Allergic rhinitis and dental-supporting tissue diseases in children

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Abstract
The etiology of dental-supporting tissue diseases in children is multifactorial and not merely related to oral hygiene. Therefore, in the present study, we investigated the relationship between children <18 years old with allergic rhinitis (AR) and the risk of dental-supporting tissue diseases.

Data from the National Health Insurance Research Database (NHIRD) of Taiwan were used to conduct a retrospective longitudinal cohort study. The study cohort comprised 378,160 patients with AR (AR group) and 378,160 patients without AR (non-AR group), who were selected through frequency matching based on age, sex, and the index year. The study patients were followed until dental-supporting tissue diseases occurrence, withdrawal from the National Health Insurance program, or December 31, 2013. Cox proportional hazards regression analysis was conducted to calculate the risk of dental-supporting tissue diseases in the AR group after adjustment for age, sex, and relative comorbidities.

The adjusted HRs of periodontal, pulp, and periapical diseases in AR children were higher than those in the non-AR controls (1.51, 95% CI: 1.50 to 1.53; 1.06, 95% CI: 1.05 to 1.07, respectively). The AR to non-AR HRs of these inflammatory dental diseases were particularly higher in children <6 years old and in boys. The HRs of periodontal, pulp, and periapical diseases were greatest in those with >5 AR-related medical visits/year (5.57, 95% CI: 5.50 to 5.66; 4.06, 95% CI: 4.00 to 4.12, respectively).

Children with AR had a greater risk of inflammatory dental-supporting tissue diseases, particularly those <6 years old with primary teeth, boys, and those with severe persistent AR.

Abbreviations: AR = allergic rhinitis, CI = confidence interval, HR = hazard ratio, ICD-9-CM = International Classification of Diseases, 9th Revision, Clinical Modification, IL = Interleukin, NHI = National Health Insurance, NHIRD = National Health Insurance Research Database, SD = standard deviation.

Keywords: allergic rhinitis, children, cohort study, periodontal disease, pulpal and periapical disease

Bullet points
- The present study is the first to utilize a large sample to investigate the association between childhood AR and inflammatory dental-supporting tissue diseases, including periodontal, pulp, and periapical diseases.
- Children with AR had a greater risk of inflammatory dental-supporting tissue diseases, particularly those <6 years old with primary teeth, boys, and those with severe persistent AR.
- Identifying risk factors for inflammatory dental-supporting tissue diseases is important not only to prevent
inflammation in the oral cavity but also to prevent multiple inflammation-related systemic diseases.

1. Introduction

Periodontal, pulp, and periapical diseases, inflammatory dental-supporting tissue diseases, are common dental disorders in children. The periodontal, pulp, and periapical tissues are closely associated with development, anatomy, and function. Pulp and periapical disease may affect the initiation and progression of periodontal disease. In contrast, patients with chronic periodontitis usually have pulp tissue changes such as inflammation, edema, necrosis, fibrosis, and calcification. The exchange of immune cells and inflammatory mediators between periodontal, pulp, and periapical tissues is considered to happen through the vascular system in the apical foramen. Besides, a variety of systemic diseases are reported to interfere with periodontal, pulp, and periapical health.

Chronic periodontitis and pulpitis may increase systemic inflammation, and are linked to multiple systemic diseases such as type 2 diabetes, metabolic disease, obesity, cardiovascular disease, fatty liver disease, and cancer. Therefore, identifying risk factors for periodontal, pulp, and periapical diseases in the pediatric population is important not only to prevent inflammation in the oral cavity but also to prevent multiple inflammation-related systemic diseases. However, despite increasing research on the link between dental-supporting tissue diseases and systemic diseases over the past few decades, the fundamental biological mechanisms of the association are not fully elucidated.

A few studies show the relationship between periodontal disease or salivary micro-flora change and allergic disease. Allergic rhinitis (AR) is the most common allergic disease. Children with AR are frequently observed to have gingival and periapical inflammation. Interleukin (IL)-5, 12, 13, and IL-16, involved in the regulation of the onset and development of AR, also play a role in the pathogenesis of periodontal diseases. Several studies have discussed the relationship between AR and dental caries in children. Although some studies with inconsistent results have focused on adult patients, there is no study demonstrating the relationship between AR and periodontal, pulp, and periapical diseases in childhood.

