Changing infant feeding practices and declining incidence of coeliac disease in West Somerset

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Abstract
An association was investigated between changing infant feeding practices and a declining incidence of childhood coeliac disease and transient gluten intolerance (TGI) in West Somerset, England during 1971–92. Dietary histories of 18 patients with coeliac disease were compared with 23 controls during 1971–80 and eight patients with coeliac disease and 39 controls during 1981–92. Our findings showed that the declining incidence of coeliac disease and TGI were associated with changing infant feeding practices, characterised by the later introduction of dietary gluten, an increased use of baby rice and gluten free foods for weaning, and an increased incidence of initial breast feeding.

(KeyEvent: coeliac disease; declining incidence; infant feeding)

The incidence of coeliac disease in childhood has declined in England, Scotland, and Ireland over the past two decades. However, in Italy, the incidence has been unchanged, and probably in Finland, where patients increasingly present with mild or atypical symptoms and signs of coeliac disease, at school age or during adolescence. In Sweden, the incidence of coeliac disease has increased since 1982. The reasons for these conflicting trends are unclear, with genetic and environmental factors being implicated, including the age of introducing dietary gluten in infancy. In this study we have investigated the influence of changing infant feeding practices on the incidence of childhood coeliac disease and transient gluten intolerance (TGI) in West Somerset, during 1971–92.

Patients and methods
West Somerset had a mean annual population of 263 000 (51 600 <14 years) during 1971–92 and a mean number of live births of 2700 per annum.

During this period, 39 children with a history of coeliac disease (22 boys, 17 girls; median age = 16 months, range = 3 months–13 years 8 months) were admitted to the children’s unit, Taunton and Somerset Hospital for further investigation. Peroral small intestinal biopsy specimens showed villous flattening by light microscopy, and they responded clinically to treatment with a gluten free diet. An initial diagnosis of coeliac disease was made and later confirmed on 16 patients (six boys, 10 girls), according to criteria of the European Society for Paediatric Gastroenterology and Nutrition (ESPGAN). A further 10 patients (five boys, five girls) were diagnosed with coeliac disease according to revised ESPGAN criteria. The age of presentation of coeliac disease in all patients was the time of their initial biopsy and the cumulative incidence of coeliac disease was the number of patients per 1000 live births, calculated for each birth cohort.

Dietary histories during infancy were obtained on patients and controls by a paediatric dietician. The numbers of infants initially breast or bottle fed, the age of introduction of dietary gluten, and the types of solids used for weaning were determined. Controls were children admitted to hospital in whom coeliac disease was excluded and 23 were studied between 1971–80 (16 boys, seven girls; median age = 15 months) and 39 between 1981–92 (19 boys, 20 girls; median age = 22 months).

Histological sections of biopsy specimens from patients were examined by light microscopy and graded by experienced observers (DNC, IKM) as follows:

Grade 1—villi of normal height, normal epithelial cell appearances, and a normally cellular lamina propria.

Grade 2—villous flattening varying from mild to moderate, loss of epithelial cell height, and increased cellularity of the lamina propria.

Grade 3—a flat mucosal surface with a cuboidal epithelium and heavy cellular infiltration of the lamina propria.

Results
COELIAC DISEASE
Of 39 patients first diagnosed with coeliac disease (22 boys, 17 girls), 29 had grade 3 histopathological changes by light microscopy and 10 had grade 2 changes. After two years on a gluten free diet, 29 (17 boys, 12 girls) had biopsy specimens taken from them again and had a normal mucosa. They were challenged with gluten and 16 (six boys, 10 girls) had a positive challenge with grade 3 changes and 13 (11 boys, two girls) a negative challenge with grade 1 changes. Ten patients (five boys, five girls) were diagnosed with coeliac disease after a single biopsy showing grade 3 changes, without a gluten challenge.

Altogether 26 children with coeliac disease were diagnosed between 1971–92 (11 boys, 15 girls; median age = 15 months; range = 4 months–165 months (13 years 9 months)); 18 during 1971–80 (median age = 13 months; range = 4 months–156 months (13 years)), and eight during 1981–92 (median age = 21 months; range = 9 months–165 months). The annual incidence of coeliac disease peaked in...
and gluten was introduced at a median age of 3 months (range = 1–9 months). The incidence of TGI declined during the early 1970s and no patients were diagnosed after 1977.

