Effects of a new, concentrated wheat fibre preparation on intestinal transit, deoxycholic acid metabolism and the composition of bile

S N MARCUS AND K W HEATON

From the University Department of Medicine, Bristol Royal Infirmary, Bristol

SUMMARY When the cholesterol saturation index of bile is reduced by wheat bran there is generally a fall in the deoxycholic acid content of bile. As the same effects occur with senna, bran might act on bile simply via its accelerating effect on colonic transit. We have studied the effects of a new, concentrated, wheat fibre preparation (Testa Triticum Tricium, Trifyba, which is 80% dietary fibre) upon bile composition, deoxycholic acid metabolism and intestinal transit time, and have assessed whether these effects are related. Twenty constipated volunteers were prescribed Testa Triticum Tricium in doses (10–32 g/day) sufficient to relieve their symptoms for at least six weeks. Before and at the end of this period, duodenal bile was sampled to enable measurement of deoxycholic acid pool (by isotope dilution), total bile acid pool, bile acid composition and cholesterol saturation index. Whole gut transit time fell from 120±SD35 to 68±35 hours. At the same time, biliary %deoxycholic acid fell from 26·6±12·0 to 23·0±11·8 (p=0·002), the total bile acid pool expanded from 2·36±0·88 to 2·75±0·90 g (p=0·008) and cholesterol saturation index fell from 1·13±0·32 to 1·07±0·29 (p=0·04). In subjects with initial cholesterol saturation index over 1·0 (n=12), it fell from 1·33±0·25 to 1·22±0·21 (p=0·008). There was no significant correlation between change in saturation index and change in %deoxycholic acid or deoxycholic acid pool, nor between any of these parameters and change in transit time. Testa Triticum Tricium reduces the cholesterol saturation index of supersaturated bile but this action appears to be independent of its effect on colonic transit and of changes in deoxycholic acid metabolism.

Several studies have shown that the administration of wheat bran lowers the cholesterol saturation of gall bladder bile, at least when this is initially supersaturated.1–4 The mode of action of bran has not been established. Initially, it was suggested that it acted by reducing the level of deoxycholic acid in bile1 and in five studies reported between 1973 and 1978 bran did lower the proportion of deoxycholic acid in the bile acid pool.1–3 5 6 No such effect was found in two later studies.4 7 In one of them7 the subjects were young and slim and they began with low levels of deoxycholic acid and were already on a high fibre diet. It was not surprising, therefore, that there was no change in the bile acid composition or the cholesterol saturation of their bile. In the other study,4 however, there was a clear fall in cholesterol saturation despite no change in deoxycholic acid levels. In view of the conflicting evidence, it seemed appropriate to examine in more detail the effects of wheat bran or, at least, of wheat fibre, upon deoxycholic acid metabolism, ensuring that some subjects had initially supersaturated bile.

If bran does act by reducing the deoxycholic acid content of bile this may be because of its laxative properties, such that, with faster transit of colonic contents, there is less time available for the formation or absorption of deoxycholic acid. We have previously shown that a chemical laxative, sennoside B, reduces both the deoxycholic acid pool and the cholesterol saturation of bile and that these changes correlate with each other and with changes in transit time.8 Lactulose has similar effects9 but the reduction in deoxycholic acid was attributed to acidification of the right colon.

The aim of the present study was to determine whether a new concentrated wheat fibre preparation was effective in lowering the cholesterol saturation index of bile and whether these effects were
mediated through alterations in deoxicholic acid metabolism and specifically in the size of the deoxicholic acid pool. The latter has not previously been measured in subjects with supersaturated bile before and after supplementation with wheat fibre. We also wished to look for correlations between wheat fibre induced changes in whole gut transit time and changes in bile composition and deoxicholic acid metabolism.

Methods

SUBJECTS AND DESIGN OF STUDY

Constipated subjects were recruited in response to posters in local chemist shops and health centres, in accident and clinic waiting areas of the Bristol Royal Infirmary and Dental Hospital, and in WRVS stalls. Advertisements were also inserted in the University Newsletter and health authority gazette and letters of appeal were sent to members of the hospital voluntary service, St John Ambulance Brigade and Red Cross Society.