Because there are 3 stages, including primary, transition, and permanent, of teeth development in children, whether AR is an independent risk factor for dental-supporting tissue diseases in childhood is of great interest. This large, population-based study aimed to explore how AR correlates with childhood inflammatory dental-supporting tissue disease, including periodontal, pulp, and periapical diseases.

2. Material and methods

2.1. Data source

The National Health Insurance (NHI) program was established in Taiwan in 1995. This program has covered the reimbursement of medical care for approximately 23 million people, accounting for over 99% of the population of Taiwan. The NHIRD contains all the registry data of the insured, diagnostic codes of International Classification of Diseases, 9th Revision, Clinical Modification (ICD-9-CM), as well as details of outpatient and inpatient visits, procedures, prescriptions, and medical expenditure. This study used a dataset from the NHIRD, containing a randomly selected sample of half of all insured children in Taiwan. Based on the Personal Information Protection Act, de-identification was performed before the release of the dataset to researchers; thus, informed consent was not required for this study. This study received approval from the institutional review board of China Medical University Hospital (CRREC-103-048).

2.2. Study design and subjects

This was a retrospective cohort study using the NHIRD. Between 2000 and 2012, patients <18 years old with newly diagnosed AR (ICD-9-CM code 477) were selected as the AR cohort. The comparison non-AR cohort with no AR diagnostic codes was selected by 1:1 matching based on a propensity score. The propensity score was calculated using a logistic regression model to estimate the probability of disease assignment based on baseline variables, which included age, sex, index year, urbanization, and the presence of the following comorbidities: chronic sinusitis (ICD-9-CM 473), obstructive sleep apnea (ICD-9-CM 372.23), asthma (ICD-9-CM 493), hypertrophy of the tonsils and adenoids (ICD-9-CM 474.1), and obesity (ICD-9-CM 278). Confounders such as age, sex, urbanization, and comorbidities were adjusted for in the analysis.

2.3. Confounders

Periodontal, pulp, and periapical diseases are multi factorial diseases. Well known risk factors for oral diseases include poor oral hygiene, frequent consumption of a sugary diet, female, and urbanization.

Obesity has been reported to be associated with periodontal disease and may be related to the frequent consumption of a sugary diet. There were no data on the frequency of sugary diet consumption. Therefore, obesity was used instead of frequency of sugar diet consumption. Chronic sinusitis and asthma are common comorbidities in children with AR. The etiology and medication, such as antibiotics and inhaled steroids, might be associated with oral disease.

There is some evidence on the association between periodontal disease and obstructive sleep apnea. These comorbidities are thought to be confounding factors that increase the risk of periodontal, pulp, and periapical diseases.

Dental facial anomaly and children with disability could increase the risk of dental disease because of difficulty in maintaining dental health. Individuals with congenital anomalies (ICD-9-CM 740-759) and mental retardation (ICD-9-CM 317-319) were excluded. Individuals diagnosed with dental diseases and dentofacial anomalies before 2000 were also excluded.

2.4. Outcome measurement

The risk of periodontal, pulp, and periapical diseases was compared in the AR and non-AR cohorts between 2000 and 2013. All physician-diagnosed dental diseases were determined.
using diagnostic codes for periodontal (ICD-9-CM 523), pulp, and periapical diseases (ICD-9-CM 522).

2.5. Statistical analysis

Baseline characteristics of AR and non-AR cohorts were compared using standardized mean difference. Values of standardized mean differences \( \leq 0.01 \) indicated a negligible difference in mean values between the AR and non-AR cohorts. We used a Cox proportional hazard regression model to compare the risk of periodontal, pulp, and periapical diseases in AR and non-AR cohorts, adjusted for age, sex, urbanization, and comorbidities. We also compared the risk of development of these diseases given the average frequency of AR-related medical visits. The cumulative incidence curves of periodontal, pulp, and periapical diseases in the study cohort were estimated using Kaplan-Meier analysis. All analyses were performed using SAS software, version 9.1 (SAS Institute, Cary, NC, USA), and a \( P \) value \(< .001\) was considered statistically significant.

