Pediatric Idiopathic Anaphylaxis: 3 Case Reports and A Systematic Review of Literature

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Abstract

Background: Idiopathic Anaphylaxis (IA) is a diagnosis of exclusion and represents a major diagnostic and management challenge, especially in children. There are no current guidelines for diagnosis and management of pediatric IA. We aim to present a series of 3 cases of pediatric IA followed by a systematic review of the literature.

Methods: We describe 3 cases of pediatric IA and conducted a systematic review of original articles published in the past 22 years regarding diagnosis and management of pediatric IA.

Results: The current proposed diagnostic approach and treatment regimens are based on a few small studies. Future large-scale studies are required. IA is a diagnosis of exclusion and should be made only after extensive evaluation excluding potential triggers of anaphylaxis as well as non-allergic conditions with a similar presentation. The current proposed treatment regimens recommend prophylactic treatment with continuous prednisone for patients with frequent episodes. However, daily treatment with systemic steroids have well recognised serious adverse effects such as obesity, poor growth, osteoporosis and behavioural disturbances that are of paramount importance in the pediatric age group. More recently the use of biologics was suggested to benefit patients with IA although the optimal management protocol is not yet established.

Conclusion: Future studies are needed to optimize diagnosis and treatment strategies in pediatric cases of IA. Omalizumab may be a promising novel therapeutic option for pediatric IA.

Introduction

Idiopathic Anaphylaxis (IA) is a diagnosis of exclusion after known causes for anaphylaxis and other diseases that mimic anaphylaxis have been ruled out [1]. The condition was first described in 1978 [2]. The exact prevalence is not known but has been estimated to be between 20,000 and 47,000 cases in the United States [3].

IA accounts for up to 60% of cases of anaphylaxis in adults and 10% of pediatric cases [4]. Given that as per the definition there are no identifiable triggers of IA, there are substantial challenges related to the diagnosis and management of these cases. There are currently no guidelines regarding the diagnostic approach including assessment for underlying diseases and use neither of confirmatory tests nor on the best management of pediatric cases of IA. The current approach to treatment is based on disease frequency; short-term treatment for infrequent attacks, but prophylactic treatment with glucocorticoids is recommended for patients with frequent episodes. It is well known however that long-term treatment with systemic glucocorticoids is associated with multiple side effects that may be particularly worrisome in the pediatric population including growth failure and adrenal suppression. Although several case series have been published regarding IA, only a few small descriptive studies have focused on pediatric idiopathic anaphylaxis. We aim to present a series of 3 cases of pediatric IA followed by a
systematic review of the literature published in the past 22 years regarding IA in children, with a focus on diagnosis and management.

**Methods**

We searched the PubMed database for original scientific studies pertinent to the clinical diagnosis and management of idiopathic anaphylaxis in children. We used the search terms “Idiopathic anaphylaxis” AND “Children” and then limited the results to articles published between May 12, 1995 and May 12, 2017, that were written in English or French.

The abstracts of the resulting papers were reviewed and those relevant to the diagnosis and management of idiopathic anaphylaxis in children were included. The initial PubMed search yielded 58 articles, which was reduced to 19 after applying previously mentioned filters and then further narrowed to 8 articles after the first and last authors (SD and MBS) assessed the abstracts and discussed their relevancy to the systematic review. (Figure 1)

