; in the superficial type there is focal parakeratosis in the edge of the lesion, spongiosis and superficial perivascular lymphohistiocytic infiltrate, while in the deep form there is no epidermal changes, superficial and deep perivascular mononuclear cells, melanophages, slight vacuolization and necrotic keratinocytes in the dermo-epidermal junction [18].

As for treatment, antihistamines are effective when there is itching, especially in children. The use of antibiotics or antifungals, in the absence of demonstrated disease, has been useful in some isolated cases, as well as administration of systemic corticosteroids [19,20].

Subacute Cutaneous Lupus Erythematosus

The subacute cutaneous lupus erythematosus may present with polycyclic annular lesions. They can accompany a commitment systemic lupus erythematosus. This variant has pronounced photosensitivity and, regression leave a hypochromia without atrophy. The periphery can have blistering or crusting.

Histopathology of these lesions shows thickening of the basal membrane, hydropic degeneration of the basal layer, and superficial lichenoid lymphocytic infiltrate with melanophages (Figs 6a and 6b) [4]. They are very important measures designed to avoid exposure to sunlight, such as the use of sunscreens. Topical treatments include the use of corticosteroids and tacrolimus. For systemic treatment are useful hydroxychloroquine to 6.5 mg/kg/day, prior ocular control and every 4 to 6 months (color vision and visual field) or thalidomide at a dose of 50 to 100 mg/day [4].

CONCLUSION

1. The appearance of annular lesions in childhood is always a diagnostic challenge.
2. The clinical history, appearance and characteristics of lesions by the valuable contribution of histopathology, allow us to know the different entities and separate benign self-limiting pathologies that do not require treatment of those who needed.
3. For all of the above, when we confronted a ring, annular or figurative lesion in children we paust y attention to the age of the onset injury, lesion characteristics, duration and location, evolution and histological findings to establish a diagnostic orientation and an etiological treatment.

REFERENCES


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