Granuloma Annulare

Granuloma annulare is a benign self-limited disease, first described by Colcott-Fox\(^1\) in 1895 and Radcliffe-Crocker in 1902.

EPIDEMIOLOGY

Granuloma annulare is a relatively common disorder. It occurs in all age groups but is rare in infancy. The localized annular and subcutaneous forms occur more frequently in children and young adults. The generalized variant is more common in adults. Many studies report a female
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preponderance, but some have found a higher frequency in males. Granuloma annulare does not favor a particular race or geographic area, with the possible exception of the perforating type, which may be more common in Hawaii.

Most cases of granuloma annulare are sporadic. Occasional familial cases are described with occurrence in twins, siblings, and members of successive generations. Attempts to identify an associated HLA sub-type have yielded disparate results in different population groups.

ETIOLOGY AND PATHOGENESIS

The etiology of granuloma annulare is unknown, and the pathogenesis is poorly understood. Most cases occur in otherwise healthy children. A variety of predisposing events and associated systemic diseases is reported, but their significance is unclear. It is possible that granuloma annulare represents a phenotypic reaction pattern with many different initiating factors.

Predisposing Events

Non-specific mild trauma is considered a possible triggering factor because of the frequent location of lesions on the distal extremities of children. An early study of subcutaneous granuloma annulare found a history of trauma in 25 percent of children, but this observation has not been replicated. Trauma is also a suspected factor in auricular lesions. Granuloma annulare has occurred after a cat bite and an octopus bite, and insect bite reactions have been implicated. There is a report of perforating granuloma annulare in long-standing tattoos. Widespread lesions have developed after waxing-induced pseudofolliculitis and erythema multiforme minor, and in association with systemic sarcoidosis. Severe uveitis without other evidence of sarcoidosis was found in eight patients with granuloma annulare.

INFECTIONS AND IMMUNIZATIONS

There are several reports of the development of granuloma annulare within herpes zoster scars, sometimes many years after the active infection. It is also described after chickenpox. Generalized, localized, and perforating forms of granuloma annulare may occur in
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association with human immunodeficiency virus (HIV) infection. Adenovirus was isolated from a lesion in one HIV-positive patient. Epstein-Barr virus was excluded as a causative agent in these cases. However, in other instances generalized granuloma annulare has been linked to viral infections, including Epstein-Barr virus infection, chronic hepatitis B, and hepatitis C. Vaccinations for tetanus, diphtheria toxoid, and hepatitis B vaccination have been implicated as triggering factors, although vaccination sites were spared in one case of generalized granuloma annulare.

Lesions compatible with granuloma annulare may occur in patients with active tuberculosis. There are also reports of granuloma annulare after tuberculin skin tests and bacille Calmette-Guérin immunization. Evidence of Borrelia burgdorferi infection was detected in two reports, but this association was not confirmed in a serologic study. A case in which chronic relapsing granuloma annulare flared during scabies infestation was attributed to the Koebner phenomenon.

SUN EXPOSURE

Granuloma annulare with a predilection for sun-exposed areas and seasonal recurrence has been described. Photosensitive granuloma annulare has been observed in patients with HIV infection. One patient developed generalized disease after psoralen plus ultraviolet A light therapy, but it is of note that phototherapy and psoralen/ultraviolet A phototherapy have also been used to treat generalized granuloma annulare.

Actinic granuloma, also known as annular elastolytic giant cell granuloma, develops on photodamaged skin and is believed to represent a granulomatous reaction to actinic elastosis. Its relationship to granuloma annulare is debated.

DRUGS

Granuloma annulare-like drug reactions are reported for gold therapy and treatment with allopurinol, diclofenac, quinidine, intranasal calcitonin, and amlodipine. An interstitial granulomatous drug reaction linked to the use of angiotensin-converting enzyme inhibitors, calcium channel blockers, and other medications is considered a distinct entity but may mimic
granuloma annulare.

