Fine and ultrafine particles of the diet: influence on the mucosal immune response and association with Crohn’s disease

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Crohn’s disease is a modern Western disease characterised by transmural inflammation of the gastrointestinal tract. It is of unknown aetiology, but evidence suggests that it results from a combination of genetic predisposition and environmental factors. Bacterial-sized microparticles (0.1–1.0 µm) are potent adjuvants in model antigen-mediated immune responses and are increasingly associated with disease. Microparticles of TiO₂ and aluminosilicate accumulate in macrophages of human gut-associated lymphoid tissue where the earliest signs of lesions in Crohn’s disease are observed. Dietary microparticles are of endogenous or exogenous origin. Endogenous microparticles dominate and are calcium phosphate (most probably hydroxyapatite), which precipitates in the lumen of the mid-distal gastrointestinal tract due to secretion of Ca and phosphate in the succus entericus. Exogenous dietary microparticles are contaminants (soil and/or dust) and food additives. TiO₂, for example, is a food colourant, and aluminosilicates are anticaking agents, although some aluminosilicates occur as natural contaminants. Food additives alone account for ingestion of approximately 10¹² particles/person per d. Possible mechanisms for the role of exogenous and endogenous dietary microparticles in promoting tolerogenic or immune responses of gastrointestinal mucosal phagocytosis are discussed. In a double-blind randomised pilot study we have shown that a diet low in Ca and exogenous microparticles appears to alleviate the symptoms of ileal Crohn’s disease, with a significant (P = 0.002) improvement in the Crohn’s disease activity index. A multi-centre trial and further mechanistic studies at the cellular level are underway.

Dietary microparticles: Crohn’s disease

Fine (0.1–1.0 µm diameter) and ultrafine (<0.1 µm diameter) particles of the diet, collectively termed microparticles, are increasingly associated with the modern Western lifestyle. The present review considers dietary sources of microparticles, their cellular uptake and effects, and their possible association with the inflammatory bowel disease, Crohn’s disease.

Crohn’s disease

Crohn’s disease is a chronic relapsing inflammatory bowel disease that is characterised by transmural inflammation anywhere in the gastrointestinal tract, but typically in the distal small intestine (ileum), and/or caecum and large intestine (Table 1). It is often characterised by ‘skip lesions’, i.e. segments of bowel are affected with normal areas in between. The disease is commonly treated with drugs (e.g. azathioprine, corticosteroids), or liquid polymeric or elemental diets. Surgical removal of diseased bowel is necessary when management has failed, but the disease often recurs, and is then almost always proximal to the anastomosis.

The severity of the disease varies, but it commonly presents with periods of active disease and then remission, although any exacerbating factors are not clear. Both genetic (Hampe et al. 2001; Hugot et al. 2001; Ogura et al. 2001) and environmental factors (Fiocchi, 1998) contribute to the disease. The latter is clearly important since Crohn’s disease is almost exclusively a disease of developed countries, but presents also in immigrants from under-
developed countries. Crohn’s disease has increased in incidence, especially since the Second World War, and the prevalence is presently about 1:1000 of the population in the Western world (Calkins & Mendelhoff, 1995).

The first histological signs of disease overlie intestinal lymphoid aggregates (Peyer’s patches) and are commonly attributed to the translocation of bacteria and/or macromolecules across the M-cell-rich mucosa (Neutra, 1998), leading to abnormal inflammatory responses. Such observations certainly explain the presence of skip lesions in Crohn’s disease and the predominance of distal disease. In addition, these data suggest that macrophages, which phagocytose macromolecules, microparticles and bacteria, may be the primary cell type that is involved; a view that has received much recent support from cellular (Powell et al. 2000; SM Evans and JJ Powell, unpublished results), immunocytochemical (Ellis et al. 1998) and molecular (Hampe et al. 2001; Hugot et al. 2001; Ogura et al. 2001) studies. Recent molecular and genetic studies (Hampe et al. 2001; Hugot et al. 2001; Ogura et al. 2001) implicate a mutated NOD2 protein in disease susceptibility, this protein being an Apaf-1-like molecule with both nucleotide-binding and caspase-recruitment domains, which is exclusively expressed in monocytes. However, environmental factors that trigger and exacerbate disease have not been identified.

