Keloid risk in patients with atopic dermatitis: a nationwide retrospective cohort study in Taiwan

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ABSTRACT

Objective The pathogenesis of keloid is largely unknown. Because keloid and atopic dermatitis have overlapping pathophysiological mechanisms, we aimed to evaluate keloid risk in patients with atopic dermatitis.

Study design Population-based retrospective cohort study.

Setting The Taiwan National Health Insurance Research Database was used to analyse data for people who had been diagnosed with atopic dermatitis.

Participants We identified 8371 patients with newly diagnosed atopic dermatitis during 1996–2010. An additional 33,484 controls without atopic dermatitis were randomly identified and frequency matched at a one-to-four ratio.

Primary and secondary outcome measure The association between atopic dermatitis and keloid risk was estimated using Cox proportional hazard regression models.

Results After adjustment for covariates, the atopic dermatitis patients have a 3.19-fold greater risk of developing keloid compared with the non-atopic dermatitis group (3.19 vs 1.07 per 1000 person-years, respectively). During the study period, 163 patients with atopic dermatitis and 532 patients without atopic dermatitis developed keloid. Notably, keloid risk increased with severity of atopic dermatitis, particularly in patients with moderate to severe atopic dermatitis.

Conclusions Our results indicate that patients with atopic dermatitis had a higher than normal risk of developing keloid and suggest that atopic dermatitis may be an independent risk factor for keloid.

INTRODUCTION

Keloid is a pathological disorder as a result of dermal injury and excess production in the extracellular matrix (ECM) (mainly type I collagen and proteoglycans). Keloids are characterised by concomitant annoying symptoms such as pain and pruritus. The recurrence rate after surgical excision is high. The pathogenesis of keloids is largely unknown,1–3 and no standard treatment has emerged. Although the cause of this disorder is not well understood, known risk factors for keloids are age 10–30 years, female gender,1,2 and dark skin pigmentation (eg, in Asian, African-American and Hispanic people). Notably, the risk of keloid development is highest in populations with the darkest complexions.4,5

In a recent case–control study performed in Northeast China, comparisons of blood test results for 283 subjects with keloids and 290 controls suggested that keloid development may be associated with the single-nucleotide polymorphisms, a disintegrin and metalloprotease 33 (ADAM33).6 Earlier studies have also shown that ADAM33 is associated with immune-mediated disorders such as atopic dermatitis (AD).7,8 In the first epidemiological study of the relationship between ADAM33 gene polymorphisms and AD risk, Matsusue et al reported a significant association between ADAM33 gene polymorphisms and AD in comparisons of 140 children with AD and 258 healthy controls recruited from June 2006 to January 2007. We hypothesised that AD and keloids may develop through common pathways, particularly in the ADAM33-related gene. Since no population-based data were available for investigating the association between AD and keloids, this study retrospectively analysed data from the Taiwan National Health Insurance Research Database.
Insurance Research Database (NHIRD) to investigate this relationship.

**MATERIALS AND METHODS**

**Source of data**

The data used in this population-based cohort study were obtained from the NHIRD established by the Taiwan national healthcare system. This encrypted secondary database, which was implemented in March 1995 contains medical reports for almost 99% of the 23.74 million residents of Taiwan. The NHIRD contains records of medical services and enrolment data for all beneficiaries. The database is publicly available to researchers, but its use is strictly regulated by Taiwan Personal Electronic Data Protection Law. The NHIRD has been used in many studies published in scientific journals. The current study used the Longitudinal Health Insurance Database 2010, in which diseases are coded according to the International Classification of Diseases, Ninth Revision, Clinical Modification (ICD-9-CM).

The study was proceeded according to Declaration of Helsinki guidelines. Because all NHIRD data had been deidentified and personal information traced was anonymised before data analysis, informed consent was not needed.

