Insights into pediatric pollen food allergy syndrome

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Pollen food allergy syndrome (PFAS), previously known as oral allergy syndrome (OAS), is an immunoglobulin E–mediated food allergy caused by cross-reactivity between pollen and plant allergens. Patients with pollen allergies may react to fruits, vegetables, and/or tree nuts that share epitopes of prior sensitized allergenic pollen.1 OAS is a term based on a characteristic clinical manifestation of oropharyngeal symptoms. OAS is primarily caused by plant foods which cross-react with pollen, but it can be caused by animal food allergens. Thus, OAS is currently considered a collection of symptoms of food allergy or latex fruit syndrome in addition to PFAS.

The prevalence of PFAS is difficult to estimate for several reasons including a lack of awareness of patients and physicians. Since the prevalence of pollinosis has increased with climate change,2 the incidence of PFAS is also expected to increase.

Little is known about PFAS prevalence by age. Among British children with seasonal allergic rhinitis (AR), the prevalence of PFAS was reportedly 17% and 78% in those under 5 and older than 10 years, respectively.3 This large difference in prevalence is probably due to eating more fruits, vegetables, and/or nuts, and being more aware of symptoms after age 5. Recent study showed no age-based difference in PFAS prevalence among patients with pollen allergies except those under 5 or over 60 years of age.4

The incidence of PFAS in children from some countries was reported as 9%–43%, depending on the geographical distribution of types of pollen and the characteristics of study subjects (Table 1).3,8

The prevalence of AR in Korean children is expected to be 21%; of them, 37.0% showed pollen sensitization.5 The PFAS prevalence is 42.7% among Korean children with pollinosis.4 This incidence in hospital-based patients may be higher than those of general population. With those data, the estimated prevalence of PFAS would be 3.3%. Well-known causes of PFAS are birch, alder, timothy grass, orchard grass, mugwort, and ragweed pollen.

Class 1 food allergens, which induce sensitization via the gastrointestinal tract, are stable to heat, acid, and digestive enzymes, and are usually responsible for systemic reactions. Class 2 food allergens are highly cross-reactive with pollen, sensitized through the respiratory route, and generally unstable and degraded easily by heat and digestive enzymes. Symptoms of PFAS include pruritus, paresthesia, and angioedema of the lips, oral mucosa, tongue, and palate. However, other cutaneous, respiratory, gastrointestinal, neurologic, and cardiovascular symptoms include pruritus, paresthesia, and angioedema of the lips, oral mucosa, tongue, and palate. However, other cutaneous, respiratory, gastrointestinal, neurologic, and cardiovascular symptoms.5

Clinical manifestations can vary depending on sensitized pollen (panallergen) type and implicated food allergens from mild oral symptoms to systemic reactions including anaphylaxis.

Table 1. Prevalence of pediatric pollen food allergy syndrome

| Country | Subjects, numbers | Age (yr) | Prevalence (%) | frequently Implicated fruit | Implicated pollen | Comments |
|---------|-------------------|---------|----------------|---------------------------|------------------|----------|
| Italy6) | Seasonal AR, 1360 | 4–18    | 24             | Kiwi                      | Timothy grass    |          |
| Australia5) | AR with pollen sensitization, 66 | 4–17 | 12 | Watermelon | Perennial rye grass | Sydney region |
| Britain3) | Seasonal AR, 54 | <5      | 17             | Hazelnut, apple, kiwi     | Birch            |          |
|          |                   | 5–10    | 50             |                           |                  |          |
|          |                   | >10     | 78             |                           |                  |          |
| Mexico7) | Seasonal AR, 267 | 6–14    | 8.9            | Pine apple                | Oak              |          |
| Korea8)  | Pollen allergy, 648 | ≤18 | 42.7          | Peach, apple              | Birch, alder, oak |          |
| Korea8)  | AD with birch sensitization, 186 | 2–6 | 36.6          | Kiwi, apple               | Birch            |          |
|          |                   | 7–18    | 50.5           | Apple, kiwi, peach        |                  |          |

AR, allergic rhinitis; AD, atopic dermatitis

1) OAS is a term based on a characteristic clinical manifestation of oropharyngeal symptoms. OAS is primarily caused by plant foods which cross-react with pollen, but it can be caused by animal food allergens. Thus, OAS is currently considered a collection of symptoms of food allergy or latex fruit syndrome in addition to PFAS.

