Ultrastructure of mouse intestinal mucosa and changes observed after long term anthraquinone administration

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SUMMARY In an attempt to study the relative toxicity of anthraquinonic laxatives on intestinal mucosa, we compared in mice the effects of fruit pulp containing sennosides A and B with those of a free anthraquinone, 1-8 dihydroxyanthraquinone. Observations have been made with transmission electron microscopy (EM) after 16 weeks of treatment with the two drugs. Although the doses used in this study were equipotent in terms of laxative activity, no damage to the intestinal tissue was observed with the sennosides. A number of changes, however, were detected in intestinal nervous tissues of all the animals treated with 1-8 dihydroxyanthraquinone, mainly in the form of vacuolisation of the axons, formation of lysosomal structures and in some cases appearances of fibrillar degeneration.

A large number of people habitually take laxatives over long periods of time. A small group of patients are serious abusers of laxatives, for complicated psychological reasons: they usually conceal the habit and may be difficult to diagnose. Laxatives of the anthraquinone group are presumed to be severe cell poisons following prolonged intake. Histological and ultrastructural studies in patients with chronic laxative misuse have indicated damage of the myenteric plexus and of the colonic epithelium.1 2

Damage of the myenteric plexus was reported by Smith1 in the mouse after intraperitoneal, or oral administration of syrup of senna. Unfortunately neither the quantity of syrup used, nor the strain of the animals were stated. Nevertheless, the mouse seems to be a good animal model for the study of laxative toxicity.

In previous experiments on mice (Swiss strain) we were unable to find any histological, or ultrastructural changes in the jejunal and colonic mucosa after oral administration of a standardised senna powder for 11 weeks at a dose equivalent to sennosides A and B 10 mg/kg – approximately 50-fold the therapeutic dose in man.3 Light microscope examination showed no difference between untreated and treated animals: autonomic nerve fibres and interstitial cells were normal in the two groups. Electron microscope (EM) examination showed no ultrastructural differences: enterocytes and nerve structures had the same aspect in control and treated animals, both in the jejunum and in the colon.

The differences in our results from those of others could be explained by the fact that syrup used by Smith may have contained free anthraquinones produced by hydrolysis of glycosides. Instability of syrup of senna has been pointed out by some authors.4 5 Unlike glycosides, free anthraquinones are absorbed from intestinal mucosa.6 They are then concentrated in the bile as conjugates and split into free anthraquinones by intestinal bacterial enzymes.7 High concentrations of anthraquinones in the intestinal wall associated with a possible direct effect on the jejunal may be more toxic to intestinal cells than glycosides. This difference in the metabolic pathway between glycosides and free anthraquinones has been reported by several groups8-12 and it has been suggested that the enhanced absorption of anthraquinones makes them much more toxic than glycosides.13

In order to test this hypothesis, we have compared in the mouse the effects of sennosides A and B with those of a synthetic aglycone at doses producing the same laxative activity.

Methods

ANIMALS
Experiments were performed on 30 NMRI strain
male mice (EVIC CEBA breeding, France), weighing 24 g at the start of the study. Before and during the experiments the animals were kept in polypropylene cages, except during laxative activity measurements (see below). Food (UAR. Ref. 104) and water were given ad libitum.

TREATMENT

Fruit pulp containing 0-17% sennosides A and B* or 1-8 dihydroxyanthraquinone† was used for dosing the animals. The 30 animals were divided into three groups of 10 and received by gastric tube 0-3 ml/20 g body weight/day for 16 weeks either: arabic gum suspension 2-5% (Group I); or fruit pulp suspension in arabic gum 2-5% containing sennosides 0-626 mg/ml (0-187 mg/20 g) (Group II); or Danthron suspension in arabic gum 2-5% containing 1-8 dihydroxyanthraquinone 16-7 mg/ml (5-01 mg/20 g) (Group III).

EXPERIMENTAL PROCEDURES

General appearance, behaviour and weight of the animals were recorded daily.

Measurements of laxative activity

Faecal output was measured daily during the first four weeks of the study. During six hours after dosing the animals were placed in separate cages fitted with a wire-mesh floor in order to recover faeces separately. Laxative activity was measured by Lou and Fairbairn method22 as modified by Brittain and d'Arcy,23 by counting the number of stools of soft consistency. Laxative activity was expressed as the percentage of soft stools relative to the total number of stools after six hours. Results are expressed as mean ± SD. Unpaired t test was used to determine the statistical significance between treated and control groups.