Shepherd et al. (1987) presented their findings of ‘pigment cells’ at the base of Peyer’s patches (Fig. 1) in all healthy and diseased samples of human small intestine. Subsequent analysis indicated that these cells contained Ti, Al and Si (Shepherd et al. 1987). In a follow-up study we showed that these elements were present as inorganic microparticles, i.e. anatase (a specific polymorph of TiO2), kaolinite (a polymorph of aluminosilicate) and other silicates (Powell et al. 1996). These microparticles are of dietary origin, accumulate at the base of Peyer’s patches and have been implicated elsewhere in inflammatory disease in susceptible individuals (see p. 125). For these reasons, we have investigated a possible association between inorganic microparticles of the diet and Crohn’s disease. More recently, we have widened these investigations to include endogenous lumen microparticles (calcium phosphate) that are degradable and are not accumulated, but still may transiently and frequently interact with mucosal phagocytes.

### Table 1. Features of Crohn’s disease

<table>
<thead>
<tr>
<th>Features</th>
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<tbody>
<tr>
<td>Peaks in young adults</td>
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<tr>
<td>Rare in developing countries</td>
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<tr>
<td>Increasing in immigrants to UK</td>
</tr>
<tr>
<td>Skip lesions</td>
</tr>
<tr>
<td>Predilection for ileum and/or colon</td>
</tr>
<tr>
<td>Treatment</td>
</tr>
<tr>
<td>Drugs</td>
</tr>
<tr>
<td>Diet</td>
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<td>Surgery</td>
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**Fig. 1.** High-powered light micrograph (×400 original) showing the base of a human intestinal lymphoid aggregate. The dark pigmented cells (P) represent CD68+ macrophages containing numerous granular microparticles of predominantly aluminosilicates and titanium dioxide.

**Microparticle-related diseases**

It has become clear that microparticles are associated with increased mortality and morbidity (Samet et al. 2000). Data commonly refer to organic atmospheric particles, such as diesel exhaust, and there are less clear data for inorganic particles, such as aluminosilicates and TiO2, except in geochemically-related diseases, Exposure to freshly-fractured silica is associated with coal miners’ pneumoconiosis (Liou et al. 1996), while the aluminosilicate, erionite, causes mesothelioma (Metintas et al. 1999). More interestingly, podoconiosis, which is an obstructive lymphopathy also known as non-filarial elephantiasis, is caused by soil microparticles taken up into the lymphatics of bare-footed agriculturists in tropical Africa (Price, 1990; Harvey et al. 1996). These biologically-active soil microparticles are the result of neovolcanic action, and those taken up into the lymphatics are rich in Al, Si and Ti. Only susceptible individuals go on to develop a latent granulomatous inflammatory disease of the lower limbs, whilst most individuals remain disease free. The potential analogy with microparticles and Crohn’s disease of the intestine is interesting.

The pattern of increasing dietary exposure to TiO2 in the Western world (Barksdale, 1966) parallels the prevalence of Crohn’s disease, and this relationship has stimulated us to focus on intestinal responses to this particle type (Powell et al. 2000). Although TiO2 is generally considered biologically inert, there are examples of inflammatory reactions to the TiO2 layer that forms on the surface of Ti metal hip prostheses (Kohilas et al. 1999). The surface layer (TiO2) migrates and is taken up by the local cellular environment, possibly with adsorbed self-proteins, thus inducing local immune responses (Ellingsen, 1991). The general evidence of lack of responsiveness to TiO2 in animal
studies should not be confused with possible abnormal immune responsiveness in genetically-diverse human subjects.

**Microparticles in the gastrointestinal lumen**

Microparticles are bacterial-sized non-biological particles which fall into two categories in the gastrointestinal tract, i.e. those of exogenous or endogenous origin. Endogenous microparticles are formed by precipitation in situ in the gastro-intestinal lumen and appear only to be calcium phosphate (most probably hydroxyapatite). Precipitation of other metal ions, such as Fe and Al, is carefully prevented due to the secretion of low-molecular-weight ligands, proteins and especially mucins into the intestinal lumen (Powell et al. 1999a,b). In contrast, while Ca is maintained in a soluble form in the upper gastrointestinal tract, it co-secretions with phosphate in the mid-distal gastro-intestinal tract forming calcium phosphate microparticles. This process is not a result of non-absorbed dietary Ca, but chiefly due to re-secretion of Ca and phosphate in the succus entericus of the gastrointestinal mucosa. In contrast, exogenous microparticles are of direct dietary origin and are mainly aluminosilicates and TiO₂.

