Does dietary fibre stimulate intestinal epithelial cell proliferation in germ free rats?

R A GOODLAD, B RATCLIFFE, J P FORDHAM, AND N A WRIGHT

From the Cancer Research Campaign Cell Proliferation Unit, Department of Histopathology, Royal Postgraduate Medical School, DuCane Road, London, and Polytechnic of North London, Holloway Road, London, and AFRC Institute of Food Research, Reading, Berks

SUMMARY The aim of the present experiment was to investigate the role of hind gut fermentation in the proliferative response of the intestinal epithelium to dietary fibre. We have previously shown that refeeding starved rats with an elemental diet supplemented with fermentable dietary fibre (but not inert bulk) is capable of stimulating intestinal epithelial cell proliferation throughout the gastrointestinal tract. Three groups of 10 germ free (GF) rats and three groups of 10 conventional (CV) rats, were used. All groups were starved for three days and then refed for two days with either an elemental diet (Flexical); Flexical plus 30% kaolin; or Flexical plus 30% of a fibre mixture. Cell production was determined by the accumulation of vincristine arrested metaphases in micro-dissected crypts. There was no significant difference between refeeding the rats with an elemental diet alone or with kaolin supplementation, however, the addition of fibre in CV rats was associated with a significant increase in intestinal crypt cell production rate in both the small intestine (p<0.01) and the colon (p<0.001). This marked proliferative effects of fibre was abolished in the GF rats. It can be concluded that it is the products of hind gut fermentation, not fibre per se that stimulate intestinal epithelial cell proliferation in the colon and small intestine.

The term ‘dietary fibre’ is generally used as a broad (but useful) description for any non-starch polysaccharide not degraded by mammalian endogenous secrections; thus encompassing the glucose chains of cellulose and non-cellulosic polysaccharides, soluble and insoluble simple sugars, hemicellulose, pectic substances and lignin. Fibre is broken down by the microflora of the hind gut of monogastric animals such as man to produce short chain fatty acids (SCFA’s), mainly in the form of acetic, propionic, and butyric acid, which can be metabolised by the colonic epithelium and by the liver, and may constitute about 10% of the daily energy intake, even in the fibre deficient ‘Western’ diet.

The intestinal epithelium is a very dynamic tissue which is capable of adapting to a wide variety of situations by adjusting the rates of cell production in its well defined reproductive zones. Feeding the low bulk chemically defined ‘elemental diets’ is associated with intetinal atrophy and decreased cell proliferation, especially in the distal gut. Although this has been attributed to the lack of non-absorbable ‘bulk’, it is more likely to be the consequence of the removal of fermentable fibre. Refeeding starved animals can potentiate the effects of different dietary components, and provides a very useful model for the study of intestinal adaptation. Refeeding starved rats with a fibre free elemental diet supplemented with fermentable fibre stimulates intestinal epithelial cell proliferation in the colon and in the small intestine, whilst the addition of inert bulk has no such effect. Fibre, but not bulk, also has a similar effect in mice fed continuously.

These proliferative effects could either be the result of the direct effects of fibre itself or to the products of its breakdown by the complex microbiological flora of the caecum and colon. If the latter was indeed the case one would expect that the proliferative effect on the intestinal epithelium would be abolished in GF animals. The present study describes the proliferative effects of refeeding GF rats, or rats with a CV intestinal flora, a fibre free elemental diet supplemented with kaolin or with a fermentable fibre mixture.
Methods

EXPERIMENTAL PLAN
Three groups of CV rats and three comparable groups of GF rats were starved for three days and then refed for two days. The first pair of groups were refed with 15 g/rat/day of Flexical (Mead Johnson, Slough), the second were refed with flexical plus 30% w/w kaolin and the third pair of groups were refed with Flexical plus 30% of a fibre mix. The fibre mix comprised of 10% of the dietary mucilage from the ispaghula husk (Reckitt & Coleman) and 90% of the wheat grain fibrous extract, Trifyba (Labaz Sanofi. Wythenshawe, Manchester, hemicellulose 40%, cellulose 20%, lignin 15%, and pectin 5%). The animals had free access to water at all times.

After two days of refeeding the rats were weighed, injected with vincristine and killed at timed intervals by ether anaesthesia followed by exsanguination. The intestines were removed, rinsed, blotted, weighed and one centimetre samples from defined sites were fixed in Carnoy’s fluid.