The group consisted of 16 women and four men, aged 38–69 years (mean 49 years) with body mass or Quetelet index 18–33·6 (mean 24·7), who admitted to three bowel actions per week or less, or who often strained to pass scybalous faeces and whose whole gut transit time was 70 hours or longer. Volunteers admitting to alternating constipation and diarrhoea were excluded. Those who were already taking laxatives, bran or bran containing cereals were instructed to stop them for at least four weeks before entering the study. Ten subjects were taking other drugs (four diuretics, two antihypertensives, two antidepressants, one each temazepam, ibuprofen, non-cyclical progesterone and non-cyclical combined oestrogen and progesterone preparation) but the dosages remained unaltered during the study. Baseline measurements of transit time, bile composition and deoxicholic acid pool size and kinetics were carried out. Dietary intake was assessed by an experienced dietitian using a questionnaire and a 48 hour weighed record. The data obtained were analysed using a computer program compiled from standard food tables.

Subjects were then prescribed a concentrated wheat-fibre preparation, Testa Triticum Tricium (Trifyba-Labaz), containing 80% dietary fibre by weight, which is manufactured by treating wheat bran enzymatically to remove starch, protein, and phytic acid. Subjects were instructed to take one or more sachets, each containing 3·5 g of Testa Triticum Tricium, with meals, in a dose sufficient to relieve their constipation (10–32 g daily) for a minimum of six weeks. Adequate response to treatment was monitored by one of us (SNM) by regular telephone calls. At the end of this period the transit time, biliary studies and dietary assessment were repeated.

In six of the volunteers, baseline measurements while they were constipated were obtained several weeks before starting Testa Triticum Tricium. In the meantime they had been treated with sennoside B (15–45 mg daily) for at least six weeks and had been restudied. They were included because it was known or expected that their baseline bile was supersaturated with cholesterol, which had not been the case with most of the subjects up till that time. It was considered important to have an adequate group of subjects whose bile was usually supersaturated. Unfortunately, all but one of these subjects refused to undergo a ‘wash-out’ period without taking any form of laxative because they were unhappy about becoming constipated again. In order to ensure that, in these five subjects, the effects of senna laxative on the bile had worn off and that any changes in bile composition could be credited to Testa Triticum Tricium, the latter was prescribed for a minimum of 12 weeks before bile was collected for repeat analysis.

Subjects had blood taken for liver function tests and these were normal. Fasting lipids were also measured and these were normal or only slightly raised. Ultrasound scan of the gall bladder showed the absence of gall stones in all. Four subjects were excluded from the study, one because of gall stones, one because of non-compliance and two because of failed intubation. The protocol of the study was reviewed and accepted by the District Ethical Committee of the Bristol and Weston Health Authority.

WHOLE GUT TRANSIT-TIME

Transit time was measured by the single stool method with two modifications. While constipated, subjects ingested a gelatine capsule containing 20 radio-opaque Portex markers at breakfast time on alternate days for one week (rather than on three consecutive days) and then collected their next two stools. Each day the markers were of a different shape and were taken in a predetermined order. Stools were excreted directly into gusseted polyethylene bags which were sealed and stored at −20°C until radiographed. Whole gut transit time was calculated from each stool by counting the two most abundant markers and applying the formula:

\[
\text{Whole gut transit time (h)} = \frac{t_1 + t_2}{s_1 + s_2}
\]

where \(t_1\) and \(t_2\)=time in hours from the ingestion of the two markers and \(s_1\) and \(s_2\)=the number of each...
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BILE COMPOSITION AND THE POOL SIZE AND KINETICS OF DEOXYCHOLIC ACID

Three hours after the last meal of the day 0.37 MBq (10 μC) 24\(^{14}\)C deoxycholic acid, with a specific activity of 5.33 MBq/mg and radiochemical purity on thin layer chromatography of 96% (Radiochemical Centre, Amersham), was injected intravenously and, after an overnight fast, bile rich duodenal fluid was collected by means of duodenal intubation and cholecystokinin injection. The period of fasting was standardised for each volunteer.\(^{12, 13}\) In 11 volunteers 5 ml aliquots of duodenal bile were collected on four consecutive mornings to enable determination of deoxycholic acid pool size, input and fractional turnover.\(^{14}\) In the remaining nine subjects who were unwilling to have repeated intubations, deoxycholic acid pool was determined by a slightly modified version of the ‘one shot’ technique.\(^{15}\)