EM examination

The animals were killed after 16 weeks of treatment by decapitation and tissue specimens from each animal from the jejunum and the colon were taken for examination under EM. Jejunal tissue was sampled from a zone located at 20 cm distal to the pylorus. Colonic tissue consisted of the part of ascending colon distal to the caecum.

The specimens were fixed in glutaraldehyde 3% solution in 0-1 M cacodylate buffer, then rinsed in a solution consisting of 1 volume cacodylate buffer and three volumes of bi-distilled water. The samples were postfixed in 1% osmium tetroxide, dehydrated in alcohol and embedded in epoxy resin. Ultra-thin sections from three blocks from each animal were cut on LKB microtome and stained with uranyl acetate and lead citrate. Five grids (with four or five ultra-thin sections on each one) from each block were examined in a RCA EMV 3 G electron microscope. The examinations were performed by the same observer (Ph G), who was unaware of the experimental conditions.

Results

No animals died during the study. The general appearance, behaviour, and weight gain were normal in the three groups during the 16 weeks of the study. At the end of the study, the weights of the animals were respectively 48-1±4-7 g, 46-9±2-8 g, and 47-5±3-2 g (mean±SD) in groups I, II and III.

LAXATIVE ACTIVITY

The proportion of soft stools was significantly higher (p<0.001) in the animals given laxatives, but there was no significant difference (p>0.05) between the two treated groups. The (soft stools/total stools) × 100 ratios were 5±3 for group I, 31-7±8-7 for group II and 35-7±9-8 for group III.

EM RESULTS

Ultrastructure of jejunum and colon in controls

(group I)

Jejunal villi

The enterocytes were columnar in form with their typical apical microvilli. Mucus cells, argentaffin cells, Paneth cells, components of the lamina propria and muscle layers revealed their well known and usually described structures. The nerve elements were myelin free fibres with abundant neurotubules and neurofilaments (Figs 1 and 2).

Colon

The ultrastructure of the crypts of Lieberkühn, of the lamina propria and of the muscle layers were similar to those generally observed. Innervation of the tissue with myelin free fibres was again observed.

Changes observed in treated animals

(groups II and III)

The only differences detected in intestinal ultrastructure between controls and treated animals were in the intrinsic nervous tissue. In treated animals, there was no observable increase in smooth reticulum, or in the number of lysosomes, and no abnormalities in the microvilli of the brush border. Desquamation, a normal phenomenon in renewal cycle, was not increased in the treated animals. The mucus cells were abundant and exhibited intensive...
Fig. 1  Normal mouse nervous tissue: jejunum. General view of transversal section of axonal fibres. (×5000 original magnification).

Fig. 2  Normal mouse nervous tissue: colon. Neuronal and axonal components. (M) Mitochondria; (NS) Neurosecretions and (NT) Neurotubules. (×12000 original magnification).

Fig. 3  Nervous tissue of mouse given Tamarine: colon. Neuronal and axonal components. All have normal ultrastructure. (M) Mitochondria; (N) Nucleus; (NS) Neurosecretions and (NT) Neurotubules. (×10000 original magnification).

Fig. 4  Nervous tissue of mouse given Tamarine: colon. Longitudinal section of axons. All components are normal. (C) Collagen; (FIB) Fibrocyte; (M) Mitochondria; (NS) Neurosecretions; (NT) Neurotubules. (×10000 original magnification).
secretory activity. The Paneth cells in the jejunum were comparable with those of the control group. The presence of mitoses showed the regularity of the cell renewal cycle. No differences in the ultrastructure of jejunum and colon were detected between control animals and those receiving sennosides (group II). No sign of ultrastructural lesions of the nerve tissues were observed. The neurotubules and neurofilaments were abundant in the axons and the overall cell was perfectly normal (Figs. 3, 4, 5).

In the group treated with 1-8 dihydroxyanthraquinone (group III), a number of changes occurred in the nerve tissue, mainly in the form of vacuolisation of the axons and formation of lysosomal structures (Figs. 6, 7, 8, 9). The nature, extent and intensity of the lesions varied from very minor abnormalities to the appearance of fibrillar degeneration. The latter was rare, but all animals given 1-8 dihydroxyanthraquinone had neuronal damage and 40% of them exhibited severe neuronal changes. These changes occurred with a similar frequency and severity in the small intestine and in the colon.