ANIMALS
The rats were the offspring of a colony of GF Lister Hooded or a colony of genetically similar CV animals. The mean weight at the beginning of the experiment was 200 g, with no significant difference between any of the groups. All rats were between five and six weeks old at the start of the experiment. The animals had been maintained on a standard laboratory diet (PRD, Spillers) post weaning. Individuals were randomly assigned to a particular dietary treatment within either GF or CV environments.

HOUSING
Both CV and GF rats were housed in stainless steel isolators of the type described by Gustafsson. Rats were in groups of 10 per isolator. The cages were stainless steel with wire grid floors and were maintained without bedding.

GERM FREE TECHNIQUES
All CV and GF rats were maintained in isolators which were supplied with filtered, sterile air. All diets were sterilised by irradiation at a level of 50 kGy from a 60Co source and water was autoclaved. The isolators containing the GF rats were maintained with appropriate sterile techniques; GF status was monitored by taking swabs from the isolators and examining them by the techniques described by Fuller.

CRYPT CELL PRODUCTION RATE
Rats were injected with vincristine sulphate (1 mg/kg intraperitoneally; Tillots Laboratories, Henlow, Bedfordshire, UK) at 0900 hours and killed at timed intervals 30 to 180 minutes later. Samples of the small intestine and colon (defined by their percentage length of the entire small intestine or colon) were fixed in Carnoy’s fluid and stored in 70% (v/v) ethanol. They were later stained with the Feulgen reaction. Intestinal and colonic crypts were displayed by microdissection and gently squashed with a coverslip. The number of arrested metaphases in 10 small intestinal crypts or 20 colonic crypts were counted and the mean values plotted against time since injection. The slope of the line was then fitted by linear regression to give the crypt cell production rate (CCPR) and its standard error.

STATISTICAL ANALYSIS
All results are presented as the mean (standard error) of the mean. Data were tested by a two-sided t test, or by two way analysis of variance where the data was classified by two factors, diet and microflora. In the latter case it is possible for one of the two factors to alter the effect of the other; this is measured by an interaction effect. Lines were fitted by least squares linear regression.

Results
The effects of the different diets and of intestinal flora on total body weight, intestinal length, and the wet weight of the main regions of the gastrointestinal tract are presented in the Table, which also summarises the results of two-way analysis of variance on the above data. The GF groups were significantly heavier than the CV rats, but these animals had the grossly distended fluid filled caeca associated with GF status which would have contributed to the increased body weight. There was a significant tendency for the body weight to be less in those groups fed the supplemented diets.

Two way analysis of variance showed a small effect of diet on stomach weight, which was less in the fibre supplemented group. No effect of microflora was noted. The small intestine, caecum and colon were all significantly heavier in the GF groups. The hypertrophy of the caecum was particularly pronounced. No significant effect of diet was noted in the small intestine and caecum, but a very highly significant effect was seen in the colon, where the colon weight increased in the fibre fed animals, especially the CV ones. A significant interaction between the effects of diets and microflora was observed. The colon length mirrored the colon weight changes.

The crypt cell production rate of the proximal small intestine of the CV group fed fibre was significantly increased when compared with the group fed the elemental diet alone or plus kaolin. No such effect was seen in the GF animals (Figure).
This proliferative effect of fibre in the CV rats was more pronounced in the distal than in the proximal small intestine, with the cell production rates of the fibre fed group being almost doubled (p<0.01) when compared with the unsupplemented group. The proliferative effect of fibre was even more pronounced in the colon of the CV rats where there was a sixfold increase in the CCPR (p=0.001). No such effect was seen in any of these sites in the GF groups.

