Erythema Annulare Centrifugum = اﻠﻨاﺒذﺔ اﻠﺤﻠﻘﻴﺔ اﻠﺤﻤاﻤﻰ
Erythema Annulare

Centrifugum and

Other Figurate

Erythemas

The figurate erythemas include a variety of eruptions characterized by annular and polycyclic lesions. Classification of this group has always been controversial; the literature abounds with contradictions, uncertainties and a bewildering array of synonyms. Darier in 1916 was the first to use the term erythema annulare centrifugum
(EAC), although similar lesions had been described previously under other names. Table 42-1 lists the figurate erythemas and the differential diagnoses to consider.

### EPIDEMIOLOGY

EAC is an uncommon disorder. No epidemiologic data are available. There are only two large series in the literature: 66 cases identified clinically and 73 first diagnosed histologically. EAC appears to have no predilection for either sex or for any age group. Erythema gyratum perstans is an older synonym for familial EAC.

### ETIOLOGY AND PATHOGENESIS

Almost certainly EAC is not a single disease but a clinical finding with many causes. Both the annularity and the peripheral spread of EAC have attracted speculation as to a possible mechanism. Most hypotheses have centered around interactions among inflammatory cells, their mediators, and ground substance as foreign antigens diffuse through the skin.
CENTRIFUGUM AT A GLANCE

- Clinical pattern of annular expanding erythematous rings, which enlarge rapidly, fade, and then disappear, as new lesions appear.

- Diagnosis of erythema annulare centrifugum is one of exclusion.
  - Superficial and deep variants can be separated clinically and histologically. Deep form is often lupus tumidus.

- No single cause.

Migrating Erythemas

DISORDER

KEY FEATURES
CHAPTER

Erythema annulare centrifugum (EAC)

Slowly migrating lesions; often idiopathic.

This chapter

Erythema gyratum repens

Rapidly moving; usually cancer mark.

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Erythema migrans

Annular lesions originating from tick bite; skin sign of Lyme borreliosis.

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<table>
<thead>
<tr>
<th>Disease</th>
<th>Clinical Features</th>
</tr>
</thead>
<tbody>
<tr>
<td>Annular urticaria</td>
<td>is annular and overlaps with EAC; patients have ordinary urticaria elsewhere and more pruritus</td>
</tr>
<tr>
<td>Giant urticaria</td>
<td></td>
</tr>
<tr>
<td>Bullous pemphigoid</td>
<td></td>
</tr>
<tr>
<td>Early lesions often urticarial</td>
<td>and may be annular.</td>
</tr>
<tr>
<td>Erythema multiforme</td>
<td></td>
</tr>
<tr>
<td>Target lesions, usually acral</td>
<td>, often mucosal disease; some lesions annular.</td>
</tr>
<tr>
<td>Dermatophyte infections and tinea versicolor</td>
<td>Many fungal infections are annular (ringworm); they resemble superficial EAC but the scale contains...</td>
</tr>
</tbody>
</table>
Annular psoriasis

and occasionally ordinary psoriasis may have annular lesions.

Annular lupus erythematosus

Most common in neonatal forms; Ro/La antibodies should be especially looked for in Asian children.

Erythema marginatum

Transient, rapidly spreading annular erythema; specific for rheumatic fever.

Necrolytic migratory erythema
Marker for glucagonoma; erosive perioral and

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Carrier state chronic granulomatous disease

Female carriers may have erythematous -like rash.

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Hereditary lactate dehydrogenase M-subunit deficiency

Another rare genodermatosis with annular erythematosus and scale

This chapter

Familial annular erythema

Extremely rare.

This chapter
Annular erythema of infancy

Many different disorders must rule out neonatal lupus erythematosus.

This chapter

The cause of most cases of EAC is unknown. In one series, 24 patients were closely evaluated, and in none of the cases was any definite cause found. Bacterial or candidal infections are most often suspected as triggers. Autoimmune diseases are also occasionally associated.

The hormonal changes of menses or pregnancy have been implicated as well. Although medications are often identified as causing EAC in case reports, none regularly induces such lesions. EAC may be coupled with malignant neoplasms, with the eruption disappearing after treatment of the tumor and often returning as the tumor recurs. This paraneoplastic marker must be distinguished from metastatic tumors with an annular pattern.
CLINICAL FINDINGS

History

The history is most important in exploring the differential diagnostic considerations. In general the lesions are asymptomatic but may be cosmetically disturbing.