Total bile salt, phospholipid and cholesterol concentrations were measured\(^{16}\) and the cholesterol saturation index of bile was determined by the method of Thomas and Hofmann\(^{17}\) according to the criteria of Hegardt and Dam.\(^{18}\) In the small number of samples in which the total lipid concentration was <20 mmol/l phospholipid concentration was measured using an enzymatic method.\(^{19}\)

After enzymatic deconjugation and thin layer chromatographic separation of the dihydroxy bile acids\(^{20}\) deoxycholic acid mass was determined by subtracting the 7α- from the 3α-hydroxysteroid dehydrogenase results (Sigma Chemical Co.).\(^{6}\) The mass of the separated cholic acid was determined using 3α-hydroxysteroid dehydrogenase. Radioactivity was measured by liquid scintillation counting. The counts were corrected for any effects due to quenching using quench curves calculated from external standards of \(^{14}\)C.

After this period, deoxycholic acid was removed from the pool size by adding a single dose of Testa Triticum Tricum to the diet of 20 constricted subjects for at least six weeks.

From the data, the pool size, half life and input of deoxycholic acid were determined as well as the relative proportions of the three major bile acids. By combining these data the size of the total bile acid pool was estimated, accepting that a slight systematic underestimate of cholic acid is likely using this method.

STASTICAL ANALYSIS

Student's paired t test was used for determining the significances of differences. A p value of <0.05 was taken as significant. Results are expressed as mean ±SD.

Results

DIETARY INTAKE AND BODY MASS (QUETELET) INDEX

Dietary intake (Table 1) and body mass index (24.6±3.1 before and 24.7±3.1 after) remained unaltered throughout the study.

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Table 1 Dietary intake of 20 constipated subjects before and during supplementation of their diet with Testa Triticum Tricum (Mean over 24 h ±SD)

<table>
<thead>
<tr>
<th></th>
<th>Before</th>
<th>During</th>
<th>p value</th>
</tr>
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<tbody>
<tr>
<td>Energy (kcal)</td>
<td>1734±542</td>
<td>1745±562</td>
<td>NS</td>
</tr>
<tr>
<td>Total sugar (g)</td>
<td>76±36</td>
<td>79±48</td>
<td>NS</td>
</tr>
<tr>
<td>Added sugar (g)</td>
<td>45±25</td>
<td>48±35</td>
<td>NS</td>
</tr>
<tr>
<td>Total fibre (g)</td>
<td>22±7</td>
<td>20±7*</td>
<td>NS</td>
</tr>
<tr>
<td>Cereal fibre (g)</td>
<td>10±5</td>
<td>9±5*</td>
<td>NS</td>
</tr>
<tr>
<td>Protein (g)</td>
<td>65±20</td>
<td>67±22</td>
<td>NS</td>
</tr>
<tr>
<td>Fat (g)</td>
<td>69±25</td>
<td>73±26</td>
<td>NS</td>
</tr>
<tr>
<td>Carbohydrate (g)</td>
<td>216±93</td>
<td>207±82</td>
<td>NS</td>
</tr>
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* Excludes the contribution due to Testa Triticum Tricum

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Fig. 1 Whole gut transit-time in 20 constipated subjects before and after adding Testa Triticum Tricum to their diet for at least six weeks.
WHOLE GUT TRANSIT-TIME (Fig. 1).
With Testa Triticum Tricum transit time fell in all but one subject, the mean decreasing from 120±35 h to 68±35 h (p<0.001).

RELATIVE PROPORTIONS OF BILE ACIDS IN BILE (Fig. 2)
The proportion of deoxycholic acid fell from 26-6±12-0% to 23-8±11-8% (p=0-002). Conversely the proportion of cholic acid rose from 37-6±6-2% to 40-9±8-0% (p=0-008). The proportion of chenodeoxycholic acid did not alter significantly with treatment (35-9±9-4% before and 36-2±10-7% after).