### Discussion

Early estimates of the incidence of childhood coeliac disease in the UK were based on diagnoses using clinical criteria. Since 1970, however, the diagnosis has been based on small bowel biopsy according to criteria agreed by ESPGAN, and the incidence has varied from 1:500 to 1:3000, with a mode of around 1:1000 live births.

In this study the incidence of coeliac disease in West Somerset declined from 1:1228 during 1971–80 to 1:4168 during 1981–92. A declining incidence was also reported from the Edinburgh and Lothian region of Scotland after 1976, after a more than twofold increased incidence.
Figure 3 Weaning solids first given to children with coeliac disease and controls (%).

from 1960–75. These findings could have represented a true reduction in the incidence of coeliac disease or a declining incidence from a high level in the late 1960s and early 1970s, possibly due to changing infant feeding practices. Studies of records from the Royal Hospital for Sick Children, Glasgow showed that increased numbers of children with coeliac disease were admitted during 1958–65 and 1966–73 than during 1950–57 and that patients were diagnosed earlier in the two later periods. Cereals containing gluten were also introduced earlier in Glasgow during 1958–63, and 91% of infants were given gluten by three months of age in 1965. The early introduction of gluten in infancy could also have contributed to the high incidence of coeliac disease in the West of Ireland during 1960–70, and to a reduction in the mean age of presentation of coeliac disease in England during 1964–72.

In 1974, a report from the Department of Health and Social Security (DHSS) Working Party of the Panel on Child Nutrition, encouraged breast feeding and discouraged the early introduction of cereals or other solid foods into the diet of infants before 4 months of age and the addition of cereals to bottle feeds. These recommendations were effected in West Somerset in the early 1970s and the median age of introducing gluten to patients with coeliac disease increased from 3 months during 1971–80 to 5.5 months during 1981–92. The median age of presentation of coeliac disease also increased during these periods, from 13 to 21 months. These dietary changes could have caused the declining incidence of coeliac disease in susceptible infants as well as the declining incidence of TGI. Another contributory factor could have been the increased availability of commercial gluten-free weaning foods, which rose from 72 in 1970 to 205 in 1989/90 (Coeliac Society (UK) handbook). Other infant feeding surveys also confirmed that infants were being weaned onto solids six weeks later in 1980 than in 1975, but weaning practices were relatively unchanged between 1980–90.

An increased incidence of initial breast feeding in children with coeliac disease and controls in West Somerset during 1971–92 and in other infant feeding surveys, could have contributed to the declining incidence of coeliac disease and TGI.

Breast fed children with coeliac disease and controls in West Somerset were given dietary gluten later than bottle fed children with coeliac disease and controls, and breast fed children with coeliac disease presented later than bottle fed children with coeliac disease. In another study the number of infants initially bottle fed with cows’ milk formulas during 1971–80 decreased during 1981–92. An increase in initial breast feeding may have protected the small intestinal mucosa against ‘priming’ by gluten, and possibly by other dietary antigens as well.

In Sweden, a doubling of the gluten content of proprietary baby foods was associated with an increased incidence of coeliac disease, from 1.7 children per 1000 live births during 1978–82 to 3.5 per 1000 thereafter. This occurred in spite of an increased prevalence and duration of breast feeding, and delay in the introduction of dietary gluten from 4 to 6 months. The clinical response to gluten in patients with coeliac disease is known to be dose dependent, and an overall reduction in the amount of gluten given during infancy could have been another factor contributing to the declining incidence of coeliac disease in West Somerset.

In this study, the findings suggested that changing infant feeding practices contributed to the declining incidence of coeliac disease during 1971–92 and to the disappearance of TGI. The dietary findings supported the recommendations made by the DHSS Working Party of the Panel on Child Nutrition in 1974. Also that baby rice and gluten free foods should initially be given to wean infants onto solids and that dietary gluten should be avoided until at least 5 months of age.

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References