**Study design**

We examined medical claims and identified 8371 patients who had at least one AD diagnosis (ICD-9-CM code 691.8) during 1996–2010. To ensure accurate data, the analysis was limited to patients who had received ≥3 AD diagnoses during outpatient visits or ≥1 AD diagnoses during inpatient medical service. The date of the first clinical visit for AD was defined as the index date. Patients with moderate to severe AD were defined as patients who had received systemic corticosteroids. The exclusion criterion was any diagnosis of a keloid (ICD-9-CM code 701.4) before the index date. During the follow-up period, the occurrence of keloid was defined as ≥3 keloid diagnoses in outpatient visits or ≥1 keloid diagnoses in inpatient medical service. The two cohorts were limited to patients who had an ICD-9 code for keloid assigned by a dermatologist. Another 33,484 controls without AD were randomly identified after eliminating study case cohort and frequency matched at a one-to-four ratio for age, gender and index year (year of AD diagnosis). Both cohorts were followed up until 31 December 2010, or until the occurrence of keloids. Figure 1 shows a flow chart of the study procedure.

**Ethics approval**

The study was conducted in accordance with the Declaration of Helsinki guidelines.

**Measures and definition**

As in previous NHIRD studies, the patients were classified into three urbanisation levels according to their residence addresses recorded in the databases: urban (those living in the ‘city’), suburban (those living in ‘Jeng’ areas) and rural (those living in ‘Xiang’ areas). In addition to demographic risk factors (urbanisation level, gender and age), concurrent allergic diseases identified by ICD-9-CM codes in the claims records before the index date included asthma (ICD-9-CM code 493), allergic rhinitis (ICD-9-CM code 477) and allergic conjunctivitis (ICD-9-CM code 372.05, 372.10 and 372.14). Baseline comorbidities were also identified by ICD-9-CM codes,

![Flow diagram summarising the process of the enrolment. LHID, Longitudinal Health Insurance Database.](http://bmjopen.bmj.com/content/8/8/e022865)

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**Figure 1** Flow diagram summarising the process of the enrolment. LHID, Longitudinal Health Insurance Database.
including hypertension (ICD-9-CM codes 401–405), chronic obstructive pulmonary disease (ICD-9-CM code 491, 492, 494 and 496), hyperlipidaemia (ICD-9-CM code 272), chronic kidney disease (ICD-9-CM code 582, 583, 585, 586 and 588), diabetes mellitus (ICD-9-CM code 250), chronic liver disease (ICD-9-CM code 571), stroke (ICD-9-CM code 430–438), mental disorders (ICD-9-CM code 290–319), coronary artery disease (ICD-9-CM code 410–414) and alcohol-attributed disease (ICD-9-CM codes 291.0–9, 303, 305.0,357.5, 425.5, 535.3, 571.0–3980.0 and V11.3). Concurrent allergic diseases and comorbidities were ascertained at least more than two outpatient claims during study periods.

Statistical analysis
We used \(\chi^2\) test to compare distributions of categorical demographic and clinical characteristics between the AD and non-AD control cohorts. Student’s \(t\)-test and Wilcoxon rank-sum test were used to compare mean age and follow-up time (y) between the two cohorts, as appropriately. In the patients with AD, survival time was calculated until hospitalisation, an ambulatory visit for keloids or the end of the study period (31 December 2010), whichever came first. Keloid incidence rates per 1000 person-years were compared between the two cohorts. Kaplan-Meier analysis was used to estimate cumulative incidence of keloids, and the differences between the curves were compared by two-tailed log-rank test. Univariable and multivariable Cox proportional hazard regression models were used to calculate HRs and 95% CIs for keloids in patients with AD when the proportional hazards assumption was satisfied. Multivariate Cox models were adjusted for age, gender, urbanisation level, concurrent allergic diseases and relevant comorbidities. Statistical Analysis Software, V.9.4 was used for data analyses; a \(p\) value less than 0.05 in a two-tailed test was considered statistically significant.

Patient and public involvement
There were neither patients nor the public involved in this study.