**Discussion**

The results of this study confirm the proliferative effect of fibre on intestinal epithelial cell production in CV rats, and the lack of any proliferative response in the GF rats implies that it must be the fermentation of fibre in the hind gut that is the cause of this proliferation. The main product of the bacterial breakdown of fibre are the short chain fatty acids (SCFA's), and there is some evidence that these are indeed trophic to the intestine in vivo.6 14 21 22

The changes in crypt cell production noted in the small intestine and colon were very pronounced, which was reflected by the significant changes in intestinal weight observed, despite the short duration of refeeding. Two way analysis of variance showed the presence of significant changes in stomach weight with diet, which reflected a slight increase in tissue weight with fibre ingestion. All areas of the intestine were heavier in the germ free rats, and this led to a significant microflora effect in all areas studied except the stomach. The most dramatic effect of germ free status was the massive increase in caecal weight, which increased 2.4 times and even this still does not reflect the magnitude of the caecal volume increase; Gustafsson30 quotes a five-fold volume increase in the caecal contents of the GF rats (for a doubling of tissue weight) and a 30-fold increase in the luminal mucus content. Caecal enlargement is also seen in CV rats fed large amounts of non-absorbed large molecules, or in those fed antibiotics35 and this does not appear to depend on the luminal osmolarity.34

It is interesting that no effect of diet on the caecum was noted, whereas the colon length and weight both showed a highly significant effect of both diet and microflora. This increase in colon weight noted in the CV rats, although quite dramatic, however, was not nearly as pronounced as the increase in CCPR; nevertheless, if the experiment had continued for longer the weight change could perhaps have been equally pronounced. Alternatively the changes

<table>
<thead>
<tr>
<th>Measurement</th>
<th>Diet</th>
<th>Diet+ kaolin</th>
<th>Diet+ fibre</th>
<th>Diet</th>
<th>Microflora</th>
<th>Interaction</th>
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<tbody>
<tr>
<td>Body weight (g)</td>
<td>226.4-6 (6-9)</td>
<td>211.2-7 (7-7)</td>
<td>211.9-9 (7-0)</td>
<td></td>
<td></td>
<td>*</td>
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<tr>
<td>CV</td>
<td>246.8-5 (5-4)</td>
<td>231.3-3 (9-0)</td>
<td>221.2-7 (8-0)</td>
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<td>†</td>
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<tr>
<td>GF</td>
<td></td>
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<td></td>
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<tr>
<td>Weight stomach (g)</td>
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<td></td>
<td></td>
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</tr>
<tr>
<td>CV</td>
<td>1.183 (0.038)</td>
<td>1.113 (0.030)</td>
<td>1.108 (0.034)</td>
<td></td>
<td></td>
<td>*</td>
</tr>
<tr>
<td>GF</td>
<td>1.194 (0.030)</td>
<td>1.103 (0.030)</td>
<td>1.120 (0.030)</td>
<td></td>
<td></td>
<td>†</td>
</tr>
<tr>
<td>Length small intestine (cm)</td>
<td>1.046 (1.5)</td>
<td>1.031 (1.3)</td>
<td>9.66 (9.0)</td>
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<tr>
<td>CV</td>
<td>1.061 (1.7)</td>
<td>1.077 (1.6)</td>
<td>11.05 (7.8)</td>
<td></td>
<td></td>
<td>†</td>
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<td>Weight small intestine (g)</td>
<td>6.186 (0.253)</td>
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<tr>
<td>CV</td>
<td>6.624 (0.150)</td>
<td>6.513 (0.340)</td>
<td>6.675 (0.321)</td>
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<td>†</td>
</tr>
<tr>
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<td></td>
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<tr>
<td>Weight caecum (g)</td>
<td>0.810 (0.036)</td>
<td>0.726 (0.039)</td>
<td>0.714 (0.031)</td>
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<td>1.795 (0.038)</td>
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<td>†††</td>
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<tr>
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</tr>
<tr>
<td>Length colon (cm)</td>
<td>1.35 (0.4)</td>
<td>1.41 (0.2)</td>
<td>1.47 (0.3)</td>
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<td>†††</td>
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<tr>
<td>CV</td>
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<td>1.55 (0.2)</td>
<td>1.80 (0.5)</td>
<td></td>
<td></td>
<td>†††</td>
</tr>
<tr>
<td>GF</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Weight colon (g)</td>
<td>0.906 (0.030)</td>
<td>0.992 (0.049)</td>
<td>1.221 (0.060)</td>
<td></td>
<td></td>
<td>†††</td>
</tr>
<tr>
<td>CV</td>
<td>1.085 (0.031)</td>
<td>1.143 (0.040)</td>
<td>1.155 (0.051)</td>
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<td>†††</td>
</tr>
<tr>
<td>GF</td>
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</table>