**Sources of endogenous microparticles (calcium phosphate)**

The gastrointestinal homeostasis of Ca is complicated. In healthy individuals, net absorption occurs in the proximal small intestine, while net secretion occurs in the mid and distal small intestine, with variable flux in the colon. Ca secretion therefore occurs primarily in the jejunum and ileum, and is a non-saturable process that mostly follows a Na-dependent paracellular pathway (Favus, 1985). Since water and electrolyte fluxes greatly influence the secretion of Ca (Phillips & Giller, 1973; Powell et al. 1999), there are large ranges of lumen Ca concentrations both within and between individuals (Phillips & Giller, 1973; Powell et al. 1994). A median value of 4.2 mM-Ca in the proximal ileum is quoted (Geigy, 1968). In animals with normal Ca status the precipitate makes up the majority of the inorganic particulate content of the gastrointestinal lumen. This Ca species is acid soluble (Schedl et al. 1968), and would be re-dissolved in the acidic phagosomes of phagocytic mononuclear cells of the intestinal mucosa. This process leads to rapid cell death, probably apoptosis (SM Evans and JJ Powell, unpublished results), as elevated intracellular Ca is involved in a number of apoptotic pathways (Krebs, 1998; Tagliarino et al. 2001). It is possible, therefore, that calcium phosphate microparticles, carrying antigen on their surface or co-precipitated within their matrix, may induce immune tolerance rather than active immunity, as the phagocytic antigen-presenting cell undergoes cell death during antigen presentation.

A consequence of this induced apoptosis, however, appears to be the concomitant release of the pro-inflammatory cytokine interleukin (IL-1β) which, possibly, is essential for cell–cell signalling during the apoptotic process (Powell et al. 2000; SM Evans and JJ Powell, unpublished results). Surprisingly, our present data suggest that this common pathway of cell death and IL-1β release is not mediated through caspase 1 (IL-1 converting enzyme), since inhibition of caspase 1 activation inhibits IL-1β secretion but does not affect cell death (P Ashwood and JJ Powell, unpublished results). Clearly, other caspases are involved, and further investigations are underway. Nonetheless, the observation that particulate calcium phosphate induces IL-1β secretion in peripheral and intestinal mononuclear cells (SM Evans and JJ Powell, unpublished results) suggests that in the already-inflamed bowel, such as in Crohn’s disease, this process could add to the burden of pro-inflammatory cytokines.

**Sources of exogenous dietary microparticles**

Exogenous microparticles occur as natural dietary contaminants (typically dust and soil) and man-made food additive or pharmaceutical excipients. An average individual’s intake in the UK amounts to >10¹² microparticles/d (see p. 126), and these microparticles are resistant to gastrointestinal degradation. Furthermore, microparticles have large adsorptive negatively-charged surfaces that bind biomolecules of the gastrointestinal lumen. Once ingested, some microparticles are absorbed across the gastrointestinal mucosa and some accumulate in the macrophages (pigment cells) described earlier.
In the UK the daily dietary consumption of TiO$_2$ and aluminium sodium silicate (E554) is estimated as 5.4 and 0.5 mg/person per d respectively (Ministry of Agriculture, Fisheries and Food, 1993). TiO$_2$ added to food has a mean diameter of 0.2 µm, and thus the above-average daily intake translates to $10^{12}$ particles/person per d (Churg, 1996). Pharmaceutical and micronutrient supplements, which frequently contain TiO$_2$ and aluminosilicates as excipients, further increase the intake of microparticles, but estimates of this additional intake are not yet available.

**Titanium dioxide.** TiO$_2$ is an intense whitening and brightening agent used in the food and pharmaceutical industries (Table 2). In foods, it is primarily used to restore whiteness in creamy products (e.g. salad dressings) and in confectionery, where it sometimes forms a barrier between layers of different colours (e.g. soft-centred sweets with a crisp shell). Typical sources of TiO$_2$ are confectionery, white-coloured sauces and dressings, and non-dairy creamers. In the pharmaceutical industry TiO$_2$ is predominantly used as an opacity agent.

Quantitative evaluation of foods containing TiO$_2$ is difficult; first, because food labels do not have to provide quantitative data on food additives and second, because there are exceptions to the legislation that states food additives must be identified on food labels. For example, labelling is not required when additives perform no additive function in the final product and are used in compound ingredients that make up less than 25% of the finished product (Lomer et al. 2000). In addition, of course, many foods purchased as ready-to-eat do not carry food labels. TiO$_2$ (E171) can be used in specific food products at *quantum satis*, which indicates that there is no maximum level specified (Jukes, 1997). However, following good manufacturing practice, food manufacturers confirm that TiO$_2$ is used at levels no greater than that necessary to perform its function, and is generally used at $\leq 0.2$ % (w/w) of a product. Again, there are exceptions to this practice and, as illustrated in Table 2, use of this additive is highly variable, making accurate estimates of intake yet more difficult. Indeed, an individual’s dietary choices or preferences will dictate their intestinal exposure to TiO$_2$.