Results
Table 1 shows the demographic characteristics and comorbidities of patients in the AD and non-AD cohorts. In the AD cohort, 50.71% patients were younger than 40 years old, and 64.81% patients were female. Compared with the non-AD cohort, the AD cohort had significantly higher percentages of patients with asthma (25.41 vs 12.91, \(p<0.001\)), allergic rhinitis (54.78 vs 35.92, \(p<0.001\)), allergic conjunctivitis (57.22 vs 42.08, \(p<0.001\)), hypertension (34.09 vs 21.60, \(p<0.001\)), diabetes mellitus (23.51 vs 13.46, \(p<0.001\)), hyperlipidaemia (34.56 vs 21.08, \(p<0.001\)), chronic kidney disease (12.03 vs 6.20, \(p<0.001\)), chronic liver disease (35.83 vs 21.98, \(p<0.001\)), chronic obstructive pulmonary disease (26.74 vs 15.08, \(p<0.001\)), stroke (5.85 vs 2.97, \(p<0.001\)), coronary artery disease (5.02 vs 2.71, \(p<0.001\)), mental disorders (18.05 vs 11.33, \(p<0.001\)) and alcohol-attributable disease (3.72 vs 2.28, \(p<0.001\)). During a median observation time of 1.9 years, 1.95% (163) of the patients with AD had keloids (IQR=0.8–4.3). The incidence of keloids was significantly (\(p=0.022\)) higher in the AD cohort compared with the non-AD cohort (532 patients with keloids out of 33,484 controls matched for age, and gender (1.59%)) during a median observation time of 9.0 years (IQR=6.2–11.8)). The median duration of follow-up for keloids was significantly shorter in the AD group (1.9 years) compared with the non-AD group (9.0 years).

Table 2 stratifies the keloid incidence rates and HRs by age, gender and concurrent allergic disease or comorbidity. During the follow-up period, keloids developed in 1.95% (163) of the patients with AD and in 1.59% (532) of the patients without AD. The overall keloid risk was 3.19 times greater in the AD group compared with the non-AD group (3.19 vs 1.07 per 1000 person-years, respectively) after adjusting for age, gender, urbanisation level, concurrent allergic diseases (asthma, allergic rhinitis and allergic conjunctivitis) and related comorbidities (hypertension, hyperlipidaemia, diabetes mellitus, chronic liver disease, chronic obstructive pulmonary disease, chronic kidney diseases, stroke, mental disorders, coronary artery disease and alcohol attributable disease).

In both the AD and non-AD cohorts, gender-specific analyses showed that the incidence of keloids was higher in women than in men (3.52 and 2.61 per 1000 person-years, respectively, in the AD cohort; 1.27 vs 0.69 per 1000 person-years, respectively, in the non-AD cohort). However, men in the AD group had a significantly higher keloid risk compared with men in the non-AD group (adjusted HR=3.67, 95% CI 2.47 to 5.44) and women in the AD group had a significantly higher keloid risk compared with women in the non-AD group (adjusted HR=3.03, 95% CI 2.39 to 3.83).

In all age groups, the AD cohort also had a significantly higher incidence of keloids compared with the non-AD cohort. Age-specific risk comparisons showed that compared with the non-AD cohort, the AD cohort had a significantly higher keloid risk in all age groups, and the risk was highest in patients under 40 years old. Keloid risk was higher in patients with AD than in patients without AD, regardless of concurrent allergic disease or comorbidity.

Table 3 shows that keloid risk increased with severity of AD, particularly in patients with moderate to severe AD.

Figure 2 compares the Kaplan-Meier curves for the cumulative incidence of keloids between the AD and non-AD groups at the 15-year follow-up. The baseline risk of keloid development was significantly higher in the AD group (2.83%, \(p<0.001\)) compared with the non-AD group (1.59%). The Kaplan-Meier curves also showed a significantly higher cumulative incidence of keloids in the AD group compared with the non-AD group (log-rank test, \(p<0.001\)).
The above comparisons supported our study hypothesis that patients with AD have a higher than normal risk of subsequently developing keloids. This study is the first to demonstrate an association between AD and keloids in an Asian population. The keloid risk was 3.19-fold higher in the AD group compared with the non-AD group. Notably, the females in the AD group had a significantly higher incidence rate of keloids compared with males. Specifically, compared with patients in the non-AD group, the AD group had a significantly higher keloid risk in all age groups, especially in those less than 40 years old. In the AD group, keloid risk was significantly associated with the presence of comorbidities and concurrent allergic diseases and keloid risk increased with the severity of AD.

