Acute hemorrhagic edema of infancy: a worrisome presentation, but benign course

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Abstract: Acute hemorrhagic edema of infancy (AHEI) is a benign type of leukocytoclastic vasculitis. It is a benign phenomenon although it manifests with fever, large palpable purpuric skin lesions, and edema. The presentation of AHEI can often resemble that of Henoch-Schönlein purpura. Since AHEI is a self-limited disease, conservative management is the most commonly followed approach. Our patient had complete resolution of AHEI without medical treatment.

Keywords: infancy, hemorrhagic, edema

Introduction
Acute hemorrhagic edema of infancy (AHEI) is a benign type of leukocytoclastic vasculitis. It has also been described as Finkelstein’s disease, Seidlmayer’s disease, and postinfectious cockade purpura.1 AHEI is considered a benign phenomenon and it usually resolves spontaneously in a few weeks.2 We are presenting a 14 month old male with AHEI who presented with rash on the face and extremities, as well as edema in the left lower extremity.

Case report
A 14 month old boy presented to our emergency department with intermittent low grade fever of 2 days duration associated with a rash, swelling of the left leg, and an inability to bear weight. The mother stated that the patient had a runny nose and a mild cough 1 week prior to admission.

There was no diarrhea, change in behavior, toxic ingestion, medication use, or any recent vaccinations. Past medical history and family history were unremarkable and immunizations were up to date. The patient was admitted to the ward for fever and rash. On admission, his temperature was 38°C, he had a respiratory rate of 25 breaths per minute, his heart rate was 110 beats per minute, and his blood pressure was 100/70 mmHg.

On examination, the patient was not looking toxic, his neck was supple, and the neurological exam was unremarkable. The skin showed ecchymotic, purpuric, targetoid plaques with oval and round shapes on the face and extremities (Figures 1–3). The right auricle of the ear was mildly edematous and purple in color. There was mild swelling on the hands and lower extremities without skin induration and the range of motion of all joints was normal. The rest of the physical exam was unremarkable. The complete blood count and urine analysis were both unremarkable.
A decision was made to biopsy one of the lesions from the left forearm and it showed inflammatory infiltrate surrounding dermal blood vessels. The infiltrate was a mixture of neutrophils, lymphocytes, and scattered eosinophils with fragmented nuclei of neutrophils, and there was focal necrosis of the vessel wall. On direct immunofluorescence there was no immunoglobulin A (IgA) deposition. The patient was discharged after the diagnosis and the rash resolved spontaneously in 3 weeks.

**Discussion**

AHEI is more common in males,\(^2\) without racial predominance.\(^3\) The age group is typically between 4 and 24 months.\(^1\) In the majority of the cases, AHEI is a self-limited disease (1–3 weeks), but there have been reports of recurrent attacks (up to four attacks).\(^4\)

Patients with AHEI are usually nontoxic on presentation\(^2\) and they classically present with fever, large palpable purpuric skin lesions, and edema. The skin changes are characterized as well demarcated, annular, medallion-like, rosette shaped purpuric plaques, and tend to appear suddenly in the face and extremities.\(^1\) However, the skin lesion does not always present in a typical way and it has been reported that it can appear as hemorrhagic blisters.\(^3\) The edema is typically on the face, extremities, and auricles.\(^2\)

Many organisms including adenovirus, vermicelli zoster virus, cytomegalovirus, herpes simplex virus, tuberculosis, streptococci, and staphylococci are associated with AHEI.\(^4\) Almost 75% of cases of AHEI are preceded by infections like otitis media, urinary tract infection, upper respiratory infection, and pneumonia. An immune complex hypersensitivity complex has been suggested as the etiology of AHEI since vaccines and medications are trigger factors.\(^2\)

Extracutaneous and viscera involvement are rare with AHEI. However, Fiore et al documented two cases with intestinal engagement\(^6\) and Wanatabe and Sato reported a case with renal involvement and hypocomplementemia.\(^6\)

Blood laboratory results are usually nonspecific. In a complete blood count there might be thrombocytosis, leukocytosis with eosinophilia, or lymphocytosis. In addition, C-reactive protein and the erythrocyte sedimentation rate may be high.\(^7\) In some cases, patients with AHEI can present with hypocomplementemia affecting complement C4, complement C1q, and complement CH50,\(^8\) and transient abnormal liver function.\(^9\)

A skin biopsy for histopathology and immunofluorescence testing is useful if diagnosis is unclear.\(^10\) The typical description of the histology in the patient with AHEI is perivascular neutrophilic infiltration with several nuclear fragments in the vascular wall, resulting in fibrinoid necrosis.\(^7\) The rare serious complications of AHEI are renal involvement, gastrointestinal bleeding, epididymo-orchitis, and cartilage damage.\(^11\)

Since AHEI is a self-limited disease, conservative management is the most commonly followed approach.\(^2\) Steroid use is warranted when there is a severe or persistent gastrointestinal symptom or complication of the disease. The use of immunosuppressive medications like cyclophosphamide or azathioprine to prevent renal damage has been reported as futile.\(^10\) Antibiotics can be used if there is a concomitant bacterial infection.\(^2\)

AHEI can present as erythema multiforme, but the latter does not show leukocytoclastic vasculitis. AHEI can also mimic urticarial vasculitis, but the former usually has an acute presentation with rare remissions.\(^8\) The presentation of AHEI can often resemble that of Henoch-Schönlein purpura (HSP) and some authors consider AHEI as a cutaneous
variant of HSP. It is crucial to differentiate between AHEI and HSP since the latter requires systemic corticosteroid therapy. AHEI and HSP are similar in that both have skin lesions and leukocytoclastic vasculitis. However, the differences are as follows. AHEI presents in a younger population (3 to 24 months old) in comparison to HSP (3 to 6 years old). The skin lesions in AHEI manifest as ecchymotic changes with large purpura and edema involving the face, extremities and scrotum, while the purpura in HSP is usually smaller and involves the buttocks and the legs. Involvement of renal, gastrointestinal, and viscera are rare in AHEI when compared to HSP. IgA levels are usually normal in AHEI, but elevated in HSP. There is usually no IgA deposition in AHEI compared to HSP. Finally, complement C1q deposition is more diagnostic of AHEI.

**Conclusion**

AHEI is a self-limited disease that usually resolves spontaneously. It is crucial to make the diagnosis correctly to avoid unnecessary work up and therapy, while simultaneously monitoring for rare but severe complications.

**Disclosure**

The authors report no conflicts of interest in this work.

**References**


**Figure 3**

Circular and oval plaques on the upper extremities with swelling more prominent on the right side.