3. Results

A total of 378,160 AR patients were identified; 45.9\% of these patients were girls. The mean (standard deviation, SD) age at AR diagnosis was 4.85 (3.82) years, and 73.6\% were newly diagnosed before the age of 6 years (Table 1). More than half of the subjects with AR resided in urban areas (approximately 60.8\%). There were no significant differences between the AR and non-AR cohorts in terms of age, urbanization (except the highest residential area), and comorbidity of chronic sinusitis, asthma, and obesity (Table 1).

The analysis for the risk of periodontal, pulp, and periapical diseases in children without AR is shown in Table 2. Children with AR had a significantly higher risk to develop these inflammatory dental supporting tissue diseases, irrespective of differences in age, sex, or urbanization than those without AR. The adjusted hazard ratio (HR) was higher in the AR cohort than in the non-AR cohort (1.51, 95\% confidence interval: 1.50–1.53; 1.06, 95\% confidence interval [CI]: 1.05–1.07) for periodontal disease and pulp and periapical disease, respectively. The ages \(< 6, 6–11,\) and \(> 11\) years mark the 3 dentition stages primary, transitional, and permanent teeth stage, respectively. The risk of pulp and periapical diseases in the transitional teeth stage did not differ between the 2 cohorts; however, the risk for the 2 diseases was markedly higher in AR cohort than in the non-AR cohort for the other dental stages. The association between the annual frequency of medical visits due to AR and the risk of periodontal, pulp, and periapical diseases is shown in Table 3. The adjusted HR was higher for children with a high frequency of AR-related medical visits per year than for children without medical visits for AR. The dose-dependent relationship was found between a high frequency of AR-related medical visits and developing periodontal, pulp, and periapical diseases.

Regarding the effect of comorbidities, a high HR for periodontal, pulp, and periapical diseases was noted even for AR children without comorbidities of chronic sinusitis, obstructive sleep apnea, asthma, and hypertrophy of tonsils and adenoids (Table 4). Figure 1 demonstrates the Kaplan–Meier analysis for the cumulative incidence of periodontal disease, pulp, and periapical diseases for the AR cohort compared to the non-AR cohort. (log-rank test, \( P < .001\)).

4. Discussion

Chronic inflammatory diseases affecting dental tissues if not diagnosed promptly and treated appropriately may ultimately lead to tooth loss.\(^{1,2}\) Studies on the association between AR and inflammatory dental supporting tissue diseases have reported

### Table 1

Demographics between children with and without allergic rhinitis (AR).

|                      | Non-AR (\(N = 378160\)) | AR (\(N = 378160\)) | Standard difference |
|----------------------|--------------------------|---------------------|---------------------|
| Age, years, mean (SD) | 4.85 (3.92)              | 4.85 (3.82)         | 0.005               |
| Stratified age, years |                          |                     |                     |
| \(< 6\)              | 273408 (72.3)            | 278206 (73.6)       | 0.03                |
| \(6–11\)             | 75566 (20.0)             | 71291 (18.9)        | 0.03                |
| \(\geq 12\)          | 29186 (7.72)             | 28663 (7.58)        | 0.005               |
| Sex                  |                          |                     |                     |
| Girl                 | 177481 (46.9)            | 173452 (45.9)       | 0.02                |
| Boy                  | 200679 (53.1)            | 204708 (54.1)       | 0.02                |
| Urbanization\(^{*}\) |                         |                     |                     |
| 1 (highest)          | 11378 (29.5)             | 113918 (30.1)       | 0.02                |
| 2                    | 117316 (31.0)            | 110687 (30.7)       | 0.007               |
| 3                    | 73769 (19.5)             | 73364 (19.4)        | 0.003               |
| 4 (lowest)           | 75697 (20.3)             | 74791 (19.8)        | 0.006               |
| Comorbidity          |                          |                     |                     |
| Chronic sinusitis    | 1675 (0.44)              | 1661 (0.44)         | 0.001               |
| Obstructive sleep apnea | 270 (0.06)          | 1383 (0.27)         | 0.05                |
| Asthma               | 13149 (3.48)             | 13163 (3.48)        | 0.000               |
| Obesity              | 267 (0.08)               | 256 (0.07)          | 0.003               |
| Hypertrophy of tonsils and adenoids | 484 (0.11) | 1784 (0.35) | 0.049               |

\(^{*}\) SD = standard deviation.