Previous studies of the association between AD and keloids have obtained inconsistent results. For example, a case–control study of 131 patients with keloids and 258 controls recruited during 2000–2012 revealed no association between AD and keloids in the adjusted model (adjusted OR 1.27; 95% CI 0.76 to 2.13). However, the questionnaire-based reporting method used in that study may have caused information bias resulting in under-diagnosis of AD. Another limitation is that the authors did not report AD-compatible itchy flexural rash with AD, which may have also contributed to underdiagnosis of AD. Underdiagnosis of keloids is also common because patients with keloids are often immigrants, who may be unaccustomed to seeking prompt medical attention for non-critical medical problems. Additionally, the diverse ancestry of the population analysed in Hajdarbegovic et al. may have had a confounding effect on the association between AD and keloids. A final limitation of Hajdarbegovic et al is its small sample size.

**Table 1** Baseline characteristics of patients with and without atopic dermatitis

| Variables                                         | Atopic dermatitis | P values |
|---------------------------------------------------|-------------------|----------|
| Keloid patients, n (%)                            | Yes (n=8371)      | No (n=33484) | 0.022    |
| Period of developing keloid, median (IQR), years  | 1.9 (0.8–4.3)     | 9.0 (6.2–11.8) | <0.001   |
| Mean age at diagnosis of keloid, years            | 33.1 (14.8)       | 36.8 (14.4) | <0.001   |
| Age group, n (%)                                  |                   |           |
| <40                                               | 4245 (50.71)      | 16980 (50.71) |         |
| 40–59                                             | 2601 (31.07)      | 10404 (31.07) |         |
| ≥60                                               | 1525 (18.22)      | 6100 (18.22) | 1.000    |
| Gender, n (%)                                     |                   |           |
| Men                                               | 2946 (35.19)      | 11784 (35.19) |         |
| Women                                             | 5425 (64.81)      | 21700 (64.81) | 1.000    |
| Degree of urbanisation, n (%)                     |                   |           |
| Urban                                             | 5267 (62.92)      | 20463 (61.11) |         |
| Suburban                                          | 1332 (15.91)      | 5665 (16.92) |         |
| Rural                                             | 1772 (21.17)      | 7356 (21.97) | 0.008    |
| Concurrent allergic disease, n (%)                |                   |           |
| Asthma                                            | 2127 (25.41)      | 4323 (12.91) | <0.001   |
| Allergic rhinitis                                 | 4586 (54.78)      | 12028 (35.92) | <0.001   |
| Allergic conjunctivitis                           | 4790 (57.22)      | 14089 (42.08) | <0.001   |
| Comorbidity, n (%)                                |                   |           |
| Hypertension                                      | 2854 (34.09)      | 7232 (21.60) | <0.001   |
| Diabetes mellitus                                 | 1968 (23.51)      | 4508 (13.46) | <0.001   |
| Hyperlipidaemia                                   | 2893 (34.56)      | 7059 (21.08) | <0.001   |
| Chronic kidney disease                            | 1007 (12.03)      | 2076 (6.20) | <0.001   |
| Chronic liver disease                             | 2999 (35.83)      | 7361 (21.98) | <0.001   |
| Chronic obstructive pulmonary disease             | 2238 (26.74)      | 5050 (15.08) | <0.001   |
| Stroke                                            | 490 (5.85)        | 994 (2.97) | <0.001   |
| Coronary artery disease                           | 420 (5.02)        | 909 (2.71) | <0.001   |
| Mental disorders                                  | 1511 (18.05)      | 3794 (11.33) | <0.001   |
| Alcohol attributed disease                        | 311 (3.72)        | 765 (2.28) | <0.001   |

**DISCUSSION**

The above comparisons supported our study hypothesis that patients with AD have a higher than normal risk of subsequently developing keloids. This study is the first to demonstrate an association between AD and keloids in an Asian population. The keloid risk was 3.19-fold higher in the AD group compared with the non-AD group. Notably, the females in the AD group had a significantly higher incidence rate of keloids compared with males. Specifically, compared with patients in the non-AD group, the AD group had a significantly higher keloid risk in all age groups, especially in those less than 40 years old. In the AD group, keloid risk was significantly associated with the presence of comorbidities and concurrent allergic diseases and keloid risk increased with the severity of AD.

