Atopic dermatitis is associated with hidradenitis suppurativa diagnosis: A single institution retrospective cohort study

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Background: Hidradenitis suppurativa (HS) and atopic dermatitis (AD) are both chronic inflammatory skin diseases. An association between these 2 conditions can have important potential implications for elucidating pathogenesis, disease course, and treatment.

Objective: To investigate the association between AD and HS.

Methods: We performed a retrospective cohort study of patients seen at Duke University Medical Center from 2007 to 2017 who had AD compared with a control group without an AD diagnosis. The association of AD and HS was evaluated using a logistic regression model after adjusting for other confounders including age, sex, and race.

Results: Of 28,780 patients with an AD diagnosis, 325 (1.1%) were diagnosed with HS compared with 76 (0.2%) within the 48,383 patients in the non-AD group. An adjusted logistic regression model demonstrated an increased odds ratio of having HS diagnosis in the AD group as compared with the control non-AD group (odds ratio: 5.57, 95% confidence interval: 4.30-7.21, P < .001).

Limitations: This was a retrospective study performed at a single institution with the possibility of surveillance bias being present.

Conclusions: Patients with AD are more likely to be diagnosed with HS than patients without AD. Further research is needed to understand the pathophysiologic mechanism and potential treatment implications.

Key words: atopic dermatitis; barrier defect; clinical research; hidradenitis suppurativa; inflammatory skin diseases; notch signaling.

INTRODUCTION

Hidradenitis suppurativa (HS) is a chronic relapsing inflammatory condition that presents in the second or third decade of life and affects approximately 0.1%-0.7% of the population.1,2 It is diagnosed clinically based on specific criteria that include...
the occurrence of chronic, recurrent, deep-seated, painful nodules and draining tunnels in 1 or more predilection skin areas, such as the axilla and groin. Flares of this chronic cutaneous condition are extremely painful, and the acute presentation of these episodes has led to an increased utilization of health care delivery in high-cost settings, such as the emergency department and inpatient hospitalizations.3 Risk factors for HS include smoking, female sex, and African American race.4,5 Furthermore, HS is frequently associated with multiple comorbidities, such as inflammatory bowel disease, metabolic syndrome, obesity, and polycystic ovary syndrome.6,7

Similar to HS, atopic dermatitis (AD) is also a chronic relapsing inflammatory skin disease. This condition affects approximately 20% of the population, with 85% of patients having AD present before 5 years of age.6 AD typically peaks in infancy and in the third decade of life. AD results in pruritic lesions that commonly arise on the elbows, chest, face, neck, and ankles. AD is usually the first clinical manifestation of atopy and is commonly associated with asthma, allergic rhinitis, and food allergies. Similar to HS, risk factors associated with AD include African American race, female sex, and smoking.9-11 AD is known to result in an increased delivery of health care, and a recent study determined that AD’s health care resource utilization was significantly higher when compared with that of non-AD controls, mainly because of health care provider visits and associated costs.12

We are not aware of any published clinical data indicating a link between the 2 diseases. However, based on our clinical experience in our hidradenitis specialty clinic, in addition to the aforementioned factors, we postulated that AD is associated with HS.

METHODS

Data sources and study population

Patients who had AD diagnosed at Duke University Medical Center between November 1, 2007, and November 1, 2017, were identified from the Duke Enterprise Data Unified Content Explorer database, a web-based environment that allows Duke-based investigators access to information generated during patient care within the Duke University Health Care System. A random cohort of 50,000 patients seen at Duke University Medical Center during the same time period were also extracted from the Duke Enterprise Data Unified Content Explorer database and used as a control group. Identified AD and non-AD cohorts were evaluated using electronic health records going back to 1997. Age was calculated based on the date of the first encounter.