The urbanization level was categorized by the population density of the residential area into 4 levels, with level 1 as the most urbanized and level 4 as the least urbanized. The standardized difference was used to quantify differences in means or prevalence between the patients with allergic rhinitis and without allergic rhinitis for continuous and categorical matching variables. A value of standardized mean differences equals .01 or less, which indicates a negligible difference in means between the patients with and without AR.
conflicting results. The present study is the first to utilize a large sample to investigate the association between AR and periodontal, pulp, and periapical diseases in children. We found that AR children had a significantly high risk and higher cumulative incidences of periodontal, apical, and periapical diseases than those without AR. A particularly high risk for these inflammatory dental supporting tissue diseases were found in those below age of 6 years with primary teeth and boys. Further, children with more frequent medical visits for AR had higher risks for periodontal, pulp, and periapical diseases, indicating a dose-dependent effect.

There are few studies investigating the interaction between AR and periodontal disease and the findings are contradictory. One cross-sectional study in Germany and the other one in Korea revealed an inverse association between periodontal disease and AR in adults. In contrast, another 2 matched case-control studies in Taiwan showed a positive correlation between AR and periodontal disease in adults. Moreover, Ho et al revealed that men, urban citizens, and low income people with AR had higher incidence of pulpitis. Our study found that AR children had a high risk of inflammatory dental supporting tissue diseases. Moreover, we also assessed the risks of these dental surrounding tissue diseases according to different stages of teeth development. Our results revealed that younger children with primary teeth had particularly higher risks for periodontal, pulp, and periapical diseases. Because chronic periodontitis and pulpitis may increase

| Table 2 |
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| The risk of periodontal disease, and pulp and periapical disease compared to children without allergic rhinitis (AR) stratified by demographics in Cox proportional hazard regression. |

| Periodontal disease | Non-AR Event Person-years IR | AR Event Person-years IR | Adjusted HR (95% CI) |
| --- | --- | --- | --- |
| All | 56737 2717378 20.9 | 85054 2584093 32.9 | 1.51 (1.50, 1.53)† |
| Sex | 28977 1215356 23.8 | 40614 1137300 35.7 | 1.48 (1.46, 1.50)† |
| Boy | 27759 1502022 18.5 | 44440 1446793 30.7 | 1.55 (1.53, 1.57)† |
| Dentition stage† | 35604 2097124 17.0 | 55983 2032873 27.5 | 1.58 (1.56, 1.60)† |
| Primary | 13527 532125 25.4 | 19008 468329 40.6 | 1.48 (1.45, 1.51)† |
| Transitional | 7605 88129 86.3 | 10083 82891 121.4 | 1.38 (1.34, 1.42)† |
| Pulp and periapical disease | 60987 2594599 23.5 | 66223 2579405 25.7 | 1.06 (1.05, 1.07)† |
| Sex | 27810 1172544 23.7 | 28990 28990 25.2 | 1.03 (1.02, 1.05)† |
| Boy | 33177 1422055 23.3 | 37223 37233 26.1 | 1.08 (1.07, 1.10)† |
| Dentition stage# | 56421 1931383 29.2 | 61483 1952367 31.5 | 1.06 (1.05, 1.07)† |
| Primary | 3143 567084 5.54 | 3302 531293 6.22 | 1.01 (0.97, 1.06)† |
| Transitional | 1423 96133 14.8 | 1438 95745 15.0 | 0.98 (0.91, 1.05)† |
| Permanent | 1423 96133 14.8 | 1438 95745 15.0 | 0.98 (0.91, 1.05)† |

† P < .001.  
† Dentition stage is stratified by age (<6 years: primary dentition stage; 6-11 years: transitional (mixed) dentition stage; ≥12 years: permanent dentition stage).  
IR = incidence rate, per 1000 person-years; HR = hazard ratio; CI = confidence interval.  
Adjusted HR†, adjusted for age, sex and comorbidity of chronic sinusitis, obstructive sleep apnea, asthma, and hypertrophy of tonsils and adenoids.