Previous studies of the association between AD and keloids have obtained inconsistent results. For example, a case–control study of 131 patients with keloids and 258 controls recruited during 2000–2012 revealed no association between AD and keloids in the adjusted model (adjusted OR 1.27; 95% CI 0.76 to 2.13). However, the questionnaire-based reporting method used in that study may have caused information bias resulting in under-diagnosis of AD. Another limitation is that the authors did not report AD-compatible itchy flexural rash with AD, which may have also contributed to underdiagnosis of AD. Underdiagnosis of keloids is also common because patients with keloids are often immigrants, who may be unaccustomed to seeking prompt medical attention for non-critical medical problems. Additionally, the diverse ancestry of the population analysed in Hajdarbegovic et al. may have had a confounding effect on the association between AD and keloids. A final limitation of Hajdarbegovic et al is its small sample size.
| Variables            | Patients with atopic dermatitis | Patients without atopic dermatitis | Compared with non-atopic dermatitis |
|----------------------|--------------------------------|-----------------------------------|-----------------------------------|
|                      | Keloid | PYs   | Rate | Keloid | PYs   | Rate | Crude HR (95% CI) | Adjusted HR* (95% CI) | Adjusted HR† (95% CI) |
| Overall              | 163    | 50990.85 | 3.19 | 532    | 499087.75 | 1.07 | 3.84 (3.17 to 4.66)‡ | 3.80 (3.14 to 4.61)‡ | 3.19 (2.60 to 3.89)‡ |
| Gender               |        |        |      |        |        |      |                  |                      |                      |
| Men                  | 47     | 17992.05 | 2.61 | 123    | 176032.98 | 0.69 | 4.81 (3.31 to 6.99)‡ | 4.72 (3.24 to 6.85)‡ | 3.67 (2.47 to 5.44)‡ |
| Women                | 116    | 32998.80 | 3.52 | 409    | 323054.76 | 1.27 | 3.56 (2.84 to 4.45)‡ | 3.53 (2.82 to 4.42)‡ | 3.03 (2.39 to 3.83)‡ |
| Stratify by age (years) |        |        |      |        |        |      |                  |                      |                      |
| <40                  | 122    | 26311.81 | 4.64 | 75     | 63234.61 | 1.19 | 4.07 (3.27 to 5.07)‡ | 4.07 (3.27 to 5.08)‡ | 3.44 (2.74 to 4.31)‡ |
| 40–59                | 30     | 15707.99 | 1.91 | 29     | 38829.76 | 0.75 | 3.31 (2.19 to 4.99)‡ | 3.32 (2.20 to 4.99)‡ | 2.74 (1.79 to 4.17)‡ |
| ≥60                  | 11     | 8971.06  | 1.23 | 11     | 22791.71 | 0.48 | 2.87 (1.49-5.56)§ | 2.82 (1.46-5.44)§ | 2.36 (1.22-4.59)¶ |
| Concurrent allergic disease** |       |        |      |        |        |      |                  |                      |                      |
| No                   | 22     | 9661.48  | 2.28 | 139    | 194784.11 | 0.71 | 5.41 (4.02 to 7.27)‡ | 4.03 (2.55 to 6.37)‡ | 4.11 (3.27 to 5.94)‡ |
| Yes                  | 141    | 41329.37 | 3.41 | 393    | 304303.64 | 1.29 | 3.19 (2.52 to 4.05)‡ | 3.34 (2.72 to 4.12)‡ | 2.62 (2.05 to 3.34)‡ |
| Comorbidity††        |        |        |      |        |        |      |                  |                      |                      |
| No                   | 59     | 14141.34 | 4.17 | 247    | 246626.44 | 1.00 | 4.13 (2.62 to 6.52)‡ | 4.85 (3.61 to 6.52)‡ | 3.78 (2.38 to 5.98)‡ |
| Yes                  | 104    | 36849.51 | 2.82 | 285    | 252461.31 | 1.12 | 3.37 (2.74 to 4.15)‡ | 2.82 (2.22 to 3.58)‡ | 3.08 (2.48 to 3.82)‡ |

