Atopic dermatitis (AD) is a multisystem inflammatory disorder that is part of the spectrum of atopy, a series of conditions in which hyperreactivity and allergic symptoms are triggered by a series of causes including environmental allergens and irritants. Atopic dermatitis affects approximately one-quarter of children in developed countries and can have a negative impact on quality of life. In part 1 of this series addressing AD, the epidemiology and pathogenesis of AD are reviewed with an overview of skin barrier function. Later parts will address triggers, AD grading, differential diagnosis, comorbidities, and treatment options.

From Mount Sinai St. Luke's-Roosevelt Hospital and Beth Israel Medical Centers of the Icahn School of Medicine at Mount Sinai, New York, New York.

Dr. Silverberg has served as an investigator for Astellas Pharma US, Inc, and Novartis Corporation, and as a consultant for Anacor Pharmaceuticals, Inc; Johnson & Johnson Services, Inc; and Novartis Corporation.

This article is the first of a 3-part series. The second part will appear next month.

Correspondence: Nanette B. Silverberg, MD, 1090 Amsterdam Ave, Ste 11B, New York, NY 10025 (nsilverb@chpnet.org).

A topic dermatitis (AD), or eczema, is the leading dermatologic diagnosis worldwide and is vexing to patients due to the itchiness of the rash. It is the leading cause of skin disease burden worldwide with a prevalence of 229,761,000 reported cases in 2010, presenting largely in preadolescence but also persisting through adulthood. The prevalence of pediatric AD in the United States is estimated at more than 10% of children, with a 1.7 increased odds ratio in black children. Diagnosis generally is made based on the presence of a pruritic eczematous eruption with typical morphology and a personal and/or family history of atopy. Atopic dermatitis is caused by a complex interplay of skin barrier dysfunction and immune tendency toward allergy development.

Epidemiology of AD
The worldwide prevalence of AD varies by country and age group surveyed, with a higher prevalence in wealthy developed nations (eg, the United States) compared to poorer developing nations. Efforts to identify prevalence data for AD in the United States have been approached through a variety of strategies. A group in Oregon estimated the prevalence of AD in children aged 5 to 9 years to be 17.2% via a survey of parents (N=1465) and 11.8% with doctor-diagnosed

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eczema. In the same study, the question “Has a doctor ever said that your child has eczema?” was found to have a 91.3% predictive correlation.3 Analysis of the 2003 National Survey of Children’s Health demonstrated the overall US prevalence of pediatric AD to be 10.7% in 102,353 children 17 years or younger, with a range of 8.7% to 18.1% by region.6

In its evaluation of the worldwide prevalence of AD, the International Study of Asthma and Allergies in Childhood ranked the United States 17th.2,8 The prevalence of AD in developed countries such as the United States is fluid and is expected to increase if the trends from the last 20 years remain true. In an assessment of the National Health Interview Survey data from 1997 to 2011 based on responses to the question, “During the past 12 months, has your child had eczema or any kind of skin allergy?”, the Centers for Disease Control and Prevention identified an increase in the prevalence of AD in patients aged 0 to 17 years from 7.4% in 1997-1999 to 12.5% in 2009-2011.9 Rising prevalence seems to be paired with rising incidence in the total number of severe intractable cases, reduced clearance at the approach of grade school, or cases persisting into adulthood.

Racial Disparity in AD
Racial disparity worldwide and migration are thought to contribute to the prevalence of and therapeutic need for AD. For example, in the United Kingdom, the prevalence of AD in London-born Afro-Caribbean children versus white children (total cross-section, N = 693 junior school children) was 16.3% and 8.7%, respectively.10 In the United States, black children were more likely to have AD than white children (odds ratio, 1.7).6 Asian and black children also were more likely to present to a physician for treatment of AD than white children.6,10-13

Definition and Diagnostic Considerations
According to Hanifin,14 “Eczema represents a family of inflammatory skin conditions characterized by pruritic, papulovesicular, sometimes weeping dermatitis. All demonstrate the histological hallmark of spongiosis, which helps to distinguish the eczemas from papulosquamous diseases such as psoriasis.”14 Atopic dermatitis is a variant of eczema; however, most laymen identify eczema and AD as being one and the same.