*Significantly different (p<0.05); †Significantly different (p<0.01); ††Significantly different (p<0.001). Standard error of the means are shown in parentheses.
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Figure  Effects on crypt cell production (CCPR) of refeeding various diets to starved rats. Conventional (CV) or germ-free (GF) animals were refeed with an elemental diet, or with the same diet supplemented with either kaolin or a fermentable fibre mixture. The sites in the intestine were identified by their relative position in the gut. *Significantly greater than the respective group fed the elemental diet (p<0.05); **Significantly greater than the respective group fed the elemental diet (p<0.01); ***Significantly greater than the respective group fed the elemental diet (p<0.001).

observed may represent the peak of an adaptive response, and a lesser effect would be seen in a chronic system; however consistent longterm changes would still be expected.17

The proliferative effect is only one of the many biological effects of fibre. Poorly fermentable fibre may abrade the intestine and stimulate increased cell loss, as cellulose powder has been reported to decrease villus length in GF mice, whilst it increased lengthb and turnover rate27 in CV mice. The GF intestinal villi are usually enlarged, perhaps because of decreased cell loss.28 The use of villus length is not, however, the ideal estimator of intestinal morphology. Fibre may also dilute or bind carcinogens and alter bile acid and cholesterol levels. A further effect of fibre, especially when from the brassicas, is to stimulate the induction of xenobiotic metabolising enzymes.24 Other repercussions of fibre intake include delaying gastric emptying and consequently damping the glycaemic response, stool softening, reducing digesta transit times and increasing stool output. Some fibres, especially those rich in phytic acid, bind minerals and thus render them unavailable to the animal. The microbial breakdown of fermentable fibre leads to the production of SCFA’s which in turn will also have many effects; these include lowering luminal pH (which in turn increases SCFA uptake, as they are more readily absorbed in the unionised form), and increasing water absorption. In addition fibre can alter the intestinal flora, stimulate mucous production, increase vitamin K absorption, increase mucosal blood flow, increase motor activity,26 and alter urea and ammonia concentration.1

The extent of fibre fermentation in the colon, of monogastric animals, including man is only recently being appreciated.12 Although the metabolism of short chain fatty acids is not as important as that seen in fore-gut fermenters where over 70% of the daily energy intake may be absorbed in the form of SCFA’s (and the more fermentable the diet the greater the epithelial cell proliferation in the fore-gut),12 recent estimates suggest that even on the relatively low fibre diet of the Western world SCFA production contributes significantly to the daily energy intake in man.1 It should be borne in mind that SCFA production will occur even when no fibre is ingested, as intestinal mucus and other materials will still arrive at the hind gut, thus the SCFA concentration (but not production) in dogs fed only on meat is similar to that seen in hind gut fermenting herbivores.31

The weight of bacteria in the colon is equivalent to, or greater, than that of several major body organs, and there are more bacteria in the colon than there are cells in the human body.30 The bacterial content of the colon is around 10^{10}-10^{12} per ml whereas in the terminal ileum it is 10^2-10^4, therefore although there is some fermentation in the ileum,30 SCFA levels are very low, so that the proliferative response observed is unlikely to be a direct effect of SCFA production,
suggesting a systemic effect which may be direct or may be moderated via some other humoral agent. There is considerable evidence that many adaptive responses are strongly influenced by blood borne agents.11

It has been reported that SCFA infusion into the colon stimulates mitotic indices in the jejunum,12 but in vitro studies show no direct effect upon isolated intestinal tissue.13 We have previously shown a good correlation between plasma enteroglucagon and PYY in animals fed fibre mixtures of varying degrees of fermentability, and the changes in these hormones in GF rats will be described later.

The biological significance of this proliferative effect of fermentable fibre and SCFA’s is not as yet clear, as it is only one of the many effects of fibre now known. None the less, this effect warrants further study, especially in light of the observed relationship between hyperplasia and the promotion of carcinogenesis in experimental animals.14 Feeding carcinogen treated rats a high fibre diet has been shown to enhance colon carcinogenesis;15 16 17 nevertheless, other workers have found the converse.18 19 Although such models usually only study the end yield of a complex multistage system, they none the less sound a note of warning, particularly in light of the current advocacy of high fibre diets.

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References