DEOXYCHOLIC ACID POOL (Fig. 3)
With treatment the deoxycholic acid pool did not alter significantly (0-64±0-38 g before and 0-66±0-43 g after).

TOTAL BILE ACID POOL (Fig. 4)
Testa Triticum Tricum increased the total bile acid pool by 16% from 2-36±0-88 to 2-75±0-99 g (p=0-008).

CHOLESTEROL SATURATION INDEX (Fig. 5)
In 12 subjects whose initial cholesterol saturation index was >1-0, Testa Triticum Tricum lowered the index in nine, the group mean decreasing from 1-33±0-25 to 1-22±0-21 (p=0-008). Eight of the same 12 subjects had an initial cholesterol saturation index >1-2, which on treatment fell in all eight from a mean of 1-46±0-79 to 1-28±0-13 (p<0-0001). Taking the whole group of 20, the cholesterol saturation index fell from 1-13±0-32 to 1-07±0-29 (p=0-04).

DEOXYCHOLIC ACID KINETIC STUDIES
In the 11 subjects in whom it was measured, there

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**Fig. 2** Relative proportions of bile acids in bile acid pool in 20 constipated subjects before and after adding Testa Triticum Tricum to their diet for at least six weeks.
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There was a significant inverse correlation between change in deoxycholic acid pool and change in chenodeoxycholic acid % and between change in deoxycholic acid % and change in cholic acid %.

Discussion

As expected, Testa Triticum Tricum reduced the cholesterol saturation of bile when this was initially

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**Fig. 3** Deoxycholic acid (DCA) pool in 20 constipated subjects before and after adding Testa Triticum Tricum to their diet for at least six weeks.

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was no overall change in deoxycholic acid half-life (4.3±2.2 to 3.5±1.4 days; NS).

Deoxycholic acid input increased slightly from 0.14±0.09 to 0.15±0.08 g daily (p=0.014). Such a small increase is unlikely to have been of biological significance.

**Correlations (Table 2)**

Correlations between changes in transit and changes in bile composition before and after Testa Triticum Tricum were determined. There were no significant correlations between change in cholesterol saturation index and change in deoxycholic acid % or deoxycholic acid pool, nor between change in transit-time and change in deoxycholic acid % or deoxycholic acid pool, nor between change in transit time and change in cholesterol saturation index.

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**Fig. 4** Total bile acid pool in 20 constipated subjects before and after adding Testa Triticum Tricum to their diet for at least six weeks.
Testa Triticum Tricum cannot be blamed on a poor effect on colonic transit, because subjects have responded very well in this respect in both the present study and in a previous study in constipated subjects.21

Our other studies have suggested that there are correlations between changes in whole gut transit time, biliary deoxycholic acid and the cholesterol saturation of bile.8 Such correlations were not evident in the present study. Perhaps, these variables are dissociated in the presence of wheat fibre products. In the present study, one subject's transit time remained long yet her biliary deoxycholic acid fell (from 19.4 to 13.5%), while Pomare and Heaton8 reported three subjects whose transit times actually increased on bran yet their biliary deoxycholic acid levels fell.

Testa Triticum Tricum was less effective than expected from published bran studies in reducing deoxycholic acid levels in bile. Indeed there was no reduction at all in the size of the deoxycholic acid pool while deoxycholic acid intake actually rose slightly. As the product is an effective laxative, this relative ineffectiveness could be taken as evidence that bran lowers biliary deoxycholic acid by some means other than by speeding up colonic transit. It has been proposed that, like lactulose, bran may act by acidifying the right colon, as short chain fatty acids are released when it is fermented by bacteria; the lowered pH could inhibit 7α-dehydroxylase, the enzyme which aids the formation of deoxycholic acid from cholic acid.9 If this is the mode of action of bran, it raises the question why Testa Triticum Tricum should be less active than bran itself in being fermented into short chain fatty acids. Testa Triticum Tricum is quite powdery and small particles should be more readily attacked by bacterial enzymes. It differs from bran in containing no starch whereas bran is about 23% starch.10 This could be important if it is the undigested starch rather than the fibre in bran which is responsible for its effects on bile acid metabolism. Certainly, starch would be expected to be readily fermented by bacteria.

