Topical methotrexate alters solute and water transport in the rat jejunum in vivo and rabbit ileum in vitro

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SUMMARY The topical effect of methotrexate (MTX) on small intestinal hexose and ion transport has been studied using an in vivo steady state jejunal perfusion technique in the rat, and short circuited rabbit terminal ileum in Ussing chambers in vitro. In rat jejunum, perfusion with MTX (1 μmol/l) caused significant reductions in water, sodium, and glucose absorption within 110 minutes of exposure. Fructose absorption was, however, unimpaired. The same concentration of MTX, when added to the mucosal side of distal rabbit ileum caused significant increases in transmucosal potential difference, short circuit current and the unidirectional flux of chloride from serosa to mucosa. In the presence of a subphysiological magnesium concentration (0.3 mmol/l), MTX resulted in the abolition of net sodium absorption and the conversion of net chloride absorption to secretion. We conclude that MTX has a topical effect on small intestinal transport which is independent of its effect on crypt cell kinetics.

The early use of folic acid antagonists in cancer chemotherapy was associated with severe intestinal toxicity and extensive ulceration of small and large intestine occurred as a result of depressed crypt cell regeneration. In experimental animal, high dose or prolonged continuous methotrexate almost invariably results in hypoplastic villous atrophy, which causes malabsorption of a wide range of nutrients and drugs. In clinical practice, low dose methotrexate may also be associated with enterotoxicity and children with acute leukaemia receiving maintenance low dose MTX have impaired d-xylose absorption. Similar observations have been made in the rat where an eight day course of low dose aminopterin (a close analogue of methotrexate) resulted in hexose malabsorption but no morphological change. Although severe enterotoxicity on low dose methotrexate is unusual, protracted diarrhoea has been reported and it has been postulated that the drug may be responsible for poor weight gain in some patients and possibly the malabsorption of other drugs.

The present study was undertaken to determine whether enteral methotrexate in low doses, unlikely to cause changes in villous architecture, is associated with disturbances in intestinal transport which might contribute to diarrhoea. Effects on water and hexose transport were investigated using a steady-state perfusion technique in rat jejunum in vivo and effects on ion transport studied using short circuited rabbit distal ileum in vitro.

There is good evidence that hypomagnesaemia is a complication of diarrhoeal disorders, particularly those associated with an impaired intake of oral fluids and nutrients. Such circumstances may occur during cytotoxic therapy and the need for magnesium replacement may not always be fully appreciated as plasma Mg²⁺ concentrations are a poor indicator of total Mg²⁺ status. Furthermore, subphysiological concentrations of Mg²⁺ in vitro enhance the secretory response to Mg²⁺. The effects of MTX on ion transport were therefore determined under conditions of both physiological and subphysiological Mg²⁺ concentrations in vitro.

Methods

RAT STEADY STATE JEJUNAL PERFUSION IN VIVO

Perfusions

Male Wistar rats weighing 250–300 g were fasted overnight but allowed water ad libitum. Anaesthesia was induced and maintained using intraperitoneal pentobarbitone (6 mg/kg followed by 0.3 mg doses
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as required). Rectal temperature was maintained at 37°C using overhead electric lamps. The abdomen was opened by a midline incision and a length of jejunum extending 15–20 cm distally from the duodenojejunal junction was isolated. This was cannulated at both ends and the luminal contents gently flushed out using a 0.9 NaCl solution at 37°C. The jejunal loop was then returned to the abdomen. Warm perfusion fluid was infused continuously at a rate of 0.2 ml/min (Harvard 975 syringe pump). After an initial equilibration period of 50 minutes, five consecutive 20 minute collections of the effluent from the distal cannula were obtained. One group of animals was perfused with a basic solution which contained (mmol/l): NaCl 145; KCl 4; NaHCO₃ 25; glucose 2; and polyethylene glycol (PEG 4000) 3 g/l labelled with 20 μCi ¹⁴C PEG (Amersham International Ltd.) as a non-absorbable marker. pH was adjusted to 7.0 by gassing with CO₂; osmolarity of the solution was 290 mosmol/l. Further groups were perfused with the basic solution to which one of the following was added: MTX 1 μmol/l (MTX Parenteral Solution, Lederle); folic acid 1 μmol/l; fructose 20 mmol/l + MTX. Each animal was perfused with one solution only. At the end of each experiment the perfused loop was dissected, stripped of mesentery, opened along its mesenteric border, gently blotted dry and weighed. Each loop was then transferred into neutral buffered formalin solution and subsequently prepared for histological examination by light microscopy.