Exogenous microparticles and intestinal cells

Submicron-sized particles can be surprisingly well absorbed in the gastrointestinal tract, with notable uptake occurring at the M-cell-rich mucosa overlying intestinal lymphoid aggregates (Neutra, 1998). In addition, uptake by enterocytes and paracellular passage have been demonstrated, indicating that microparticle translocation may also occur across the regular intestinal epithelium (Desai et al. 1996). Although this process occurs more diffusely than at the surface of intestinal lymphoid aggregates, total uptake may be important due to the large surface area of the epithelium (Desai et al. 1996). Exogenous aluminosilicates (mostly kaolinite) and TiO$_2$ (anatase) are then passed to, and retained by, underlying phagocytes (pigment cells; see p. 124) of the mucosa. Both kaolinite and anatase are highly resistant to chemical degradation and are biologically inert, so neither of these microparticles appear to stimulate mononuclear cells *in vitro* or *in vivo*. However, the effects of microparticles on intestinal phagocytes should not be considered in isolation.

In contrast to most soil-derived microparticles, man-made microparticles (e.g. food additives) have active charged surfaces that avidly adsorb lumen biomolecules as they traverse the gastrointestinal tract (Govers et al. 1994). A ‘Trojan horse’ mechanism may operate whereby lumen toxins (e.g. bacterial lipopolysaccharide) or antigens (e.g. bacterial and food proteins) are transported into the intestinal mucosa on the surface of dietary microparticles. The effect of particulates on intestinal cellular responses to antigen has not yet been investigated, but data from cell lines suggest that antigen presentation may be markedly altered.

The processing of particle-bound antigen is qualitatively and quantitatively distinct from that of soluble antigen (Kovacsovics-Bankowski et al. 1993; Sedlik et al. 1997;)

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Table 2. The varying contents of titanium dioxide added to foods; examples for each group are shown for the minimum (min) and maximum (max) levels of titanium dioxide found in an analytical survey (Lomer et al. 2000)

<table>
<thead>
<tr>
<th>Food group</th>
<th>Food product*</th>
<th>TiO$_2$ (mg/portion)</th>
<th>Portion size (g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Confectionery</td>
<td>Mini Eggs (Cadbury Ltd; min)</td>
<td>0–55</td>
<td>15–30</td>
</tr>
<tr>
<td></td>
<td>Cake icing (Supercook; max)</td>
<td></td>
<td></td>
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<tr>
<td>Sauces</td>
<td>Mustard (Colman’s of Norwich; min)</td>
<td>0–225</td>
<td>30</td>
</tr>
<tr>
<td></td>
<td>Italian Dressing (Hellmann’s; max)</td>
<td>(&lt;1 tsp)</td>
<td>(1 tsp)</td>
</tr>
<tr>
<td>Non-dairy creamers</td>
<td>Coffeemate (Nestlé; min)</td>
<td>0–1–3.5</td>
<td>4.5</td>
</tr>
<tr>
<td></td>
<td>Teamate (Nestlé; max)</td>
<td>0–0.5</td>
<td></td>
</tr>
</tbody>
</table>

* tbs, tablespoon; tsp, teaspoon.

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1 Cadbury Ltd, Bourneville, Birmingham, UK; Supercook, Leeds, UK; Colman’s of Norwich, Norwich, Norfolk, UK; Hellmann’s, Unilever Bestfoods (UK) Ltd, Crawley, West Sussex, UK; Nestlé SA, Vevey, Switzerland.
Table 3. Typical food sources of aluminosilicates

<table>
<thead>
<tr>
<th>Aluminosilicates</th>
<th>E554, E556, E559</th>
</tr>
</thead>
<tbody>
<tr>
<td>Icing sugar</td>
<td></td>
</tr>
<tr>
<td>Savoury rice and noodles</td>
<td></td>
</tr>
<tr>
<td>Non-dairy creamers</td>
<td></td>
</tr>
<tr>
<td>Chewing gum</td>
<td></td>
</tr>
<tr>
<td>Powdered foods</td>
<td></td>
</tr>
<tr>
<td>Vending machine drinks</td>
<td></td>
</tr>
</tbody>
</table>