| Table 3 |
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| The risk of periodontal disease, and pulp and periapical disease among average frequency for medical visits of allergic rhinitis in Cox proportional hazard regression. |

| Average frequency for medical visit, per years | Event Person-years IR Adjusted HR (95% CI) |
| --- | --- | --- |
| Periodontal disease | Non-AR | 56736 2717378 20.9 | 1.00 (Reference) |
| <3 | 42612 2094560 20.3 | 0.97 (0.95, 0.99)† |
| 4-5 | 9875 180652 54.7 | 2.78 (2.72, 2.84)† |
| >5 | 32567 308881 105.4 | 5.57 (5.50, 5.65)† |
| P for trend | <.001 |
| Pulp and periapical disease | None | 60987 2594599 23.5 | 1.00 (Reference) |
| <3 | 27587 2094560 20.3 | 1.57 (1.56, 1.59)† |
| 4-5 | 9875 180652 54.7 | 1.76 (1.72, 1.80)† |
| >5 | 30364 288618 105.4 | 4.06 (4.00, 4.12)† |
| P for trend | <.001 |

† P < .001.  
† Adjusted for age, sex and comorbidity of chronic sinusitis, obstructive sleep apnea, asthma, and hypertrophy of tonsils and adenoids.  
IR = incidence rate, per 1000 person-years; HR = hazard ratio; CI = confidence interval.  
Adjusted HR, adjusted for age, sex and comorbidity of chronic sinusitis, obstructive sleep apnea, asthma, and hypertrophy of tonsils and adenoids.
systemic inflammation and many systemic diseases, such as type 2 diabetes, metabolic disease, obesity, cardiovascular disease, fatty liver disease, and cancer, aggressive treatment of childhood AR may be beneficial for oral health improvement and for prevention of certain systemic diseases.\[4,5\]

Inhalation of corticosteroids has been reported to increase the incidence of caries and periodontal disease because of a resultant change in oral pH, local deposition of steroids in the oral cavity, and their effect on oral mucosa\[4,23\]. However, there are limited studies on the association between intranasal steroids and dental inflammatory disease. The mainstay treatment of AR includes oral or intranasal antihistamines and intranasal corticosteroids.\[25\] The management of AR depends on its severity and duration. Those with persistent and severe AR are at a higher risk

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### Table 4

|                                | Non-AR Event | Person-years | IR  | AR Event | Person-years | IR  | Adjusted HR† (95% CI) |
|--------------------------------|--------------|--------------|-----|----------|--------------|-----|-----------------------|
| **Periodontal disease**        |              |              |     |          |              |     |                       |
| Chronic sinusitis              |              |              |     |          |              |     |                       |
| No                             | 66145        | 3228924      | 20.5| 105447   | 31.4         | 31.4| 1.51 (1.50, 1.53)**   |
| Yes                            | 302          | 12253        | 24.7| 3477     | 9115         | 35.1| 1.60 (1.43, 1.81)**   |
| Obstructive sleep apnea        |              |              |     |          |              |     |                       |
| No                             | 66397        | 3239850      | 20.5| 108641   | 3446643      | 31.5| 1.51 (1.50, 1.53)**   |
| Yes                            | 50           | 1327         | 37.7| 283      | 6525         | 43.4| 1.12 (0.82, 1.51)     |
| Asthma                         |              |              |     |          |              |     |                       |
| No                             | 64970        | 3150083      | 20.6| 87352    | 266063       | 37.8| 1.51 (1.49, 1.59)**   |
| Yes                            | 1477         | 91094        | 16.2| 21572    | 787105       | 27.4| 1.68 (1.60, 1.77)**   |
| Hypertrophy of tonsils and adenoids |              |              |     |          |              |     |                       |
| No                             | 66383        | 3237761      | 20.5| 108559   | 3442232      | 31.5| 1.51 (1.50, 1.59)**   |
| Yes                            | 64           | 3417         | 18.7| 365      | 10936        | 33.4| 1.64 (1.25, 2.15)**   |
| **Pulp and periapical disease** |              |              |     |          |              |     |                       |
| Chronic sinusitis              |              |              |     |          |              |     |                       |
| No                             | 72843        | 3070794      | 23.7| 84812    | 3335548      | 25.4| 1.06 (1.05, 1.07)**   |
| Yes                            | 244          | 12038        | 20.3| 2286     | 101373       | 22.6| 1.01 (0.88, 1.15)     |
| Obstructive sleep apnea        |              |              |     |          |              |     |                       |
| No                             | 73058        | 3081474      | 23.7| 86950    | 3430035      | 25.4| 1.06 (1.05, 1.07)**   |
| Yes                            | 29           | 1358         | 21.4| 148      | 6886         | 21.5| 1.17 (0.78, 1.76)     |
| Asthma                         |              |              |     |          |              |     |                       |
| No                             | 71034        | 2997545      | 23.7| 66529    | 2670281      | 24.9| 1.06 (1.05, 1.07)**   |
| Yes                            | 2053         | 85287        | 24.1| 20569    | 766640       | 26.8| 1.09 (1.04, 1.14)     |
| Hypertrophy of tonsils and adenoids |              |              |     |          |              |     |                       |
| No                             | 73012        | 3079669      | 23.7| 86853    | 3425832      | 25.4| 1.06 (1.05, 1.07)**   |
| Yes                            | 75           | 3163         | 23.7| 245      | 11089        | 22.1| 1.07 (0.82, 1.40)     |