![Fig. 1: Results of the systematic review using PubMed database. Excluded papers were either review articles rather than original papers or not relevant to the diagnosis or management of idiopathic anaphylaxis in children.](image-url)
<table>
<thead>
<tr>
<th>Type of Study</th>
<th>Author, Journal Pub yr</th>
<th>Peds/Adults</th>
<th>Findings</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Retro-spective chart review</td>
<td>Krasnick et al 1996</td>
<td>A &gt; C</td>
<td>Charts of 225 patients diagnosed with idiopathic anaphylaxis (IA) were reviewed at a single center and 61 patients were available for long-term F/U. <strong>Diagnosis:</strong> based on history, PE, serology, radiology, SPT food and aeroallergens, challenges with food additives. Age range was between 10-68 years old, <strong>mean 39 yo,</strong> the majority was female. <strong>Treatment</strong> based on frequency of symptoms: <strong>Infrequent:</strong> treatment of acute attacks only with Epi, Pred 60 mg, antihistamine. <strong>Frequent:</strong> Pred 60-100 mg x 1 week, alternate day, tapering by no more than 5 mg per month; daily antihistamines and sympathomimetics (albuterol). <strong>Long-term F/U</strong> 65% of patients with frequent attacks and 91% of patients with infrequent attacks went in remission. Significant decrease in ED visits (except IA-AE-F), hospital admission and ICU admission after evaluation and treatment by allergy service. Reduction 184 740 $ saved by program for 546 patient-years (estimated 11 million $ saved for 33 000 cases of IA in US). Cromolyn not effective. Ketotifen effective in reducing amount of prednisone and in some stop prednisone. Greatest clinical benefit IA-G (F and IF): significant decrease in ED visit.</td>
<td>Small N of pediatric patients. No specific information regarding children in this study (percentage, clinical presentation, management?). High remission rate for patients with infrequent episodes. Spontaneous remission vs aggressive treatment of acute episodes decreases frequency and severity of future episodes. A number of patients in remission were still on daily antihistamines which may have an effect on disease control. No control group with placebo for ethical reasons.</td>
</tr>
<tr>
<td>Case series</td>
<td>Ditto et al, Ann All, 1996</td>
<td>Adults &gt; Children</td>
<td>335 patients were evaluated for episodes of anaphylaxis of unknown origin. N=132 were evaluated. <strong>Diagnosis:</strong> R/O food, drug, exercise 4 hours prior to Rx, R/O pheochromocytoma and mastocytosis, SPT to aeroallergens if atopic history, complement levels R/O AE. <strong>Treatment</strong> according to previous protocol and based on frequency of symptoms: <strong>Infrequent:</strong> treatment of acute attacks only with Epi, Pred 60 mg, antihistamine. <strong>Frequent:</strong> Pred 60-100 mg x 1 week, alternate day, tapering by no more than 5 mg per month; daily antihistamines (hydroxyzine) and sympathomimetics (albuterol). <strong>N=14/335 children (4.2%) were included</strong> N=4 between 0-9 yr (1.2%) N=28 between 10-19 yr (8.4%) <strong>Increasing incidence in children (9 new cases since previous study)</strong> <strong>increasing prevalence vs increasing awareness</strong> Children on same treatment protocol as adults and similar response as adults 64% frequent episodes requiring prednisone (3/56 cortico-steroid dependent IA) vs 36% infrequent. Remission rate overall 65% (57/132) Time to remission 1-14 years 48% atopy, 23% pre-existing urticaria/AE, 5% unrelated food allergies, 11% exercise induced episodes.</td>
<td>Small N of pediatric patients. No specific description of pediatric cases. Food SPT, laboratory test (CBC, complement and serum chemistry) non contributory - not recommended except in selected patients. No control group.</td>
</tr>
<tr>
<td>Citation</td>
<td>Study Details</td>
<td>Children</td>
<td>Adults &gt; Children</td>
<td></td>
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| Ditto et al, JACI 1997           | 22 cases of IA in children (9 new, 13 previously described in study published by Ditto in 1996) were evaluated | Diagnosis: Thorough history and PE, R/O food, drug, exercise 4 hours prior to Rx, R/O pheochromocytoma and mastocytosis, SPT to Aeroallergens if atopic hx. Treatment according to previous protocol and based on frequency of symptoms: | 13/81 (16%) children less than 14 yrs Mean age 30 +/- 17.3 yrs Age range 5-73; Median 24 years 68% F.  
N=81 cases of IA1990-1995  
13/81 (16%) children less than 14 yrs Mean age 30 +/- 17.3 yrs  
Age range 5-73; Median 24 years 68% F.  
Median N of episodes during a year was 2, range 1-130  
Median F/U = 24 months  
N=74/81 F/U until 12/1995  
Remission 56/74 (76%)  
N18/81 (22%) frequent episodes and proposed to start maintenance w corticosteroids  
12/18 agreed, 2/18 refused, 4/18 had history of use of cs with improvement  
Patients who received corticosteroids had a significant decrease in attacks between 1 year before and after F/U  
15% other anaphylaxis (food, exercise, drugs)  
48% atopy  
58% acute or CSU or AE |
| Ditto et al, JACI 1997           | Diagnosis: Thorough history and PE, R/O food, drug, exercise 4 hours prior to Rx, R/O pheochromocytoma and mastocytosis, SPT to Aeroallergens if atopic hx. Treatment according to previous protocol and based on frequency of symptoms: | Infrequent: treatment of acute attacks only with Epi, Pred 60 mg, antihistamine  
Frequent: Pred 60-100 mg x 1 week, alternate day, tapering by no more than 5 mg per month; daily antihistamines (hydroxyzine) and sympathomimetics (albuterol)  
Age at onset 3 - 17 yo, majority were female F > M 13:9  
- 12/22 frequent attacks, 9/22 infrequent, 1/22 undifferentiated somatoform idiopathic anaphylaxis  
- 2/22 CSD-IA (Tx with Ketotifen, 1 went in remission, 1 tapering daily corticosteroids)  
- 4/22 isolated throat angioedema Tx (inhaled beclomethasone dipropionate) with success  
- 3/22 malignant IA uncontrolled with alternate day prednisone (1/3 in remission, 1/3 Ketotifen: improved, 1/3 US-IA (non-organic disease), no improvement despite high dose CS) | N=74/81 F/U until 12/1995  
Remission 56/74 (76%)  
N18/81 (22%) frequent episodes and proposed to start maintenance w corticosteroids  
12/18 agreed, 2/18 refused, 4/18 had history of use of cs with improvement  
Patients who received corticosteroids had a significant decrease in attacks between 1 year before and after F/U  
15% other anaphylaxis (food, exercise, drugs)  
48% atopy  
58% acute or CSU or AE |
| Ditto et al, JACI 1997           | Charts of 8 children with IA were evaluated  
No identifiable allergen, no other systemic disease  
7/8 infrequent 1/8 frequent (tx with steroid; off steroids since 6 months)  
All cutaneous sx 6/8 resp sx (wheezing or AE w breathing difficulties) 6/8 diarrhea  
Many had life-threatening events: hypotension or LOC  
Other difficulties: Non compliance, non-belief, taken out of school  
Can start at any age, majority after 10 y of age | N=74/81 F/U until 12/1995  
Remission 56/74 (76%)  
N18/81 (22%) frequent episodes and proposed to start maintenance w corticosteroids  
12/18 agreed, 2/18 refused, 4/18 had history of use of cs with improvement  
Patients who received corticosteroids had a significant decrease in attacks between 1 year before and after F/U  
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| Hogan, Annals 1998               | Small study  
None in remission  
2 young patients at time of onset (11 months) |  
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Small study  
None in remission  
2 young patients at time of onset (11 months) |
Retro-Spective chart review + Follow-up
Lieberman et al Annals 2006
Adults > Children
To examine natural history, clinical manifestations ad factors compliance to Tx
Retrospective chart review 25 years + F/U question
N=601 cases of anaphylaxis of unknown origin
Age range 1-79 years, mean 37 yrs, F 62%
Causes in 41% of patients (food, drug, exercise)
Mastocytosis N=3/601
IA 59% increasing N of idiopathic anaphylaxis
Attacks decrease in time
Small N of children, no specifics about pediatric population

