Food related antibodies in headache patients

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SUMMARY

Highly sensitive and specific methods for assaying IgE and IgG4 for antibodies in serum have been developed in order to test a recent suggestion that food allergy is a major cause of migraine. Sera were collected from 208 adults—74 with dietary migraine, 45 with non-dietary migraine, 29 with cluster headache and 60 controls. No significant differences were identified between any of the groups with the one exception that cluster headache patients had significantly raised levels of total serum IgE: this difference may be explained by the high proportion of smokers. Mean IgG4 titres to cheese, milk and chocolate were very similar in the dietary and non-dietary migraine patients and the control subjects, though lower in the cluster headache patients. There was, therefore, no evidence that the food intolerance often associated with migraine headaches is associated with a conventional allergic mechanism.

A minority of migraine patients report that headaches can be precipitated by specific foods; the commonest of these among 494 migraine patients attending the Princess Margaret Migraine Clinic, Charing Cross Hospital, London, were cheese (19% sensitive), chocolate (18% sensitive) and citrus fruit (11% sensitive). These particular foods are all believed to have a high content of amines such as tyramine, but evidence that headaches can be reliably reproduced by the administration of pure tyramine remains conflicting.

Immunologically mediated skin and gastrointestinal responses to foods are now well established and there is evidence of circulating immune complexes and complement activation before attacks of common migraine, though not in classical migraine. Although elevated total serum IgE levels were no commoner in migraine or tension headache patients, evidence was recently presented that two thirds of severely affected migraine patients were allergic to some foods. Although these authors confirmed the clinical observation by the use of the Radio-allergosorbert test (RAST), the reported IgE levels seemed rather low, perhaps partly due to non-specific binding in the in-vitro test.

This paper re-investigates the role of IgE-mediated food allergy in migraine by using a highly sensitive IgE RAST technique to determine circulating levels of specific IgE to four foods commonly supposed to "trigger" migraine. Specific IgG4 levels to the same foods were also quantified because it has been suggested that IgG4 is the short-term sensitising IgG described by Parish and that it is responsible for the late response seen in some cases of asthma.

Patients and methods

Serum samples were taken from a total of 148 headache patients attending the Princess Margaret Migraine Clinic at Charing Cross Hospital. Of these 119 had migraine fulfilling Vahquist's criteria, 60 with classical and 59 with common migraine. Seventy-four patients (14 men and 60 women) reported that headaches were definitely precipitated by cheese, chocolate or citrus fruit, and 45 (14 men and 31 women) that headaches could not be precipitated by such foods. A disproportionate number of the "dietary" patients were selected for inclusion in this survey. 29 patients (24 men and 5 women) with cluster headache were included as a control group. In a second series of experiments 60 normal subjects (30 men and 30 women) drawn at random from a population survey, were compared with a random selection of patients from the dietary and non-dietary migraine groups and with all the subjects in the cluster headache group.

IgE measurements

Total serum IgE was determined by conventional radio-immunoassay (RIA) using the fast double-antibody separation technique and results expressed in international units per ml. Phadebas RAST (Pharmacia GB Ltd, Hounslow, England) was used to quantify circulating IgE antibodies against three common UK inhalant allergens and four foods—Timothy grass pollen (Phleum pratense), house-
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dust mite (*Dermatophagoides pteronyssinus*), cat epithelium, cow's milk, cheese, cocoa-bean and orange. The manufacturer's assay procedure was modified by increasing the first incubation from three hours to overnight and the patients' serum volume from 50 μl to 100 μl: this resulted in a two-fold increase in assay sensitivity and a relative decrease in non-specific binding to allergen-discs. Non-specific binding varied from 0-6% to 1-3% according to allergen and was allowed for in the calculations. Results were expressed as the percentage of a weakly-positive serum (Phadebas RAST “D” reference serum); 50-199% being interpreted as “weakly-positive” and over 200% as strongly positive.

IgG4 measurements

Immunosorbent-purified anti-IgG4 (Dutch Red Cross, Amsterdam), labelled with iodine-125 by the chloramine T method to a specific radioactivity of approximately 15 μCi/μg, was used to quantify both total and specific IgG4 antibody. Briefly, total and serum IgG4 was determined by a one-hour RIA in which a 50 μl IgG4 sample competed with 100 μl radio-labelled anti-IgG4 for binding to 100 μl of CNBr-activated microcrystalline cellulose-IgG4 particles.