*Model adjusted for age and gender.
†Model adjusted for age, gender, urbanisation, concurrent allergic diseases (asthma, allergic rhinitis and allergic conjunctivitis) and relevant comorbidities (hypertension, diabetes mellitus, hyperlipidaemia, chronic kidney diseases, chronic liver disease, chronic obstructive pulmonary disease, stroke, coronary artery disease, mental disorders and alcohol attributed disease).
‡P<0.001.
§P=0.002.
¶P=0.011.
**Patients with any concurrent allergic disease, including asthma, allergic rhinitis and allergic conjunctivitis, were classified as the allergic disease group.
††Patients with any examined comorbidities, including hypertension, diabetes mellitus, hyperlipidaemia, chronic kidney diseases, chronic liver disease, chronic obstructive pulmonary disease, stroke, coronary artery disease, mental disorders and alcohol attributed disease, were classified as the comorbidity group.
PYs, person-years; rate, incidence rate in per 1000 person-years.
Although the exact mechanisms underlying the relationship between AD and keloids are unclear, several possible links have been hypothesised. First, thymic stromal lymphopoietin (TSLP), an interleukin (IL)-7-like cytokine, is believed to cause Th2-type inflammatory responses.22 23 Upregulation of Th2 cytokines may be linked to atopic diseases.24 The TSLP is also reportedly linked to AD. In 2010, Lee et al collected sera from 232 children, including 75 with AD. Comparisons showed higher serum TSLP in children with AD compared with normal controls. Studies of patients with AD also indicate that skin lesions are associated with increased TSLP gene expression.25 In another recent study, Shin et al2 showed that TSLP expression was higher in keloidal tissue compared with normal tissue. Their data showed that by inducing production of collagen and tissue growth factor beta (TGF-β), TSLP contributes to excessive deposition of ECM. Since TSLP expression is apparently associated with keloids in patients with AD, keloids and AD may be induced through similar inflammatory pathways.

Periostin, which is in the fasciclin family of proteins, has important functions in the ECM and is also a matricellular protein associated with allergic inflammation.26 Kou et al reported that periostin expression in serum or in inflamed sites of patients with AD was associated with the clinical severity of AD, which suggests that periostin mediates AD.27 28 Furthermore, periostin can activate TGF-β and NF-κB to induce IL-6 in eosinophils and lead keratinocytes to produce TSLP. Periostin in fibroblasts can then be induced by activated eosinophils through TGF-β reciprocally. Besides, Zhou et al29 reported that increased expression of periostin was found in keloid scars. Since periostin participates in the epithelial–mesenchymal

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**Table 3** Incidence rates and HRs of keloid risk in patients with different severity of atopic dermatitis

| Variables              | N     | keloid | Rate  | Crude HR (95% CI) | Crude HR (95% CI) | Adjusted HR* (95% CI) | Adjusted HR* (95% CI) |
|------------------------|-------|--------|-------|-------------------|-------------------|-----------------------|-----------------------|
| Without atopic dermatitis | 33484 | 532    | 1.07  | 1.00 (reference)  | 1.00 (reference)  | 1.00 (reference)       | 1.00 (reference)       |
| With atopic dermatitis  | 8371  | 163    | 3.19  | 3.84 (3.17 to 4.66)† | 3.19 (2.60 to 3.89)† | 3.19 (2.60 to 3.89)†  | 3.19 (2.60 to 3.89)†  |
| Mild                   | 5193  | 81     | 2.58  | 3.10 (2.42 to 3.97)† | 1.00 (reference)  | 2.54 (1.98 to 3.27)†  | 1.00 (reference)       |
| Moderate to severe     | 3178  | 82     | 4.19  | 5.02 (3.93 to 6.42)† | 1.62 (1.19-2.21)‡ | 4.39 (3.38 to 5.70)†  | 1.73 (1.26 to 2.36)†  |

*Model adjusted for age, gender, urbanisation, concurrent allergic diseases (asthma, allergic rhinitis and allergic conjunctivitis) and relevant comorbidities (hypertension, diabetes mellitus, hyperlipidaemia, chronic kidney diseases, chronic liver disease, chronic obstructive pulmonary disease, stroke, coronary artery disease, mental disorders and alcohol attributed disease).

†P<0.001.
‡P=0.002.

IRR, incidence rate ratio; PYs, person-years; rate, incidence rate in per 1000 person-years.

