An unsuspected case of wheat induced asthma

ALAN J WILLIAMS, SUSAN E CHURCH, RONALD FINN

From the Royal Liverpool Hospital, Liverpool

The aetiology of asthma is still unknown, but there are well recognised trigger factors. The diagnosis of occupational asthma is rewarding; removing the patient from the work environment leads to an improvement in their asthma in most cases. In patients with inhalant asthma from the house dust mite prolonged allergen avoidance has been shown to improve symptoms. The role of food in asthma, however, remains undefined. Patients with immediate and severe asthma symptoms after ingesting a specific food are likely to recognise the problem easily and thus avoid the food in the future. Difficulty arises if the provoked asthma reaction is delayed after ingestion of the food (“late reaction”). In these circumstances the patient is less likely to notice the relationship, especially if the food is ingested every day. The following case history documents this problem in a patient whose asthma was finally recognised to be wheat induced.

Case history

A 60 year old woman was referred with a six year history of breathlessness (MRC grade 3), chronic wheeze with nocturnal exacerbations, and a non-productive cough. She was a non-smoker and had no history of eczema or seasonal rhinitis. Her treatment consisted of daily inhaled bronchodilators (salbutamol 1200 μg/day, beclomethasone dipropionate 500 μg/day) by pressurised aerosol. Her inhaler technique was faultless.

Investigations showed air flow limitation with a peak flow (PEF) of 300 l min⁻¹ (69% of predicted normal), but greater than 15% reversibility after inhaled salbutamol. A diagnosis of asthma was made. On careful inquiry the patient volunteered that her symptoms may have been aggravated by foods containing wheat, since she had noticed an improvement in her asthma after starting a weight reducing diet.

She was therefore given instruction to exclude wheat products from her diet, and when she was reviewed three months later her PEF had risen to 400 l min⁻¹ and she had of her own volition stopped all treatment.

Three weeks later she was electively admitted to hospital while continuing her wheat free diet. Skinprick tests (prepared by Bencard) for 24 common inhaled or food allergens showed positive (weal at least 3 mm greater than control) reactions to 14; the maximal weal diameter was 5 mm for grass pollens, with a 4 mm weal recorded for the following foods: potato, wheat grain, flour, egg, milk, cheese, chocolate, mixed beans, cod, lobster, nuts, and coffee. The total serum IgE concentration was raised at 172 U/l (normal range up to 100); but radioallergosorbent tests (RAST) for specific IgE for the grass pollen, house dust mite, cat fur, mould, egg white, and wheat gave negative results (RAST score 0 on the 0–4 scale). When a solid phase enzyme linked immunosorbent assay (ELISA) was used for detecting serum food antibodies (IgG) to gliadin, ovalbumin, and milk, antibodies to ovalbumin and milk were found in her serum. Serum IgG, IgA, and IgM, concentrations were normal, as was serum complement (C3 and C4).

On admission to hospital she was put on a “normal” wheat containing diet (150 g wheat/day) for six days. When she started this her peak flow fell linearly to a nadir of 250 l min⁻¹ (fig 1). At this time she began wheezing and needed to start treatment with salbutamol. During this period her FEV₁ fell from 2.1 (predicted 2.2 l) to 1.75 and her forced vital capacity (FVC) fell from 2.8 (predicted 3.0 l) to 2.2 l. The wheat diet was discontinued and she was started on an oligoallergenic diet of lamb, pears, and water only. After this her peak flow rate steadily rose to baseline levels within four days, when she was then rechallenged with dietary wheat with the same effect (fig 1).

After the PEF had returned to normal she was given separate double blind challenges with 30 g/day of wheat or potato placebo disguised as a soup. Once again with dietary wheat there was a steady fall in PEF comparable to the results of the open challenge (fig 2). A return to baseline treatment was continued with elimination of wheat from her diet.

Address for reprint requests: Dr R Finn, Royal Liverpool Hospital, Liverpool L7 8XP.

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Fig 1  Daily peak flow recordings after the introduction of a normal wheat containing diet and the subsequent response to an exclusion diet.
peak flow (prechallenge) was attained eight days after cessation of the wheat challenge. During wheat challenge total blood eosinophil count rose from 0-6 to 78 × 10^9/l, but there was no change in concentration total of any of the serum immunoglobulins or complement. The result of placebo challenge is shown in figure 2.

Discussion

Wheat has been known for centuries to induce wheezing in some people with asthma.4 4 Although asthma is probably one of the most prominent symptoms in patients with food intolerance, wheat induced reactions are found infrequently.5 For instance, in children clinical wheat sensitivity in asthma has varied in the reported series from 0%6 to 16%7 and seems likely to be of the order of 5%.8

In adults, no figures can be given since there have been no formal prevalence studies. Thus in the general population the frequency of wheat as a cause of asthma is not known, although it is probably relevant in only a small percentage of patients. Nevertheless, for those affected individuals it would be important to make the diagnosis since dietary wheat exclusion, as in this case, may lead to improved quality of life and less dependence on treatment. In our case the only clue to the possibility that wheat was important came from the patient's own observation that improvement had occurred during a period of dietary restriction. In past studies skinprick and RAST IgE tests performed to identify specific foods responsible for asthma have proved disappointing.5 9 In this case a positive skin reaction to wheat was only one of several positive reactions to foods, which were indistinguishable by weal diameter. The RAST test for specific serum IgE for wheat gave a negative result, as did a test for serum IgG antibodies to gliadin and we cannot therefore support a true "food allergic" or immunological mechanism. Circulating immune complexes of IgE or IgG with wheat, however, have not been sought and an immune mechanism still seems possible.10

References

3 Floyer Sir J. A treatise on the asthma. 3rd ed. London: Wilkins and Innes, 1726.

Notices

British Lung Foundation grants

The British Lung Foundation invites applications for grants of up to £30,000. Grants may cover the cost of personnel or equipment. Application forms can be obtained from the British Lung Foundation, 12a Onslow Gardens, London SW7 3AP. Completed application forms must be submitted by Monday 6 April 1987.

Chicago Lung Conference on Asthma

The Chicago Lung Association and Chicago Thoracic Society invite submission of papers for the 1987 Chicago Lung Conference on Asthma to be held October 8 and 9 1987. Topics and chairmen are: pharmacology (L Diamond), late asthmatic response (G Larson), neurohumoral interactions (A Leff), thermally induced asthma (R McFadden), pathogenesis of bronchial hyperreactivity (C Murlas), cell to cell interaction in airways (J Nadal), muscle physiology and protein biochemistry (N Stephens), airway responses in animal models (A Wanner). The programme will consist of state of the art presentations by the chairmen. Each lecture will be followed by 10 minute abstract presentations with five minutes discussions. The conference will emphasise basic and clinical research. The audience is limited to about 150 people. Further details from Dr Christopher Garrard (chairman of program committee), Chicago Lung Association, 1440 West Washington Boulevard, Chicago, Illinois 60607-1878, USA.
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A J Williams, S E Church and R Finn

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