**DIABETES MELLITUS AND THYROID DISEASE**

Development of granuloma annulare in patients with diabetes mellitus is extensively documented. Whether this is a true relationship has long been debated. The link is primarily with type 1 insulin-dependent diabetes, but cases are also reported with type 2 non-insulin-dependent disease. Localized and generalized as well as subcutaneous nodular and perforating forms of granuloma annulare have been observed. Granuloma annulare rarely pre-dates the onset of diabetes. The histopathologic similarity between granuloma annulare and necrobiosis lipoidica diabeticorum and the co-existence of both conditions in occasional diabetic patients suggest a true association. However, most patients with granuloma annulare do not have diabetes mellitus. Studies attempting to establish a causal correlation have yielded conflicting results.

Granuloma annulare has also occurred in a number of patients with thyroiditis, hypothyroidism, and thyroid adenoma.

**MALIGNANCY**

An association between granuloma annulare and malignancy in adult patients is reported primarily with Hodgkin and non-Hodgkin lymphoma, including mycosis fungoides, Lennert lymphoma, B-cell disease, and T-cell large cell lymphocytic leukemia. It is reported less commonly with solid tumors, particularly of the breast. The skin lesions of cutaneous lymphoma and other hematologic malignancies may mimic granuloma annulare both clinically and histopathologically. It may be difficult to distinguish whether they represent true granuloma annulare with atypical lymphocytes, or cutaneous lymphoma obscured by a granulomatous infiltrate.

Pathogenetic Mechanisms
The pathogenetic mechanisms that result in foci of altered connective tissue surrounded by a granulomatous inflammatory infiltrate are not understood. Proposed mechanisms include (1) a primary degenerative process of connective tissue initiating granulomatous inflammation, (2) a lymphocyte-mediated immune reaction resulting in macrophage activation and cytokine-mediated degradation of connective tissue, and (3) a subtle vasculitis or other microangiopathy leading to tissue injury.

CLINICAL FINDINGS

History

The typical history is of one or more papules with centrifugal enlargement and central clearing. These annular lesions are often misdiagnosed as tinea corporis and treated unsuccessfully with topical antifungal agents. Subcutaneous nodules may raise suspicion about malignancy or rheumatoid disease.

Granuloma annulare is usually asymptomatic. Mild pruritus may be present, but painful lesions are rare. Nodular lesions on the feet may cause discomfort from footwear. Cosmesis is often a concern for adolescent and adult patients, particularly with generalized disease.

Cutaneous Lesions

Clinical variants of granuloma annulare include the localized, generalized, subcutaneous, perforating, and patch types. Linear granuloma annulare, a follicular pustular form, and papular umbilicated lesions in children have also been described. There is overlap between the different variants, and more than one morphologic type may co-exist in the same patient.

LOCALIZED TYPE
The most common form of granuloma annulare is an annular or arcuate lesion. It may be skin colored, erythematous, or violaceous. It usually measures 1 to 5 cm in diameter. The annular margin is firm to palpation and may be continuous or consist of discrete or coalescent papules in a complete or partial circle. The epidermis is usually normal, but surface markings may be attenuated over individual papules. Within the annular ring, the skin may have a violaceous or pigmented appearance. Solitary firm papules or nodules may also be present. Papular lesions on the fingers may appear umbilicated.

The dorsal hands and feet, ankles, lower limbs, and wrists are the sites of predilection. Less commonly, lesions occur at other sites, including the eyelids. The palms and soles are occasionally involved. Localized annular lesions may co-exist with the subcutaneous or patch forms.

GENERALIZED TYPE

The generalized form of granuloma annulare is said to comprise 8 percent to 15 percent of cases. The majority of patients are adult, but it may be seen in childhood. Unlike in localized disease, the trunk is frequently involved, in addition to the neck and extremities. The face, scalp, palms, and soles may also be affected.