Case report
Warrier, Annals 2009
Child
Omalizumab in IA; 1st case report in child
12 yo uncontrolled asthma requiring multiple courses of oral steroids, AR,
multiple FA, recurrent episodes of anaphylaxis of unknown origin (4 to 5 per month);
Atopic; Total IgE 1848 IU/ml
Tx Omalizumab 375 mcg q 2weeks: resolution of acute episodes, anaphylaxis
and asthma exacerbations; Improved QOL

Case series
Ivkovic PAI 2015
Children
Report of 3 patients with IA, HI and positive ASST
Age at presentation 11, 15, 12.6 years
Diagnosis: All 3 had positive ASST to assess auto-reactivity (Indication of mast-cell activating auto-antibodies against IgE or high affinity receptor)
Basophil histamine release assay was negative in 3 patients to demonstrate auto-antibody specificity
Maybe different cause for auto-reactivity
DAO (histamine-deactivating enzyme Diamine Oxidase) levels decreased;
indicating high level of histamine Intolerance
Treatment
Patient 1: generalized infrequent episodes, Epi PRN, on histamine free diet
alone no improvement, started with antihistamines (Cetirizine 10 mg QD) and improved
Patient 2: generalized infrequent, on histamine free diet, Epi PRN, lost to F/U
Patient 3: frequent episodes tx with Epi, antihistamines and steroids; advised
a histamine free diet and oral steroid on maintenance but refused; on daily
antihistamines (Fexofenadine) decrease in frequency and severity of attacks; in
remission since 6 months
First reports of positive ASST in children and decreased DAO levels (histamine intolerance)
Treatment with histamine free diet, antihistamines (Cetirizine 10 mg QD

Table 1: The articles that were reviewed are summarized in table.