Cutaneous Lesions

EAC presents as one or more lesions that begin as erythematous macules or urticarial papules and enlarge by peripheral extension to form ringed, arcuate, or polycyclic figures. They spread gradually to form large rings with central clearing, with the edges of the lesions often advancing several millimeters a day. After a variable period of time, the lesions disappear, often to be replaced by new ones. In some cases annual recurrence has been described.

Two sub-types of EAC can be identified both clinically and histologically:

- Superficial gyrate erythema has an indistinct border and trailing scale and may be pruritic.

- Deep gyrate erythema has a firm yet indurated border, lacks scale, and is less often pruritic.
Additional findings may point to an underlying disorder, but most patients have no systemic signs or symptoms.

LABORATORY TESTS

Histopathology

Superficial EAC has epidermal changes of parakeratosis and spongiosis, with a superficial peri vascular infiltrate. Deep EAC lacks epidermal damage and has intense lymphohistiocytic cuffing about both the superficial and deep vessels. There is minimal papillary dermal edema and no spongiosis. One should search carefully for mucin as a clue to lupus erythematosus.

Other Laboratory Tests

There are no other laboratory tests diagnostic for EAC.
DIFFERENTIAL DIAGNOSIS

The differential diagnostic challenge in EAC is twofold. First one must exclude other eruptions that can appear similar, such as erythema migrans as the first sign of borreliosis or erythema gyratum repens, which is generally more rapidly moving and usually reflects an underlying malignancy. In addition, there are diseases that can clinically present as EAC, but biopsy will provide a different answer. Included in this group are leukocytoclastic vasculitis, granulomatous disorders, metastases, and eosinophilic infiltrates.

Then comes the problem of the many odd disorders that cause confusion, because they seem to have distinctive features but are rarely encountered. Prime examples are annular erythemas associated with the carrier state of chronic granulomatous disease or a lactate dehydrogenase M-subunit deficiency. There are also neutrophilic and lichenoid variants of EAC. Familial EAC, originally described as erythema gyratum perstans.
Erythema Annulare Centrifugum, is rare. Finally, there is the broad spectrum of annular erythemas of infancy, including neonatal lupus erythematosus, Malassezia furfur infections, and the idiopathic variants which themselves may show eosinophilic or neutrophilic infiltrates, as well as atrophy.

Box 42-1 Differential Diagnosis of Erythema Annulare Centrifugum (EAC)

**Most Likely**

- Dermatophyte infections
- Tinea versicolor
- Erythema migrans
- Annular urticaria
- Lupus erythematosus
- Lupus tumidus for deep EAC
- Subacute lupus erythematosus for superficial EAC

- Annular psoriasis

**Consider**

- Erythema multiforme
- Granulomatous diseases (granuloma annulare, actinic granuloma, sarcoidosis)
- Bullous pemphigoid (urticarial phase)
- Leukocytoclastic vasculitis (especially in children)
- Erythema marginatum
- Erythema gyratum repens
- Necrolytic migratory erythema
- Hypereosinophilic syndrome
- Carrier state chronic granulomatous disease
- Hereditary lactase dehydrogenase M-subunit deficiency
- Familial annular erythema
- Annular erythemas of infancy

Always Rule Out

- Lupus erythematosus
- Underlying tumor or annular metastasis

PROGNOSIS AND CLINICAL COURSE

EAC tends to be a chronic disease, which waxes and wanes. The course is determined by associated diseases or triggers.

TREATMENT
Although an assiduous search for the underlying cause is the primary goal of treatment, only symptomatic relief is available. Systemic glucocorticoids usually suppress EAC, but recurrence is common when these drugs are stopped. Systemic therapy with antipruritics may help. Topical vitamin D analogues, perhaps combined with ultraviolet irradiation, are another option. Empiric use of antibiotic, antifungal, or antifungal agents has sometimes been useful.

Biologics may represent yet another option. In general, most of the therapeutic approaches used for chronic urticaria can also be tried for EAC.