Generalized granuloma annulare presents as widespread papules, some of which coalesce to form small annular plaques or larger discolored patches with raised arcuate and serpiginous margins. Lesions may be skin colored, pink, violaceous, tan, or yellow. An annular or non-annular morphology may predominate. A generalized form of perforating granuloma annulare has also been described.

SUBCUTANEOUS TYPE

The subcutaneous form of granuloma annulare occurs predominantly in children but is also
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described in adult patients. It is characterized by firm to hard, usually asymptomatic nodules located in the deep dermis and subcutaneous tissues. They may extend to underlying muscle, and nodules on the scalp and orbit are often adherent to the underlying periosteum.

Individual lesions measure from 6 mm to 3.5 cm in diameter. They are distributed most often on the anterior lower legs in a pretibial location. Other sites of predilection are the ankles, dorsal feet, buttocks, and hands.\textsuperscript{93,95} Nodules on the scalp, eyelids,\textsuperscript{97-99} and orbital rim\textsuperscript{60,100,101} may present a diagnostic challenge. Subcutaneous granuloma annulare may also be found on the penis.\textsuperscript{102}

PERFORATING TYPE

The perforating type of granuloma annulare is a rare variant characterized by trans-epidermal elimination of the necrobiotic collagen. It may be localized, usually to the dorsal hands and fingers (see eFig. 43-3.1 in on-line edition), or generalized over the trunk and extremities. It has been described on the ears and within herpes zoster scars and tattoos. Superficial small papules develop central umbilication or crusting, and there may be discharge of a creamy fluid. Lesions heal with atrophic or hyperpigmented scars. In one series, 24 percent of patients complained of pruritus and 21 percent of pain. Papular umbilicated granuloma annulare on the hands of children and a generalized follicular pustular type of granuloma annulare\textsuperscript{85} may be clinical variants.

PATCH TYPE

Macular lesions that present as erythematous, red-brown, or violaceous patches without an annular rim are reported in adult women. An arcuate dermal erythema is also observed. Generalized confluent erythema has been described in an HIV-positive patient.

Related Physical Findings

Most patients with granuloma annulare are healthy and have no other abnormal physical findings. Arthralgia is reported in association with painful lesions on the hands. Granuloma
annulare-like skin lesions and joint disease characterize a multisystem disorder described as interstitial granulomatous dermatitis with arthritis.

Oral involvement was observed in one patient with HIV-associated disease.

LABORATORY TESTS

A diagnosis of localized annular granuloma annulare is made on clinical examination, and further evaluation is rarely indicated. Biopsy to obtain a specimen for histopathologic examination is necessary when the presentation is atypical, when lesions are symptomatic, and when the diagnosis is otherwise in doubt. Histopathologic analysis may be required to confirm a diagnosis of generalized granuloma annulare or subcutaneous nodular disease on the head and orbital region.

Histopathologic Findings

The diagnosis is best made at low magnification. Changes are usually observed in the upper and middle dermis, although any part of the dermis or subcutis can be involved. The characteristic histopathologic finding is a lymphohistiocytic granuloma associated with varying degrees of connective tissue degeneration and mucin deposition. The inflammatory infiltrate may have a palisaded or interstitial pattern, or a mixture of both patterns. Occasionally, a sarcoid-like pattern with large epithelioid histiocytes is seen.

The typical appearance is of single or multiple foci of inflammation with a central core of altered collagen (necrobiosis) surrounded by a wall of palisaded histiocytes. The necrobiotic centers are usually oval, slightly basophilic, devoid of nuclei, and marked by a loss of definition of the collagen bundles and diminished or absent elastic tissue fibers. Stains for mucin and lipid often give positive results.
Interstitial granuloma annulare = Interstitial granulomatous dermatitis

Differential Diagnosis of Granuloma Annulare

ANNULAR TYPE

Consider

- Tinea corporis
- Subacute cutaneous lupus erythematosus
- Neonatal lupus erythematosus
- Annular lichen planus
- Acute febrile neutrophilic dermatosis
- Erythema chronicum migrans
- Actinic granuloma
- Necrobiosis lipoidica diabeticorum