Case presentations
Case 1
An 11 year-old boy presented initially with an episode of cough and eyelid swelling 15 minutes after completing exercise (hockey game). He had eaten pasta 2 hours prior to his exercise.
A year later he developed a similar episode with angioedema, difficulty breathing and abdominal pain after eating pizza. He had done exercise 6 hours prior to eating the pizza. At that point he was seen at the emergency department and he was treated with 3 doses of IM epinephrine. An extensive work-up was performed. The boy had symptoms of seasonal rhino conjunctivitis. Skin prick tests were performed with common aeroallergens and sensitizations to ragweed and tree pollens was confirmed. Skin prick tests with suspected food allergens were negative as well as specific IgE to possible food allergens including ω-5 gliadin and alpha-gal (Alpha gal testing was performed at Thomas Platts-Mill laboratory, University of Virginia, United States). A challenge with the suspected meal was negative.
He had an elevated total IgE (571 kU/L) and his tryptase level was within normal range (3.5 - 4.5 µg/L) at baseline. Tryptase levels were not documented during his attacks. He had normal levels of C3, C4 and C1 inhibitor esterase. (1.57 g/L, 0.20 g/L, 0.32 g/L respectively). Further investigation revealed no mutation for the cKIT 816 on peripheral blood (conduct based on PCR levels - Thomas Kristiansen laboratory, Dept. of Pathology, Odense University Hospital, Denmark).
Parents refused bone marrow biopsy.

Given the association with exercise we performed an exercise
and food-dependent exercise challenge that was normal.

He was instructed to carry an Epipen auto-injector and in the following year he had 3 other similar episodes. The last episode occurred at 15 years of age. He had no new reactions over the last 24 months.

**Case 2**

A 14 year-old girl presented with 2 episodes of anaphylaxis with abdominal cramps, flushing and hives. The first episode occurred in the context of exercise after eating tree-nuts and wheat. One month later she presented with a second episode occurring 1 hour after eating meat. She was treated with Epinephrine and Reacting. Specific IgE levels were negative to all suspected foods including tree-nuts, wheat, milk and alpha-gal. Tryptase levels during the reaction (17.2 µg/L), with baseline levels within normal range (7.5 µg/L).

Further investigation revealed undetectable levels of alpha-gal (Thomas Platts-Mill laboratory, University of Virginia, United States) and no mutation for the cKIT 816 on peripheral blood (Thomas Kristensen laboratory, Dept. of Pathology, Odense University Hospital, Denmark). Exercise challenge test (with and without prior exposure to the suspected meal) and spirometry revealed no abnormalities.

She was instructed to carry an Epipen auto-injector and we advised to begin a treatment with Reactine 20 mg daily. Parents refused a bone marrow biopsy at this point. At a 24-month follow-up she remained in remission.

**Case 3**

The third patient presented at 16 years of age with recurrent anaphylaxis. Her episodes were characterized by acute abdominal pain, vomiting, diarrhea followed by itchiness and hives. The episodes would occur 6 to 7 times per month, usually at night between 10-11 pm (2 hours after eating). She would do exercise 1 hour prior to eating (30 min on a stationary bike). The episodes seemed to occur in the context of her menstruation (a few days before, during or a few days after). Otherwise, no clear triggers could be identified. In between episodes she was symptom-free and had no constitutional symptoms.

She had an extensive work-up with negative skin prick tests and specific IgE for the relevant food allergens. Her total IgE was significantly elevated (4032 kU/L) and tryptase levels were elevated during attacks (17.4 µg/L) and normal at baseline (3.4 µg/L). There were undetectable levels of alpha-gal (Thomas Platts-Mill laboratory, University of Virginia, United States) and the cKIT816 mutation was not detected in peripheral blood (Thomas Kristensen laboratory, Dept. of Pathology, Odense University Hospital, Denmark).

Given the severity of episodes and the prominent gastrointestinal manifestations, an abdominal ultrasound was done and revealed no abnormalities. Urine collection over 24 hours for 5HIAA and VMA was normal. A bone marrow biopsy was performed revealing no abnormal mast cell infiltrates and normal staining for cKIT and tryptase.