The Hanifin and Rajka15 criteria are the major diagnostic criteria for AD but are difficult to use in clinical practice. Three of the following 4 major criteria are needed for diagnosis: (1) pruritus, which is present universally; (2) typical morphology and distribution; (3) chronic or chronically relapsing dermatitis; and (4) personal and/or family history of atopy. Additionally, 3 of the following 23 minor criteria are needed for diagnosis: xerosis; ichthyosis vulgaris, palmar hyperlinearity, or keratosis pilaris; positive skin prick test; elevated serum IgE level; early age of onset; tendency toward cutaneous infections or impaired cell-mediated immunity; tendency toward nonspecific hand or foot dermatitis; nipple eczema; cheilitis; recurrent conjunctivitis; Dennie-Morgan fold (infraorbital fold); keratocnosis; anterior subcapsular cataracts; orbital darkening; facial pallor or facial erythema; pityriasis alba; anterior neck folds; itching when sweating; intolerance to wool and lipid solvents; perifollicular accentuation; skin reactions from ingested foods or by food contact; environmental or emotional factors; and lesional/nonlesional white dermographism or delayed blanch.15-17

More pragmatic streamlined diagnostic criteria were established by Eichenfield et al.18 According to these guidelines, essential features for AD include pruritus and eczema. Important features seen in most cases and adding support to the diagnosis include early age of onset, atopy, and xerosis.18 In clinical practice, diagnosis is often made based on a pruritic relapsing condition in typical locations including the face, neck, and extensor surfaces in infants and children.

Age Considerations
Diagnosis of AD is made by 5 years of age in 85% to 90% of children who will develop the disease and by age 1 year in 60% to 65%.6,19,20 Atopic dermatitis will persist into adulthood in up to one-third of children.21,22 Infantile AD is characterized by erythematous, oozing, excoriated plaques on the cheeks (sparing the nose), scalp, trunk, and extensor surfaces. Pruritus is always seen in AD and can be a source of morbidity.16-18 Seborrheic dermatitis may complicate or overlap with AD in infancy.22

By 2 years of age, most children who are going to develop AD begin to show disease signs of childhood AD characterized by flexural lesions and lesions on the neck and in the postauricular area with sparing of the diaper area.23 Adult AD often presents as eczema of the hands and/or feet. Hand eczema in adulthood is correlated with a prior history of childhood hand eczema and/or childhood AD as well as wet work and caring for small children.24 Children with skin of color may manifest with follicular eczema as their primary disease phenotype. Facial and eyelid dermatitis are more common in Asian females, infants, and teenagers.12,25 Other disease phenotypes that are common in patients with skin of color include lichenoid AD and postinflammatory hypopigmentation.12
Pathogenesis of AD
There are 2 theories on the pathogenesis of AD known as the inside-out and outside-in hypotheses. The inside-out hypothesis suggests that allergic triggering leads to a weakened skin barrier that furthers allergen introduction and presentation, while the outside-in hypothesis suggests that the skin barrier is weakened in AD and allows for the presentation of allergens. Both theories have validity and biologic basis, and both may in fact be true in certain individuals.

The Skin Barrier: An Overview
The skin barrier is a complex set of factors present and functional at birth that seal the keratinocytes and the interkeratinocyte space so that the skin can perform key processes and functions including retention of fluid, exclusion of allergens, protection from UV light and solvents, and prevention of pathogen entry (eg, infections). The superficial stratum corneum or the cornified envelope consists of keratinocytes with intercellular strips of hydrophobic and hydrophilic substances formed by various intercellular lipids, largely ceramides, cholesterol, and free fatty acids. Keratinocytes are the first responders to a variety of environmental insults with the production of IL-18, RANTES (regulated on activation, normal T-expressed, and presumably secreted), granulocyte-macrophage colony-stimulating factor, and thymic stromal lymphopoietin. These inflammatory substances produce acute and chronic inflammation, mast cell reactivity, and T-cell activation. Corneodesmosins link the keratinocytes. Peptidases released will cleave the corneodesmosins and allow normal desquamation or shedding of surface skin, which is replaced by division of stem cells in the basal layer.

The stratum granulosum is the layer beneath the stratum corneum that co-contributes to barrier activity. The stratum granulosum is absent or reduced histologically in ichthyosis vulgaris, a form of skin dryness linked to filaggrin mutations and AD. Filaggrin breakdown creates natural moisturizing factor, a series of hygroscopic compounds that attract water into the skin. Histidine, a filaggrin breakdown product, is used by urocanic acid to process UV light insults. Filaggrin also contributes to other barrier functions including pH and stratum corneum cohesion as well as paracellular permeability of the stratum corneum. Tight junctions in the stratum granulosum include claudin-1 and claudin-6 and provide another barrier feature.