Song & Harding, 1996; Vidard et al. 1996). Particulate antigen, processed and presented by macrophages or B-cells, stimulates major histocompatibility complex (MHC) class II-restricted T-cell hybrids up to 10^5-fold more efficiently than soluble antigen (Vidard et al. 1996). Macrophages process a greater size range of particulate antigens compared with B-cells, but both these phagocytic antigen-presenting cells generate epitopes distinct from those from soluble antigen. Furthermore, Sedlik et al. (1997) showed that both T-helper 1 and 2 cytokines were produced in response to immunisation with particle-bound antigen, but in the presence of appropriate cofactors (e.g. IL-12) T-helper 1 cells were selectively activated (Sedlik et al. 1997). In addition, particulate antigen, unlike soluble antigen, may gain entry to an alternative class I MHC pathway, most notably in macrophages, and a number of mechanisms have been proposed (Song & Harding, 1996). Following phagocytosis, solubilised antigen may escape the phagosome and enter conventional cytosolic class I MHC processing. Alternatively, a post-golgi combination of phagocytosed degraded antigen with class I MHC may occur either intravesicularly or at the cell surface following regeneration. Recent data support an intravesicular mechanism (Song & Harding, 1996), and although the trafficking of nascent MHC I is rare, conventionally-coupled class I MHC–peptide may undergo pH induced de-coupling during passage of the acidic trans-golgi network, allowing peptide exchange (Stryhn et al. 1996).

Fewer studies have specifically investigated the processing of particulate antigen in vivo, although Maloy et al (1994) have shown that oral ovalbumin, delivered in poly(lactide-co-glycolide) particles, induced ovalbumin-specific cytotoxic T-lymphocyte responses and generated intestinal immunoglobulin A (Maloy et al. 1994). These processes are summarised in Fig. 2 and, clearly, the influence of the physical form of antigen in dictating cellular uptake and processing in the gastrointestinal tract should be considered.

To date much of our work has considered the influence of microparticle–toxin conjugates, rather than microparticle–antigen conjugates, on intestinal phagocyte function. Bacterial lipopolysaccharide (LPS) is abundant in the lumen of the gastrointestinal tract, and is a potent stimulant of monocytes. Monocytes recruited to the intestinal mucosa, however, undergo phenotypic and functional changes during maturation and become LPS-tolerant macrophages. One potential way to overcome LPS tolerance may be its delivery to the cell in a particulate form (e.g. as whole bacteria or adsorbed to microparticle surfaces). Pro-inflammatory cytokines such as IL-1β, tumour necrosis factor α and IL-6 are secreted by intestinal phagocytes in response to particulate LPS. However, one confounding factor is that adsorption of LPS to the surface of TiO₂ is facilitated by a Ca²⁺ bridge and, again, 4 mM-Ca has been used in much of our work. Thus, in the mixed LPS–anatase–Ca system the relative contributions of particulate Ca and particulate LPS to cellular secretion of inflammatory cytokines remains to be established. Nonetheless, the data clearly indicate that food additive TiO₂ amplifies cellular responses to Ca and/or LPS.

Thus, investigations with bacterial LPS or antigens indicate that microparticles can act both quantitatively and qualitatively as adjuvants for cytokine responses and T-cell proliferative responses induced by macrophages. Whether there is an underlying dysfunction in Crohn’s disease cells to such particulate adjuvant effects remains to be established.

Dietary exclusion of microparticles in Crohn’s disease

There are no data on dietary microparticle intakes in health or disease. However, diet has a major effect on the intestinal environment (Russel et al. 1998), and perhaps the adoption of Western dietary habits plays a part in the aetiology or pathogenesis of Crohn’s disease. Indeed, the most consistent dietary finding relates to sugar consumption, and studies have found that patients with Crohn’s disease have a significantly higher intake in comparison with control subjects (P<0.005, Martini & Brandes, 1976; P<0.05, Thornton et al. 1985). A high intake of sugar is often associated with diets high in convenience foods and confectionery, and these foods often contain dietary microparticles. Recently, we have conducted a case–control dietary study to assess differences in dietary microparticle intake, and our data confirm increased exposure to refined carbohydrate in patients with Crohn’s disease compared with healthy controls. Whether this finding will reflect differences in dietary microparticle intake is being investigated (MCE Lomer and JJ Powell, unpublished results).