She was instructed to carry an Epipen auto-injector and we advised to begin treatment with Reacting 20 mg daily that did not improve her symptoms.

Treatment with Omalizumab 300 mg monthly was started and she has been in remission since then, for one year. Four months after stopping Omalizumab treatment she has not experienced new episodes consistent with anaphylaxis. The findings for each patient are listed in (Table 2).

<table>
<thead>
<tr>
<th>Clinical characteristics</th>
<th>Patient 1</th>
<th>Patient 2</th>
<th>Patient 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age at presentation (years)</td>
<td>11</td>
<td>14</td>
<td>16</td>
</tr>
<tr>
<td>Gender</td>
<td>Male</td>
<td>Female</td>
<td>Female</td>
</tr>
<tr>
<td>Presence of atopy</td>
<td>Yes</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>Frequency of attacks</td>
<td>Infrequent</td>
<td>Infrequent</td>
<td>Frequent</td>
</tr>
<tr>
<td>SPT</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Ragweed</td>
<td>+</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>Tree</td>
<td>+</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>Grass</td>
<td>-</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>House dust mite</td>
<td>NA</td>
<td>NA</td>
<td>+</td>
</tr>
<tr>
<td>Wheat</td>
<td>-</td>
<td>NA</td>
<td>-</td>
</tr>
<tr>
<td>Soy</td>
<td>-</td>
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<td>NA</td>
</tr>
<tr>
<td>Almonds</td>
<td>-</td>
<td>NA</td>
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</tr>
<tr>
<td>Peanut</td>
<td>NA</td>
<td>NA</td>
<td>-</td>
</tr>
<tr>
<td>Celery</td>
<td>NA</td>
<td>NA</td>
<td>-</td>
</tr>
<tr>
<td>Tomatoe</td>
<td>NA</td>
<td>NA</td>
<td>-</td>
</tr>
<tr>
<td>Prick to prick</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fresh shrimp</td>
<td>-</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>Pizza</td>
<td>-</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>Fresh noodles</td>
<td>-</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>Total IgE</td>
<td>571</td>
<td>804</td>
<td>4032</td>
</tr>
<tr>
<td>(IU/ml)</td>
<td>sIgE</td>
<td></td>
<td></td>
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<tr>
<td>(IU/ml)</td>
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</table>
Diagnosis of IA

Idiopathic anaphylaxis is a diagnosis of exclusion and should be made only after extensive evaluation to exclude other potential causes of anaphylaxis and other diseases with similar manifestations. The classification of IA is based on frequency of episodes and clinical manifestations.

The majority of studies included in this review (Table 1) describe an extensive evaluation in adults and children with anaphylaxis of unknown origin including diagnostic testing for possible triggers of IgE mediated anaphylaxis such as food, drugs and exercise 4 hours prior to reaction. In patients with an atopic history, Skin Prick Tests (SPT) to aeroallergens is usually performed. C1-inhibitor levels are measured in patients presenting with angioedema to rule out hereditary angioedema. Rare diseases such as systemic mastocytosis, carcinoid syndrome or pheochromocytoma are usually considered. The extensive diagnostic work-up makes the diagnosis of IA highly probable in those series [4,5].

In a large cases series of 335 patients, including adults and children, authors concluded that SPT for foods and laboratory testing (CBC, complement and serum chemistry) were non-contributory and should be reserved for selected patients [6].

In a case series published by Hogan et al, 8 children were evaluated for IA when no identifiable allergen was found and in the absence of other systemic disease. However, the diagnostic tests performed were not laborated on in the manuscript [7].

In a Spanish series published in 2005, including mostly adults and a small proportion of children(mean age 30.0 ± 17.3 years, age range 5-73 years), all patients had an extensive evaluation as previously described as well as excluding insect bites, latex and other rare causes such as anaphylaxis attributed to Alpha gal in these 3 cases was excluded based on medical history (no data or signs of tick bites, and no symptoms related to ingestion of mammalian products). Interestingly all 3 patients had a thorough investigation as previously mentioned studies in which no systemic diseases were found [4]. More recently, 3 case reports of IA were published in which all 3 patients had a thorough investigation as previously described [5].