The skin barrier is composed of lipids and keratinocytes. Ceramides, which represent one type of lipids, are reduced in AD, causing alteration in the lamellar pattern and increased transepidermal water loss. Furthermore, the stratum corneum is thickened in AD, possibly in response to trauma, and hydration is reduced. Filaggrin (chromosome arm 1q21.3) is formed from the 400-kDa precursor profilaggrin through dephosphorylation and cleavage, and it performs an essential function in the skin barrier through its differential cleavage and breakdown as well as release of natural moisturizing factor and other compounds. Filaggrin mutations are linked to AD and ichthyosis vulgaris; however, barrier defects as evidenced by transepidermal water loss in the absence of filaggrin mutation are sufficient to allow for sensitization to allergens through the skin. Filaggrin mutations have been associated with AD development and vary in prevalence worldwide. In the United Kingdom, a prevalence study of filaggrin mutations in patients aged 7 to 9 years (N=792) demonstrated an 18.4% carrier rate in AD patients versus 12.9% in controls. A similar study in Sweden (N=3301) showed carrier rates of 13.5% versus 6.5%, respectively. Although filaggrin mutations are lower in black patients, ceramide content may be reduced in this population, demonstrating that a variety of skin barrier defects can result in AD. Carriers of filaggrin mutations are more likely to have eczema on skin exposed to environmental factors (eg, face, hands).

Barrier Defects Contributing to AD
The breakdown of the stratum corneum allows for antigen presentation to Langerhans cells, the dendritic antigen-presenting cells of the skin. Breaks in the stratum corneum may occur from scratching. These macroscopic breaks are large, whereas the breaks that otherwise occur due to barrier breakdown may be more microscopic in nature. Scratching causes aggravation of the helper T cell (TH2) response. For example, it allows the dendritic ends of Langerhans cells to be exposed to antigens. The dendritic ends capture allergens through IgE (may be elevated in AD), which is bound to the high-affinity FcER1 receptors on Langerhans cells. Rather than causing a type I hypersensitivity reaction, these Langerhans cells are activated and move to the lymph nodes where they present antigen and initiate a cascade of proinflammatory activity. This Th2 cascade includes release of cytokines such as IL-2, IL-4, IL-8, IL-10, tumor necrosis factor α, and IFN-γ.

Transepidermal water loss and barrier dysfunction contribute to disease activity and facilitate food/environmental allergen sensitization by allowing increased penetration of allergens through the skin to be presented by Langerhans cells to Th1 cells (sensitization phase). The Langerhans...
cells can reach their dendritic ends through tight junctions and into the stratum corneum, allowing them to reach surface allergens when the barrier is impaired. Ultimate expansion to systemic allergy (effector phase) occurs when dendritic cells move to draining lymph nodes, causing antigen presentation to CD4 and/or CD8 cells. Langerhans cells and dendritic cell sensitization through the weakened skin is believed to be the basis or role of barrier disruption as a trigger of atopic diseases, including AD and food and environmental allergies.

Many different forms of barrier disruption can cause a TH2 response in AD. The TH2 response triggers a constellation of proinflammatory activities including release of IL-4, associated with eosinophilia and elevated IgE levels, the latter being minor criterion in the diagnosis of AD. One mechanism by which the TH2 response is elicited may be the release of molecules such as danger-associated molecular patterns that may elicit recruitment of other inflammatory cells. Helper T cell (TH2) activity also can worsen barrier defects through IL-4 and IL-13 release, which can reduce filaggrin expression, and can aggravate barrier dysfunction in AD.

Inflammatory activation in AD also may involve inflammatory dendritic epidermal cells (IDECs). The IDECs can be tolerogenic or immunogenic mature phenotypes. The IDECs activate helper T cells (TH1), which may contribute to long-term AD activity.

Conclusion
Atopic dermatitis is a common skin condition worldwide and is characterized by the hallmark of pruritus and features that include a typical pattern, history of atopy (personal or family), and usually xerosis and early disease onset. Barrier dysfunction and immune dysregulation are prominent in AD, both of which aggravate the other and may encourage increased development of allergies and other forms of atopy over time.

REFERENCES