2) The prevalence of PFAS is difficult to estimate for several reasons including a lack of awareness of patients and physicians. Since the prevalence of pollinosis has increased with climate change, the incidence of PFAS is also expected to increase.

3) Little is known about PFAS prevalence by age. Among British children with seasonal allergic rhinitis (AR), the prevalence of PFAS was reportedly 17% and 78% in those under 5 and older than 10 years, respectively. This large difference in prevalence is probably due to eating more fruits, vegetables, and/or nuts, and being more aware of symptoms after age 5. Recent study showed no age-based difference in PFAS prevalence among patients with pollen allergies except those under 5 or over 60 years of age.

4) The incidence of PFAS in children from some countries was reported as 9%–43%, depending on the geographical distribution of types of pollen and the characteristics of study subjects. The prevalence of AR in Korean children is expected to be 21%; of them, 37.0% showed pollen sensitization. The PFAS prevalence is 42.7% among Korean children with pollinosis. This incidence in hospital-based patients may be higher than those of general population. With those data, the estimated prevalence of PFAS would be 3.3%. Well-known causes of PFAS are birch, alder, timothy grass, orchard grass, mugwort, and ragweed pollen.

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Patients with PFAS reacting to the same food may have different sensitization profiles that result in different responses to food. Bet v 1 homologues belong to PR-10, well-known causes of OASs, are heat-unstable, thus, elicit mild oropharyngeal symptoms, and account for most of the clinical cross-reactivity between birch pollen and Rosaceae fruits (e.g., apple, pear, plum), Apiaceae vegetables (e.g., celery, carrot), Betulaceae (hazelnut), and Fabaceae (soybean). However, some molecules such as Api g 1 (celery), Ara h 8 (peanut) and Gly m 4 (soybean) can be stable, and cause systemic symptoms. Apple, peach and other fruits, as well as peanuts are commonly implicated foods in Bet v 1 sensitized subjects.

Profilins also are ubiquitous, heat-labile, and easily degraded molecules by digestive enzymes, and known to be responsible allergens for the fruit-pollen, fruit-fruit, and latex-food syndrome. Musk melon (Cuc m 2) has been suggested as a clinical marker of sensitization to profilin.

Lipid transfer protein (LTP) are members of the family of PR-14, resistant to pepsin digestion, and can cause both class I and 2 food allergies. LTP is widely present in a variety of plant foods and pollens, and known to be responsible for allergic reactions to various foods including apple, peach, asparagus, barley, parsley and carrot. Several LTPs are highly cross-reactive among them, thus, LTP-sensitized patients were reported to develop allergic reactions to relevant multiple foods, which is called LTP syndrome. Since Pru p 3 in peach shares epitopes with LTPs from many fruits, nuts and vegetables, and even with mugwort (Art v 3), Pru p 3 was suggested to be a primary sensitizer. However, sensitization to LTP without sensitization to Pru p 3 was reported in patients with fruit allergies; kiwi (Act d 10), or pomegranate (Pun g 1).

In cases with plant food allergy, such as peach allergy, various reactions can be expected. Some of the subjects with PFAS are associated with birch pollen Bet v 1 (Pru p 1 in PR-10), or birch Bet v 2, mugwort and/or grass pollen (Pru p 4 in profilin), both of them generally have mild reaction. However, others are caused by LTP (Pru p 3); some of them have multiple allergies to various pollen and plant foods, others to peach only. They will have more severe systemic reactions, including anaphylaxis, especially when the cofactor is present. Therefore, patients with plant food allergy, class 1, could be diagnosed with OAS, which may lead to suboptimal treatment of potentially life-threatening food allergies.

In conclusion, although the true prevalence of PFAS is difficult to determine in the general pediatric population, PFAS can be present in a significant proportion of children, especially in those with pollen-induced AR. Therefore, it is important to take a detailed medical history in order to diagnose PFAS. Children who are unwilling to eat fruits, vegetables, and/or nuts or having food-associated oropharyngeal symptoms should be evaluated for PFAS. Although many aspects of PFAS remain poorly understood because of great heterogeneity in its causes, risk factors, natural history, comorbidities, and treatment responses, physicians are expected to properly categorize patients and provide appropriate guidance upon understanding the prevalent allergenic pollen and cross-reactive foods by geographical region.

Conflicts of interest
No potential conflict of interest relevant to this article was reported.

See the article “Pollen-food allergy syndrome in children” via https://doi.org/10.3345/cep.2019.00780.

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