Reduction of biliary deoxycholic acid is the most plausible explanation for the beneficial action of bran on the cholesterol saturation of bile.1 22 In the case of Testa Triticum Tricum the mode of action is less clear because, although there was a fall in % deoxycholic acid, there was no significant correlation between the change in % deoxycholic acid (or the size of the deoxycholic acid pool) and change in cholesterol saturation index. Other mechanisms should also be considered. One possibility is expansion of the chenodeoxycholic acid pool. We did not measure this directly but presumably it increased as its contribution to the bile acid pool stayed the same.

raised, though to a modest degree. It is difficult to compare its efficacy with that of ordinary bran because, in the published studies, there have been variations in the subjects studied, in the type and amount of bran and in the duration of its administration. Quite impressive reductions in cholesterol saturation index have been reported in some bran studies, however—for example, from 1.43 to 0.76 and from 1.01 to 0.67.4 The relatively low efficacy of

Fig. 5 Cholesterol saturation index (CSI) of bile-rich duodenal fluid in 20 constipated subjects before and after adding Testa Triticum Tricum to their diet for at least six weeks.
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Table 2 Correlation coefficients for changes in bile composition, bile acid pool sizes and whole gut transit time in 20 constipated subjects given Testa Triticum Tricium

<table>
<thead>
<tr>
<th></th>
<th>Δ CA%</th>
<th>Δ CDCA%</th>
<th>Δ DCA%</th>
<th>Δ DCA pool</th>
<th>Δ Total bile acid pool</th>
<th>Δ CSI</th>
<th>Δ Transit time</th>
</tr>
</thead>
<tbody>
<tr>
<td>Δ CA%</td>
<td>-0.614*</td>
<td>-0.529*</td>
<td>-0.028</td>
<td>0.213</td>
<td>-0.341</td>
<td>-0.127</td>
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<tr>
<td>Δ CDCA%</td>
<td>-0.345</td>
<td>-0.480*</td>
<td>0.542*</td>
<td>0.115</td>
<td>0.231</td>
<td>0.166</td>
<td></td>
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<tr>
<td>Δ DCA%</td>
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<td>Δ DCA pool</td>
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<td>Δ Total bile acid pool</td>
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<tr>
<td>Δ CSI</td>
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<td>0.088</td>
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*p<0.05 †p<0.01 ‡p<0.001
CA% percent of cholic acid in bile acid pool; CDCA% percent of chenodeoxycholic acid; DCA% percent of deoxycholic acid; CSI cholesterol saturation index of bile-rich duodenal fluid.

while the bile acid pool expanded by about 16%. The beneficial effect of expanding the chenodeoxycholic acid pool by administering the bile acid orally is of course well documented but it has not been shown that cholesterol saturation falls with such a minor expansion of the pool. Moreover, fall in cholesterol saturation is normally associated with a rise in % chenodeoxycholic acid, not found by us. The cholic acid pool probably expanded as well because the proportion of cholic acid in the bile acid pool rose and so did the total bile acid pool. Expansion of the cholic acid pool, however, has not been shown to result in a fall in the cholesterol saturation of bile.

Previously, it has been suggested that a change in the proportion of deoxycholic acid in the bile acid pool results in a reciprocal change in the percentage of chenodeoxycholic acid, because of selective inhibition by deoxycholic acid of chenodeoxycholic acid synthesis.23 The validity of this relationship is, however, controversial.24-27 The present data are equivocal in that, while there was no significant correlation between changes in the percentages of these two bile acids, there was a weak but significant inverse correlation (r=-0.48) between change in the deoxycholic acid pool and change in % chenodeoxycholic acid.

The questions raised by the present findings can only be answered when more is known of the interrelationships between colonic function and microbial metabolism of bile acids on the one hand and the effects of wheat fibre and starch on the other.

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References