Rule Out

- Interstitial granulomatous dermatitis with arthritis
- Interstitial granulomatous drug reaction
- Annular sarcoidosis
- Lymphoma

GENERALIZED TYPE

Consider

- Lichen planus
- Lichen nitidus
- Molluscum contagiosum
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Rule Out

- · Lichenoid and granulomatous dermatitis of acquired immunodeficiency syndrome
- · Sarcoïdosis
- · Blau syndrome (familial granulomatous arthritis, skin eruption, and uveitis)
- · Interstitial granulomatous drug reaction
- · Lymphoma

SUBCUTANEOUS TYPE

Consider

- · Erythema nodosum
- · Dermoid cyst
- · Rheumatoid nodules

Rule Out

- · Epithelioid sarcoma
- · Benign or other malignant tumors
- · Deep infections

PERFORATING TYPE

Consider

- · Molluscum contagiosum
- · Insect bites
- · Pityriasis lichenoides
Interstitial granuloma annulare

- Perforating collagenosis and other perforating disorders
- Foreign body granuloma
- Papulonecrotic tuberculid
- Palisaded neutrophilic and granulomatous dermatitis of immune complex disease

PATCH TYPE

Consider

- Morphea
- Erythema annulare centrifugum
- Parapsoriasis

Rule Out

- Lymphoma

Immunofluorescence testing may show deposition of fibrin, immunoglobulin M, and C3 as a variable finding around vessel walls or at the basement membrane zone; immunoglobulin M cytoid bodies are also reported. Ultrastructural changes in the connective tissue and capillaries have been described.

Special Tests

A diagnosis of granuloma annulare is made clinically or by skin biopsy. Special investigations are usually not necessary. Further evaluation to rule out systemic disease such as infection, sarcoidosis, or malignancy may be required in atypical cases of granuloma annulare. Investigation for endocrine disease is indicated if the patient has signs or symptoms of diabetes or thyroid dysfunction.
Imaging studies may be performed in subcutaneous granuloma annulare when the clinical features are not recognized or when the presentation is atypical with rapid enlargement or pain. Radiographs show a non-specific soft tissue mass without calcification or bone involvement. Ultrasonographic examination reveals a hypoechoic area in the subcutaneous tissues. Magnetic resonance imaging shows a mass with indistinct margins, isointense or slightly hyperintense to muscle with T1-weighted images and with a heterogeneous but predominantly high signal intensity on T2-weighted images.\textsuperscript{112}

**TREATMENT**

Treatment Options for Granuloma Annulare

- · Await spontaneous resolution
- · Apply topical corticosteroid with or without occlusion

**ANECDOTAL REPORTS OF BENEFIT**

- · Topical

1. o Pimecrolimus cream
2. o Imiquimod 5% cream

- · Intraleesional

1. o Interferon-γ
2. o Interferon-β
3. o Sterile water or saline
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- · Systemic

1. o Antimalarials
2. o Retinoids
3. o Corticosteroids
4. o Cyclosporin
5. o Zileuton with vitamin E
6. o Chlorambucil, niacinamide, potassium iodide, fumaric acid esters, dapsone\(^3,43,83\)
7. o Etanercept
8. o Infliximab
9. o Efalizumab

- · Other

1. o Phototherapy
2. o Skin biopsy
3. o Cryotherapy
4. o Pulsed dye or CO\(_2\) laser

**CLINICAL COURSE AND PROGNOSIS**

Most cases of localized granuloma annulare resolve spontaneously without sequelae. Lesions may clear within a few weeks or persist for several years. The majority disappear within 2 years. Recurrent lesions may develop months or even years later, frequently at the same site. Generalized granuloma annulare often runs a more protracted course. Perforating granuloma annulare results in scarring. There is one report of anetoderma and mid-dermal elastolysis after generalized granuloma annulare.