Analytical methods

The following were measured in duplicate in the initial perfusate and effluent solutions: sodium by flame photometry (Corning EEL); glucose by a hexokinase – G6PD method; fructose by initial conversion to G6PD by hexokinase and phosphoglucone isomerase, followed by glucose assay. Two hundred microlitre aliquots of solution were added to 10 ml scintillation fluid (RIA Luma) and counted for ¹⁴C in a liquid scintillation counter to determine PEG concentrations. For each perfusion period the PEG recovery was estimated and beyond the initial equilibration period the mean value for each experiment always exceeded 90%.

Calculations

Net rates of water and solute absorption were calculated using standard formulae¹³ and statistical comparisons made between groups using Student's t test.

Rabbit short circuited terminal ileum in vitro

After an overnight fast, Dutch brown male rabbits weighing between 1.8 and 2.3 kg were killed by a blow on the head and the terminal ileum mounted in Perspex half-chambers as previously described.¹² Each surface of the mucosa was bathed separately with 12 ml oxygenated (95% O₂/5% CO₂) Krebs-Hensleit solution at 37°C and pH 7.0 containing (mmol/l): Na, 143; K, 5.7; Ca, 1.9; Mg, 1.1 or 0.3; Cl, 125; HCO₃, 25; H₂PO₄, 1.2. Transmucosal potential difference (PD), short-circuit current (SCC) and uni-directional ion fluxes using ²²Na and ³⁶Cl were measured as described by Field et al.¹⁴ using the formula of Schultz and Zalusky.¹⁵

Fluxes of radio labelled sodium and chloride were measured on paired tissues taken from adjacent segments of distal ileum. Tissues were only paired if their electrical resistances differed by less than 25%. Forty minutes after mounting, 2-2 μCi of ²²Na and 2-2 μCi of ³⁶Cl (Amersham International Ltd.) were added to the mucosal solution of one tissue and to the serosal side of the other. Twenty minutes after the addition of isotopes, during which the tissues were short circuited, a 2 ml sample was removed from each unlabelled bathing solution and a 100 μl sample removed from each labelled solution for isotopic counting. Samples removed from the unlabelled solution were replaced with an equal volume of unlabelled bathing solution. Duplicate samples were taken 30 minutes later in order to determine base line fluxes. (Period A).

Under open circuit conditions, methotrexate (MTX) was added to the mucosal bathing solution to give a final concentration of 1 μmol/l. Forty minutes after the addition of MTX, the tissues were again short circuited and after a 20 minute equilibration period, fluxes were again measured from initial and final samples taken 30 minutes apart. (Period B).

Two series of experiments were performed: one in which MTX was added to tissue bathed in a solution containing a physiological concentration of Mg²⁺ in vitro (1.1 mmol/l) and a second in which tissue was bathed in a subphysiological concentration of Mg²⁺ in vitro (0.3 mmol/l). All values are expressed as means±1 SEM. Statistical comparisons of fluxes before and after MTX exposure were made using Students paired t test.

Results

Rat jejun al perfusion in vivo

Effects of methotrexate on net glucose, sodium and water transport

The inclusion of MTX (1 μmol/l) in the basic perfusate resulted in progressive decreases in the rates of net jejunal absorption of glucose, sodium
and water when compared with control animals perfused with basic perfusate alone (Fig. 1). Significant decreases were seen in net glucose absorption within 50 minutes of starting the perfusion (p<0.02), sodium by 70 minutes (p<0.01) and water by 110 minutes (p<0.01). When the experiments were concluded after 150 minute perfusion, transport rates in the animals perfused with MTX were still falling.

Histological examination by light microscopy of jejunal mucosa from MTX perfused and control animals, showed no evidence of morphological abnormality.

**Effects of folic acid (1 μmol/l) on net glucose, sodium and water transport**

To determine whether the impaired transport seen in animals perfused with MTX is likely to be caused by a direct blockade of the brush border membrane glucose carrier, further perfusions were carried out in which folic acid, an extremely close structural analogue of MTX, was added to the basic perfusate. This addition made no significant difference to the rate of glucose, sodium or water transport compared to controls.