The authors suggest that rare disorders that may mimic IA include delayed anaphylaxis to red meat. In contrast with immediate food-mediated anaphylaxis, symptoms may occur more than 2 hours after exposure. Reaction to mammalian oligosaccharide α-gal in these 3 cases was excluded based on medical history (no data or signs of tick bites, and no symptoms related to ingestion of mammalian products). Interestingly all 3 patients were found to have a positive Autologous Serum Tests (ASST), buta histamine release test was negative and the presence of circulating auto-antibodies against IgE and FcεRIα could not be confirmed.

In addition they determined the activity of histamine-inactivating enzyme (DAO) which was very low in all three patients, indicating low histamine intolerance. Histamine intolerance describes a state with decreased ability to catabolise endogenous or exogenously administered histamine, leading to histamine-mediated adverse reactions. Diagnosis is based on careful clinical history and identification of intestinal or serum activities of DAO and histamine N-methyl transferase. However, there are

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Epinephrine auto-injector PRN</th>
<th>H1 histamines Epinephrine auto-injector PRN</th>
<th>H1 histamines Epinephrine auto-injector PRN Oma lizumab</th>
</tr>
</thead>
</table>

Table 2: Patient characteristics.

Results

Diagnosis of IA

Table 2: Patient characteristics.
Discussion

Although IA accounts for only 10% of anaphylaxis cases in children, it represents a major diagnostic and management challenge given the lack of confirmatory tests inability to avoid potential triggers. Based on the existing literature, a diagnosis and management algorithm specifically targeting pediatric cases of IA is required. Based on current literature we suggest considering the following diagnostic tests when evaluating a patient with possible IA:

A thorough clinical history and physical exam should be performed followed by laboratory investigations. Immediate skin testing and/or in vitro tests for specific IgE to potentially relevant drugs or foods may help to confirm or exclude food or drug allergy as a possible trigger including sIgE to ω-5 gliadin and alpha-gal in the case of a suggestive history.

Delayed anaphylaxis to red meat is a rare disorder that may mimic IA. It should be suspected as a culprit in any case of anaphylaxis of an unknown origin, especially in events occurring 3 to 6 hours after eating, particularly when reactions start in the early morning hours. Our 3 patients had negative sIgE to the relevant oligosaccharide galactose-α1,3-galactose. Other potential causes that should be considered are exercise and food-dependent exercise-induced anaphylaxis.

Exercise and/or food-dependent exercise challenges should thus be considered in the case of a suggestive history.

Other disorders that may mimic IA are systemic mastocytosis, monoclonal mast cell activation syndrome and mast cell activation syndrome.

Elevated serum tryptase levels during attacks are suggestive for IA and patients with elevated levels both at baseline and during attacks should be evaluated with a bone marrow biopsy to rule out mastocytosis or other clonal mast cell disorders.

Given studies supporting the comparison of serum tryptase levels at baseline and during reaction to confirm the presence of anaphylaxis [12], patients with IA should have tryptase levels done at these two time points. All 3 of our patients had a normal baseline tryptase and a negative cKit 816 mutation analysis on peripheral blood. Our third patient who presented with generalized frequent episodes of IA had a normal bone marrow biopsy.

Pheochromocytoma and carcinoid syndrome are rare disorders in which patients may present with symptoms similar to anaphylaxis, such as flushing induced by release of vasoactive substances (vanillylmandelic acid or VMA in pheochromocytoma and 5-hydroxyindoleacetic acid or 5HIAA in carcinoid syndrome). Detection of these mediators in 24-hour urine may help exclude or confirm the diagnosis in patients with a suggestive history. Our third patient had a normal abdominal ultrasound as well as a normal 24 urine for 5HIAA and VMA and hence repeated tests are required if symptoms persist and suspicion is high.

Hereditary angioedema may mimic IA presenting with recurrent episodes of angioedema as principal symptom. Laboratory findings should include serum levels of C4 and C1 inhibitor in selected patients. More recently, 3 previously described case
When suggestive history to confirm or exclude pheochromocytoma/carcinoid syndrome.

To exclude or confirm systemic mast-cell disease.

More interestingly, decreased DAO levels, indicative of histamine intolerance were demonstrated in all 3 patients [5]. Whether these findings are coincidental or play a role in the underlying pathogenesis of IA remains to be further evaluated. Given the small number of patients, it would be interesting to confirm these findings in larger studies and to look at ASST and DAO levels in patients presenting with IA. (Table 3) summarizes diagnostic testing to consider in patients with possible IA.