The assay was controlled with milk-positive reference serum which bound 15% of 125I-anti-IgG4 in the assay, and hypogammaglobulinaemic serum which bound 0-8-1-2% of the tracer, depending on the allergen tested. Results were expressed in arbitrary units IgG4/ml: for example, 100 units IgG4/ml refers to a test sample which bound as much tracer as the reference serum. A figure of 15 units IgG4/ml was associated with a count rate approximately three times that associated with hypogammaglobulinaemic serum, and this was taken to be a positive result. Statistical analyses were carried out using Student's t test on log-transformed data and the null hypothesis rejected when p < 0-05.

Results

Total IgE—Geometric mean values and 95% confidence intervals for the dietary and non-dietary migraine patients and those with cluster headache are shown in fig 1: there was no significant difference between the two groups of migraine patients, but those with cluster headache were significantly higher (p < 0-01) than those of all the migraine patients taken together. There were no significant differences between 23 patients whose cluster headache was active and the remaining six whose headaches were quiescent at the time of sampling. Smoking data was available in 19 male cluster headache subjects—16 subjects smoked more than five cigarettes daily with a geometrical mean IgE titre of 65-9 units; only three subjects smoked less than five cigarettes daily, their mean IgE level was 16-8 units.

IgE to specific allergens (fig 2)

Responses to grass, mite and cat antigens were normal (averaging 20% of the subjects positive) and there were no differences between the three patient
antigens, but were more frequently strongly positive to the foods tested, but this was to an equal extent in the dietary and non-dietary migraine patients (fig 4). The cluster headache group, in contrast, had lower titres (p < 0.05) to dietary antigens and *D. pteronyssinus* than the migraine groups. The significance of this finding was explored by repeating the assay for cheese-related IgG4 in all the cluster headache patients, a random selection of those with migraine and a group of control subjects (fig 5). Detectable antibody was found in a minority of all the patient groups except those with cluster headache, which appear to have a very narrow distribution of antibody titres.

**Discussion**

Our finding that there are no differences in total or food-specific IgE levels between our dietary and non-dietary subjects makes us unable to confirm the suggestion of Monro et al. that 75% of severe migraine patients have raised levels of food-specific IgE antibodies. Our modification of the manufacturers' IgE-RAST procedure increased the percentage binding to the reference serum two-fold, thus lowering the limit for reliable detection of specific IgE. The variability of non-specific binding allowed for (1-3% for cocoa bean) was of the same order of magnitude as the results Monro et al. interpreted as positive—it is likely that we would have interpreted at least some of them as negative. We found no differences in total IgG4 in any of our subject groups. Although occasional high titres to food-related IgG4 antibodies were encountered in migraine patients, they were not
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![Graph showing cheese-specific IgG4-RAST results](image)

**Fig 5** Scattergram for cheese-specific IgG4-RAST results in which each patient group is compared to the normal group which is divided according to gender.

Differently distributed than the dietary and non-dietary patients and we found no evidence to suggest they are relevant to the pathogenesis of headache. In a recent community survey we found that 63% of women had raised IgG4 levels to egg and 40% to milk, with less than 3% having raised levels to grass, mite or cat allergens. Our findings in the cluster headache patients were unexpected. There is no clinical or epidemiological evidence of food-precipitated headache in this disease, though most patients can precipitate headaches by alcohol. Total IgE levels were significantly higher than those of either the migraine patients or controls, even when the study is confined to male subjects, but specific inhalant or food related antibody titres were not elevated. The geometric mean IgE levels in the cluster headache group (62.4 u/ml) was similar to that reached in non-atopic patients with respiratory disease where it was speculated that this may, at least part, be due to smoking habits, because men tend to have higher total IgE levels than women and usually are the heavier smokers. The explanation for the high IgE levels in our cluster headache patients may indeed lie in the disproportionate number of smokers; whether the headache is a cause or a consequence of the excessive smoking remains uncertain.

In contrast total IgG4 levels were within normal limits, but occasional patients with elevated titres to these foods, common in all other populations studied, were found.

The clinical significance of these immunological disturbances is uncertain but may be related to the excess of mast cells found in the skin in the area of the headache with degranulation during the active phase and also the favourable response often seen to corticosteroids.

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**References**

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