Discussion

Using EM examination to study the effects of anthraquinone laxatives on the ultrastructure of the intestinal tissue of the mouse, we found abnormalities of the intrinsic nerve tissue after treatment with 1-8 dihydroxyanthraquinone, the changes being evident not only in the colonic but also in the jejunal mucosa. It was difficult to judge if the lesions were focal or diffuse. It must be pointed out that during the first four weeks of the study, no diarrhoea was observed in the treated animals. The doses used in this experiment are those we usually use to study anthraquinone laxative activity in the mouse.14 The neuronal damage observed after treatment with 1-8 dihydroxyanthraquinone seems to occur after long...
term treatment at pharmacological doses. Damage of the jejunal and colonic nerves without any changes in other structures suggest that the initial action of free anthraquinones may be neuronal stimulation, perhaps leading to increased intestinal motility.

This conclusion differs from the generally accepted theories concerning the mechanism of laxative action. Many investigators no longer consider effects on propulsive activity as a primary mechanism, but rather as secondary consequences of altered electrolyte transport with fluid accumulation in the intestine.\textsuperscript{12–21} All the experiments concerned with water and electrolytes transport, however, have been conducted \textit{in vitro}, or \textit{in situ}, using very high concentrations of anthraquinone laxatives. It remains to be established that the alteration of fluid transport is the primary effect after oral administration at pharmacological doses. If neuronal stimulation is also the primary effect of anthraquinone laxatives in man and if the same lesions are likely to occur in man after prolonged treatment with usual doses of 1-8 dihydroxyanthraquinone, this could explain why patients become resistant to laxatives and need to increase the dose. Damage of myenteric neurones may affect the coordinated propulsive activity of the intestine producing a delay in transit of the intestinal contents so that the laxative efficacy of the drug is decreased: patients are then tempted to use larger doses. This is likely to produce severe electrolyte disturbances, such as increased net secretion in the small intestine and decreased absorption in the colon. Unfortunately, faecal output was not measured during the 16 weeks of the study so that it is not possible to state whether laxative efficacy decreased with time in the 1-8 dihydroxyanthraquinone treated animals.

Although the doses of sennosides and 1-8 dihydroxyanthraquinone used in this study were equipotent in terms of laxative activity in the experimental animals, no damage to the intestinal nerve tissue was observed after treatment with sennosides. Tested by Lou and Fairbairn method,\textsuperscript{22} as modified by Brittain and d'Arcy,\textsuperscript{23} laxative activity of sennosides is 24-fold higher than that of

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**Fig. 8** Nervous tissue of mouse given 1-8 dihydroxyanthraquinone showing important axonal vacuolisation. (M) Mitochondria; (NS) Neurosecretions; (NT) Neurotubules; (SMC) Smooth muscle cell. (×27000 original magnification). (Colon).

**Fig. 9** Nervous tissue of mouse given 1-8 dihydroxyanthraquinone showing axonal degeneration (large vacuolisation, lysosomal structure). (LY) Lysosome; (SMC) Smooth muscle cell. (×6500 original magnification). (Colon).
1-8 dihydroxyanthraquinone. That ratio is of the same order as the ratio of doses therapeutically used in man. The relative low potency of 1-8 dihydroxyanthraquinone has been pointed out by other authors.\(^6\)\(^8\)\(^10\) Two important aspects of sennoside chemistry make them potentially less toxic than 1-8 dihydroxyanthraquinone. Firstly, their glycoside structure prevents absorption in the small intestine so that the doses needed to produce laxative activity are smaller than those of 1-8 dihydroxyanthraquinone. Secondly, they have as glycosides, no effect on water on water. They can only act at the colonic level where they have to be hydrolysed by bacterial enzymes to exert their laxative activity. Sennosides therefore, are unlikely to exhibit any pharmacological, or toxic effect on the small intestine either topically or systemically. These considerations, together with our results on intestinal mucosa of mouse, suggest that at equivalent laxative activity, the doses needed to produce any intestinal cell toxicity should be much higher for sennosides than for 1-8 dihydroxyanthraquinone.

Reference

21 Fling E. Antidiarrheal agents and laxatives: changing concepts. *Clin Gastroenterol* 1979; 8: (1).