<table>
<thead>
<tr>
<th>Diagnostic test</th>
<th>Comment</th>
</tr>
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<tbody>
<tr>
<td>Skin prick test and/or sIgE for food, intradermal test for insect stings or drug</td>
<td>Based on suggestive clinical history, to confirm or exclude other causes of IgE-mediated anaphylaxis</td>
</tr>
<tr>
<td>sIgE for alpha-gal</td>
<td>When suggestive history to confirm or exclude alpha-gal allergy</td>
</tr>
<tr>
<td>Exercise challenge</td>
<td>When suggestive history, to confirm or exclude exercise-induced anaphylaxis</td>
</tr>
<tr>
<td>Oral food challenge plus exercise challenge</td>
<td>When suggestive history, to confirm or exclude food-dependent exercise-induced anaphylaxis</td>
</tr>
<tr>
<td>Serum tryptase</td>
<td>At baseline and during reaction; suggestive for IA when elevated during reaction; suggestive of mastocytosis when elevated at baseline and during reaction</td>
</tr>
<tr>
<td>Bone marrow biopsy</td>
<td>To exclude or confirm systemic mastocytosis</td>
</tr>
<tr>
<td>cKIT 816 mutation on peripheral blood</td>
<td>To exclude or confirm systemic mastocytosis</td>
</tr>
<tr>
<td>24 h - urine for VMA/SHIAA</td>
<td>To exclude or confirm pheochromocytoma/carcinoid syndrome</td>
</tr>
<tr>
<td>Abdominal ultrasound</td>
<td>To exclude or confirm pheochromocytoma/carcinoid syndrome</td>
</tr>
<tr>
<td>Complement levels</td>
<td>To confirm or exclude hereditary angioedema when suggestive history</td>
</tr>
<tr>
<td>C1 inhibitor levels +/- function</td>
<td>To confirm or exclude hereditary angioedema when suggestive history</td>
</tr>
<tr>
<td>Autologous serum skin test (ASST) and diamine oxidase (DAO) levels</td>
<td>To be considered</td>
</tr>
</tbody>
</table>

Table 3: Diagnostic tests to consider in a patient with possible IA.

The current proposed treatment regimens are based on small studies from long ago, and daily treatment with systemic steroids in children is associated with an increased risk of long-term side effects. As the risks of systemic steroids often outweigh the benefits in children, we do not recommend systemic steroids for treatment of IA in children. Based on recently published case reports and our own clinical experience, we would suggest the following algorithm for the management of pediatric IA:

All patients should be instructed on the management of acute episodes and should be prescribed an epinephrine auto-injector. In patients with frequent episodes, daily H1-antihistamines at regular doses should be prescribed and the patient should have close clinical follow-up. If no improvement after 1 month the physician should consider increasing the daily dose to 2 times and up to 4 times the regular dose.

If no improvement upon increased daily anti-histamines after 1 or 2 months, treatment with Omalizumab should be considered. Omalizumab is a 95% humanized monoclonal antibody that recognizes and binds to the FC portion of IgE. Allergen specific IgE plays a central role in IgE mediated anaphylaxis. Cross-linking of receptor bound IgE on mast cells and basophils by allergens leads to activation of those cells and subsequent release of inflammatory mediators causing anaphylaxis. The exact mechanism in IA remains unknown but it has been postulated that the high affinity IgE receptors may be cross-linked by autoimmune mechanisms.

Omalizumab may prevent IgE expression on effector cells and subsequent cross-linking of IgE. It has been a successful treatment in other IgE-mediated conditions such as asthma and chronic urticaria. Here we report a second case of pediatric IA successfully treated with Omalizumab. Although we could argue that time could have played a role in achieving remission, the rapid and significant improvement supports the positive therapeutic effect of Omalizumab.

Conclusion

In conclusion few studies have evaluated diagnosis and management of pediatric idiopathic anaphylaxis. Future studies are needed to optimize treatment regimens for IA in children, who are particularly at risk for potential side effects of oral steroids.

Omalizumab maybe a therapeutic option for pediatric IA as a novel approach to treatment. Based on our clinical experience we would recommend treatment of acute attacks with epinephrine for patient with infrequent episodes and a stepwise approach for patients using a potential algorithm we have constructed for the diagnosis and management of IA. The usefulness and accuracy of these algorithms should be assessed in future large-scale studies.

References


