Thermal Angioedema Induced by Hot Water

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Although angioedema is caused by C1 inhibitor deficiency, drugs and various disease, cases linked only to physical irritation are extremely rare. We report here a case of thermal angioedema caused by showering in hot water.

CASE REPORT

A 20-year-old Japanese woman presented with a 1-year history of recurrent episodes of eyelid swelling without major urticaria. The swelling occurred only when she took a hot shower or bath. Symptoms lasted for a few days and improved spontaneously. On one occasion, respiratory discomfort accompanied the eyelid swelling. The patient was consuming no agents associated with such episodes, including foods, medicines or supplements, although she did have a history of childhood atopic dermatitis and bronchial asthma. She had also suffered a single episode of oral allergy to kiwi fruit without major urticaria or angioedema. She had no history of cholinergic or physical urticaria linked to cold, solar radiation, contact heat, vibration or water.

The patient was treated with chlorpheniramine and olopatadine hydrochloride, but the swelling continued for two days. Her serum histamine levels were already elevated before challenge and did not increase markedly 20 min after the symptoms developed. Although we have no evidence to prove it, this phenomenon does suggest that antihistamines may inhibit early reactions induced by histamine, but not later reactions that are probably caused by other chemical mediators and complement activation.

The histamine levels in our patient were already elevated before challenge and did not increase markedly 20 min after the symptoms developed. Although we lack sufficient data to explain the reason for this, her histamine levels may have been constantly high. Moreover, since the half-life of histamine in the serum is very short, we may have detected a significant increase in her histamine levels if we had measured them immediately after the symptoms developed.

Zingale et al. (2) described the pathogenic characteristics in 776 cases of angioedema without major urticaria as follows: unknown origin, 38%; C1 inhibitor deficiency, 25%; exogenous stimulus, 16%; ACE inhibitor, 11%; and autoimmune disease or infection, 7%. They also reported the frequencies of each type of exogenous stimulus in 124 cases as follows: medication 45%; food 36%; both 8%; insect bite 4%; other environmental allergens 3%; and other minor stimuli, including physical irritation 3%. The details of the cases linked to minor stimuli were not described.

The histamine levels in our patient were already elevated before challenge and did not increase markedly 20 min after the symptoms developed. Although we lack sufficient data to explain the reason for this, her histamine levels may have been constantly high. Moreover, since the half-life of histamine in the serum is very short, we may have detected a significant increase in her histamine levels if we had measured them immediately after the symptoms developed.

Recent, recommendations for diagnostic testing of physical urticaria have been reported (6, 7). The result of the challenge test in our case indicated that angioedema was induced by hot showering of the whole body rather than a local increase in temperature. In addition, while the symptoms were blocked by the preventive administration of an antihistamine, antihistamine administration after symptoms had developed produced no improvements. Although we have no evidence to prove it, this phenomenon does suggest that antihistamines may inhibit early reactions induced by histamine, but not later reactions that are probably caused by other chemical mediators and complement activation.

Investigations of cases of angioedema should be tailored to each patient’s symptoms. The challenge test is important for determining the cause of the symptoms and for therapeutic planning.

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Fig. 1. Swelling of both eyelids induced by showering at 41°C for a few minutes.
The authors declare no conflict of interest.

REFERENCES