Elemental diets are effective in the treatment of active Crohn’s disease (Mansfield et al. 1995) and the maintenance of disease remission (Wilschanski et al. 1996), and our analyses show them also to be free of microparticles (MCE Lomer and JJ Powell, unpublished results). These diets are as efficacious as conventional drug therapy (O’Morain et al. 1984; Okada et al. 1990; Zöli et al. 1997), and in adolescents and children are used in preference to treatment with corticosteroids or immunosuppressants (Wilschanski et al. 1996). Dietary compliance is often a problem, as large volumes are required to meet nutritional requirements, and the feeds can be unpalatable. Once disease remission is achieved, usually within 2–8 weeks, re-introduction of normal food is a gradual process, using an ‘exclusion diet regimen’ which introduces a single food each day and excludes any food that may be associated with symptoms of abdominal pain and/or diarrhoea (Riordan et al. 1993). Generally, these re-introduction diets start with foods that are typically well tolerated in Crohn’s disease, are usually in an unprocessed form, and are thus low in microparticles.
To establish whether a reduction in lumen microparticles can contribute to disease remission a double-blind pilot dietary study was undertaken. Patients with active ileal Crohn’s disease, treated with corticosteroids, were randomised to receive dietary advice on either a diet low in microparticles (trial) or a diet unrestricted in microparticles (control) for 4 months (nine subjects per group completed the treatment; Lomer et al. 2001). Dietary advice for the low microparticle intake (trial group) included: (1) avoidance of the inorganic exogenous microparticles, TiO₂ and aluminosilicates; (2) limiting dietary Ca intake to 400 mg/d (lower reference nutrient intake; Department of Health, 1991); (3) meeting the dietary reference values for all other nutrients (Department of Health, 1991). The unrestricted microparticle diet (control group) were advised on how to meet their dietary reference values for all nutrients (Department of Health, 1991). Both groups were recommended to avoid fibrous fruit and vegetables, as indicated for patients with structuring disease (Ballegaard et al. 1997), and equal time was spent in advising all patients.

Fig. 3 summarises the data after 4 months’ treatment, when a significant ($P=0.002$) reduction in the Crohn’s disease activity index (a score derived from patient symptoms for the preceding 7 d and current clinical data; Best et al. 1976) was observed in the trial group by the end of the study, as compared with results in the control group (Lomer et al. 2001). The efficacy of this diet is now being assessed in a multi-centre randomised double-blind placebo-controlled trial, but the pilot study provides some early evidence for a role for lumen microparticles in disease exacerbation.

Indeed, diversion of the faecal stream can help maintain disease remission in patients with Crohn’s disease following surgical removal of the diseased bowel, while following re-anastomosis the disease recurs (Rutgeerts et al. 1991). As the pilot diet was low in TiO₂, aluminosilicates and Ca, the relative role of exogenous microparticles and...
calcium phosphate in exacerbating disease is unclear, but this factor is presently being addressed in the multi-centre study.

In conclusion, dietary microparticles occur in two distinct forms; exogenous microparticles are of dietary origin, are resistant to degradation, have little effect on intestinal cells per se, but can bind lumen biomolecules and act as adjuncts in their cellular stimulation. In contrast, endogenous Ca microparticles (most probably hydroxyapatite) precipitate in situ in the gastrointestinal lumen, are acid labile and broken down by phagocytes, where they can induce IL-1β release and cellular apoptosis. Whether such microparticles can contribute to the ongoing inflammation of Crohn’s disease, which is a modern Western inflammatory bowel disease driven by dysfunctional macrophages, remains to be established.

Acknowledgements

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References


Govers MJAP, Termont DSML, Aken GAV & Van der Meer R (1991) Dietary calcium phosphate in exacerbating disease is unclear, but can bind lumen biomolecules and act as adjuncts in their cellular stimulation. In contrast, endogenous Ca microparticles (most probably hydroxyapatite) precipitate in situ in the gastrointestinal lumen, are acid labile and broken down by phagocytes, where they can induce IL-1β release and cellular apoptosis. Whether such microparticles can contribute to the ongoing inflammation of Crohn’s disease, which is a modern Western inflammatory bowel disease driven by dysfunctional macrophages, remains to be established.


Song R & Harding CV (1996) Roles of proteasomes, transporter for antigen presentation (TAP), and β2-microglobulin in the processing of bacterial or particulate antigens via an alternative class I MHC processing pathway. *Journal of Immunology* **156**, 4182–4190.