Patients with AD were identified using the International Classification of Diseases Ninth and Tenth Revisions (International Classification of Diseases [ICD 9/10]) codes 691.8 and L20.9 for atopic dermatitis and related conditions and ICD diagnosis codes 692.9 and L25.9 for contact dermatitis and other eczema, unspecified cause. These codes were previously validated by Hsu et al.13 In order to increase the positive predictive value (PPV) for the diagnosis of AD, we only considered a patient to have validated AD if they also had a concomitant diagnosis of either asthma, allergic rhinitis, or food allergies with the caveat that a patient needed a minimum of 2 diagnoses of allergic rhinitis to be considered to have allergic rhinitis.13,14 In validating our own cohort, 50 patients were randomly selected from our AD cohort identified based on ICD codes. Chart reviews were performed in order to determine the AD diagnosis by using the UK criteria. This criterion specifically included history of flexural involvement, history of dry skin, onset under the age of 2, personal history of asthma, history of a pruritic skin condition, and visible flexural dermatitis to determine if a patient had the diagnosis of AD.15 We observed a PPV of 74% for the diagnosis of AD using this method, which is consistent with the range of 68%-80% previously reported in the literature.13,15 Of note, we excluded patients with a syndrome or deficiency that increases the risk for atopic dermatitis, including Jobs syndrome, Wiskott–Aldrich syndrome, ataxia telangiectasia, and Vitamin B3 deficiency.17,19 After determining the validity of this selection criterion, we identified a cohort of 28,780 patients with a diagnosis of AD (Fig 1, A).

For the non-AD control group, we proceeded to exclude patients with validated AD and any syndromes that increase the risk of AD as mentioned previously and were left with a cohort of 48,383 patients (Fig 1, B). We then identified patients with a diagnosis of HS from the non-AD control cohort and the AD cohort.
Abbreviations used:
AD: atopic dermatitis
AMP: antimicrobial peptide
CI: confidence interval
HS: hidradenitis suppurativa
ICD: International Classification of Diseases
OR: odds ratio
PPV: positive predictive value
TSLP: thymic stromal lymphopoietin

We used ICD-9/10 codes 705.83 and L73.2 to identify patients of all ages with a diagnosis of HS from this cohort. In validating our own cohort, 50 patients were randomly selected from our HS cohort identified based on ICD codes. We used the following criteria, based on previous studies, to validate the HS diagnosis: confirmation by a dermatologist, pathology results, or accuracy of HS lesion description. We observed a PPV of 76% for the diagnosis of HS by using a single ICD-9/10 code. This result is consistent with the previous finding by Strunk et al showing that the presence of at least one ICD-9 code for HS provides a 79% PPV in confirming HS diagnosis. The initial diagnosis date for HS was identified as the diagnosis date for boils in patients who initially presented with the ICD-9/10 code. For these 76 patients, the median age at HS diagnosis was 29 years (Q1-Q3: 20.2-41.2). The unadjusted odds of having an HS diagnosis for patients who had AD was significantly higher than that for patients in the non-AD control group (OR = 7.26, 95%CI: 5.65-9.32, P-value < .001).

A multivariable logistic regression model was used to study the effect of AD on the odds of having HS after adjusting for age, race, and sex and is summarized in Table II. The adjusted model demonstrated an increased OR of having an HS diagnosis in the AD group as compared with the non-AD control group (OR: 5.57, 95% CI: 4.30-7.21, P < .001), indicating that patients with AD were more likely to have HS (Table II). Furthermore, we found sex and race were significantly associated with the incidence of HS with an increased odds of having HS for female compared with male patients (OR: 3.36, 95% CI: 2.59-4.36, P < .001) and for African American race compared with Caucasian (OR: 3.52, 95% CI: 2.80-4.43, P < .001) as expected.