*P < .01.
**P < .001.

IR = incidence rate, per 1000 person-years, HR = hazard ratio, CI = confidence interval.
of using intranasal steroids. We speculate that nasopharyngeal deposition of steroids due to postnasal drip might induce the same oral diseases that inhaled steroids do, which explains our finding that AR children with more frequent medical visits have a higher risk of periodontal, pulp, and periapical diseases. Chronic polymicrobial infection to the surrounding dental tissues and eliciting a host inflammatory immune response in susceptible individuals is a central feature of periodontal, pulp, and periapical diseases. Our study showed that AR children had a higher incidence and risk of dental soft tissue inflammation. Although the pathogenesis remains unclear, dry mouth due to mouth breathing and taking an oral antihistamine and inflammatory reactions due to AR may worsen dental health. Mast cells play a crucial role in the pathogenesis of AR. Several studies demonstrate that mast cells may also play an important role in the development of pulpsitis and periodontitis during its acute stages and its subsequent transition to chronic inflammation. Histamine, released from mast cells, acts as a strong vasodilator and mediator of vascular permeability and may play a role in initiating pulp, periapical, and periodontal inflammation. There are several limitations of our study. First, the claims data do not include information on oral hygiene and the severity of periodontal, pulp, and periapical diseases. The urbanization used instead of oral hygiene due to association with oral hygiene behavior. Nonetheless, we report a positive and dose-dependent relationship between AR and development of periodontal, pulp, and periapical diseases. Second, the AR cohort was selected based on the diagnostic code of ICD-9-CM. Ideally, a skin prick test is necessary to confirm the AR diagnosis; however, the diagnoses for our study cohort were made by a licensed and well-trained physician. Third, surveillance bias should be taken into consideration for the AR cohort with a high risk of dental surrounding tissue diseases. Those with severe and persistent AR need more visits for medical care, which can trigger early screening for dental diseases. However, AR children with more medical care visits may have a higher chance to have oral health knowledge and surveillance, that may controvert this bias.

5. Conclusion
AR children had a higher risk for inflammation of dental supporting tissues, including periodontal, pulp, and periapical diseases. The risk was much higher in children aged less than 6 years with primary teeth, in boys, and in those with severe persistent AR. These findings highlight the importance of awareness and regular follow-up of dental conditions in AR children.

Author contributions
*Wan-Yu Lai and Chang-Ching Wei contributed equally. Wan-Yu Lai, Chen-Hao Mai, and Chang-Ching Wei conceptualized and designed the study. Wan-Yu Lai and Chang-Ching Wei drafted the initial manuscript. Cheng-Li Lin carried out the acquisition of data and analysis and interpretation of data. Lei Wan and critically reviewed and revised the manuscript. Chang-Ching Wei and Jeng-Dau Tsai coordinated and supervised data collection, critically reviewed the manuscript, and approved the final manuscript as submitted.

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