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**Figure 2** Cumulative incidence of keloid among patients with atopic dermatitis and the control cohort. The incidence of keloid in atopic dermatitis patients and controls was compared using the log-rank test by a Kaplan-Meier analysis.
transition, periostin is important in forming both AD and keloids.

Sirtuins, which are class III histone deacetylases, are implicated in fibrotic disease, inflammation and cellular proliferation, apoptosis and differentiation. Ming et al demonstrated that loss of sirtuin can reduce the epidermal formation of filaggrin, resulting in disruption of the epidermal barrier and a predisposition to occurrence of AD. 31 32 Bai et al showed that expression of sirtuin was suppressed in fibrotic scarring and that sirtuin deletion enhanced fibrotic wound healing. Sirtuin overexpression can attenuate the expression of ECM induced by TGF-β. Loss of sirtuin leads to both AD development and excessive fibrotic scarring. Finally, we hypothesised that individuals with AD may be predisposed to subsequent development of keloids.

TGF-β is a main fibrotic growth factor involved in wound healing and tissue repair. Uregulation of TGF-β gene with proinflammation cytokines was found in keloids, resulting in keloidal fibroblasts formation together with collagen synthesis. 37 Previously mentioned literatures showed periostin, TSLP and loss of sirtuins can activate TGF-β in AD and keloids. Therefore, patients with AD are prone to develop keloids.

Psychological stress can trigger various physiological responses in the endocrine, nervous and immune systems. Psychoemotional stress is associated with a higher than normal incidence of several skin diseases including AD and it is also known to aggravate AD. Anxiety and depression have been reported to correlate with AD. 34–36 In Chen et al, the patients with AD had a higher risk of developing major depression, any depressive disorder and anxiety disorders in later life. Other studies have reported an association between anxiety and keloids. In Furtado et al, for example, psychological stress was evaluated in 25 patients with keloids before undergoing surgical resection and postoperative radiotherapy. They found that stress had a direct correlation with development of keloids. Notably, the hypothalamic–pituitary–adrenal axis has a well-known mediating role in the eventual contributing effects of psychological stress on development of both AD and keloids. 38–45 Psychological stress also increases concentrations of nerve growth factor (NGF). Ewa and colleagues demonstrated that comparing to healthy ones, patients with AD had significant increased plasma levels of substance Pand NGF. The NGF is made by keratinocytes and contributes to neurogenic inflammation. Akashi et al found that neurogenic inflammation has an important role in keloid formation. Therefore, we suggest that AD should be considered a risk factor for keloids.

A notable strength of this study is the use of a large representative population-based dataset, which enabled analysis of all cases of AD and keloids during the study period. Moreover, the large sample size also provided sufficient statistical power to compare the complex relationships between these two diseases in the AD and non-AD groups. Notably, since most of the Taiwan population is ethnic Chinese, analysis of this population may yield valuable insights into the roles of genetic ancestry in AD and keloids. However, several limitations of this study are noted. First, the data analysed in this study were originally collected for insurance purposes rather than for research purposes. Second, data for diagnoses of AD and keloids in the study population were based on ICD-9-CM codes entered by different doctors, which may have introduced coding errors. The Taiwan Bureau of National Health Insurance has attempted to minimise this problem by performing regular audits and by imposing large fines on hospitals that make coding errors. Nevertheless, the NHIRD has been rigorously examined and extensively used in many studies in recent years. Third, despite the relatively low healthcare costs in Taiwan, some patients attempt to eliminate AD and keloids through folk remedies or through private treatment by local pharmacists rather than by medical professionals. Moreover, the NHIRD does not record all information regarding risk factors such as body mass index, blood groups, smoking, alcohol consumption and operation sites, which potentially compromises our results. Lacking information about trauma history, anatomical sites, keloid size and number, acne and the different methods of surgery is also a flaw in the report. Finally, the retrospective design of this study only revealed statistically significant relationships. Further studies are needed to validate actual causal relationships. Further clinical trials are needed to clarify the underlying mechanisms of this association.

In conclusion, this study showed that AD is associated with development of keloids and suggested that AD could be an important risk factor in the future development of keloids. Nevertheless, further studies are needed to clarify this association and to explore pathophysiological mechanisms that are common to AD and keloids.

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