DISCUSSION
We determined that patients with a diagnosis of AD have an approximately 5.57-fold increased OR of having HS as compared with those who do not have AD. This relationship is highly significant and suggests a potential unique pathophysiologic link that may be related to similar deficient notch signaling, barrier defect and antimicrobial peptide (AMP) dysregulation seen in both conditions. This association may also reflect a common genetic susceptibility shared between AD and HS. In clinical practice, a history of atopy should be considered when deciding on the frequency and use of various topical antiseptic washes typically recommended for patients with HS. Patients should be carefully monitored for clinical signs of eczema in areas affected by HS and treated accordingly. Exacerbation of barrier defect in these areas can also worsen HS by contributing to ongoing inflammatory responses. Future prospective studies are needed to understand whether hidradenitis patients with a concomitant AD diagnosis demonstrate a different disease course and treatment response compared with those of patients with hidradenitis alone. The theoretical similarities in pathophysiology need further confirmation and
could result in treatment options with similar therapeutic targets. We therefore suggest several testable hypotheses that can be considered to explain this new observation of an increased association of HS in AD patients. Recent evidence suggests that in cases of familial HS, there are loss-of-function mutations in the gamma secretase complex, which may lead to decreased proteolytic cleavage of transmembrane notch protein to its active form. The role of impaired notch signaling in HS is unclear, but in AD, notch signaling dysregulation can result in epidermal barrier defects and increased inflammation. Specifically, notch signaling plays a key role in the regulation of skin differentiation (epidermis and pilosebaceous unit) and is important in the processing of filaggrin. Mouse models with genetically suppressed notch signaling have shown AD-like skin disease and increased transepidermal water loss. Notch signals are also important in the differentiation of regulatory T cells and play a role in the feedback inhibition of innate immunity. In particular, notch dysfunction has been associated with increases in thymic stromal lymphopoietin (TSLP) in multiple disease processes including AD. Recent studies have suggested that TSLP alone is sufficient to promote the Th2 phenotype, but a combination of TSLP and toll-like receptor 3 is required to promote simultaneous Th17 phenotype, a unique profile seen in severe asthma. In HS, there is clear evidence suggesting an increased presence of Th17 cells in lesional skin in addition to an upregulation of the Th2 pathway as well as the Th1 pathway. We suspect that a combination of TSLP increase secondary to defective notch signaling, as well as other inflammatory mediators, maybe be contributing to the inflammatory cascade witnessed in HS, although this has not

![Fig 1. Atopic dermatitis. Flow diagram of cohort selection of patients with (A) and without (B) validated atopic dermatitis. AD, Atopic dermatitis.](image-url)
yet been examined. Such findings may provide a potential link in the association between AD and HS. In addition, AMP dysregulation in the setting of altered skin flora acts to promote an inflammatory response in both AD and HS; specifically, both diseases are associated with altered levels of the AMPs such as cathelicidin (LL-37) and dermcidin.\textsuperscript{39,40} In both AD and HS, a distinct wound response signature similar to what is seen at injury sites, including an upregulation of AMPs, has also been reported.\textsuperscript{23,41} In circumstances in which AMP levels were altered, the resulting phenotypic changes were shown to lead to an increase in local inflammatory mediators, further propagating tissue inflammation and damage.\textsuperscript{39,42,43} Therefore, the exact role of AMP dysregulation, similar in both AD and HS, needs to be further elucidated.

Several limitations are present that warrant consideration when interpreting the results of our study. This was a retrospective study performed at a single institution. However, the study did examine a heterogeneous population that is mostly representative of the North Carolina population. Additionally, patients may have moved throughout their lifetime and received health care from multiple medical centers. Therefore, the association between AD and HS may be even stronger, as many of our patients could have had a diagnosis of AD during childhood but did not access health care at Duke University Health Care System until a later age. Another important limitation that warrants consideration is the fact that surveillance bias may be present in our study, as patients who had AD managed by dermatologists may possibly have had a higher chance of being diagnosed with HS as compared with that in the patients in the control group. Unfortunately, we do not have provider information of encounters for each patient over time; therefore, we cannot accurately estimate the percentage of patients being seen by a dermatologist for each of the 2 cohorts (AD vs non-AD control cohort). However, the association between AD and HS was independent of the timing of the AD diagnosis.

In the future, our goal is to repeat this study with a larger US population using Truven Health MarketScan research database. Despite these limitations, this study described an association between AD and HS, as demonstrated by the increased incidence of HS witnessed in patients with AD.

### Conflicts of interest

Dr Macleod consulted for Silab when she was an employee at Duke. The MacLeod laboratory has previously received funds from Silab Company; funding from this partnership was not directly used for this study. Silab did not have any influence on the content of this project. Dr Macleod is also consulting for the LEO Foundation. The spouse of Dr Macleod is employed at Precision Biosciences and has stock options. Dr Jaleel is an investigator for UCB and reported consulting for Eli Lilly and Chemocentryx and receiving honoraria. None of the content or the decision to publish has been affected by the authors’ involvement with Eli Lilly, Silab, Chemocentryx, Precision Biosciences or the LEO Foundation. Drs Kaakati, Tanaka, Liu, Ward, and Green have no conflicts of interest to declare.

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