**Effects of addition of fructose in the presence and absence of methotrexate**

In view of the inhibitory effects of MTX on jejunal transport when perfused with an actively transported hexose (glucose), its effects in the presence of an additional, passively transported hexose were investigated. Fructose, which is passively transported by a carrier mediated process was therefore added to the perfusate and a further group of animals studied. The addition of fructose (20 mmol/l) did not result in significant changes in the rate of sodium or water absorption when compared with rates in the presence of glucose alone (Figs. 1, and 2). When animals were perfused with a solution containing glucose, fructose and MTX the inhibitory effects of MTX on water and sodium were again apparent although occurring somewhat later. By contrast the absorption of fructose remained unimpaired throughout the experiment (Fig. 2).

**SHORT CIRCUITED RABBIT TERMINAL ILEUM IN VITRO**

**Methotrexate added to mucosal solution containing Mg\(^{2+}\) 1.1 mmol/l (Table)**

Addition of MTX resulted in small but sustained increases in short-circuit current (SCC) (p<0.001) and potential difference (PD) (p<0.01) which began within 10–15 minutes of MTX addition and continued rising slowly for the duration of the experiment. A significant increase in the flux of chloride from serosa to mucosa occurred (JCl\(_{sm}\)) p<0.05, sufficient to abolish net chloride absorption (JCl\(_{net}\)) although this latter change did not reach statistical significance. No significant

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**Fig. 1** Absorption of water, sodium and glucose from rat jejunum in the presence (■) and absence (○) of methotrexate. (1 μmol/l) n=10 in each group. Mean±SE.

**Fig. 2** Absorption of water, sodium and fructose from rat jejunum in the presence (■) and absence (○) of methotrexate (1 μmol/l) n=10 in each group. Mean±SE.
The occurrence of changes in the ion concentration of Mg\(^{2+}\) in chloride but significantly (p<0.01) was an increase in the flux of ions containing sodium fluxes. Increase to absorption studies.

Discussion

This study shows that MTX has a potent topical effect on small intestinal transport, which is rapid in onset and occurs in the absence of morphological mucosal change. MTX binds dihydrofolate reductase (DHFR), preventing the conversion of folic acid to tetrahydrofolate and blocking DNA synthesis. One major effect of systemic methotrexate is therefore on the actively dividing crypt enterocyte where a high percentage of cells in the synthetic (S phase) of division and are thus susceptible to methotrexate toxicity. Within a few hours of drug administration, inhibition of mitosis is evident\(^{17,18}\) and where this is sustained an imbalance develops between the shedding of mature enterocytes and their replacement from the crypts. Prolonged exposure after high doses or continuous administration is associated with progressive villous hypoplasia and ultimately severe atrophy.

The changes in intestinal transport associated with local MTX cannot be explained on the basis of such effects on intestinal epithelial cell turnover and appear to result as a direct effect on the enterocyte. Dihydrofolate reductase is also an important coenzyme in a number of other one carbon transfer reactions and consequently the synthesis of RNA and protein in mature cells may also be influenced by MTX.\(^{19}\) Previous studies have suggested that the mature villous enterocyte is affected by this non-S phase action. Within four hours of a single high dose of aminopterin, given systemically, there was a marked reduction in mucosal oxygen consumption and glycolysis\(^{20}\) and patchy cytoplasmic vacuolation is seen in villous enterocytes within three hours of low dose MTX administration in both children with leukaemia and adults with psoriasis.\(^{17,27}\) The precise mechanism of this early systemic effect is unclear, but may be due to disruption of cellular enzyme systems as a result of the inhibition of protein synthesis.

In the rat jejunum in vivo, the progressive decrease in net sodium, water and glucose absorption associated with local exposure to MTX could not be reproduced by folic acid, a close structural analogue of MTX, suggesting that impaired absorption is not simply because of competition by MTX for glucose binding sites. Fructose was normally absorbed in the presence of MTX. This preservation of fructose absorption,
which occurs passively by a carrier mediated process, but inhibition of active glucose absorption
supports the notion that MTX does not act directly on brush border membrane transport. It seems more
likely that it inhibits one or more of those intracellular events which follow and are necessary for
continuing glucose absorption. Studies by Kaminskas support this hypothesis. In an Erlich ascites tumour cell preparation it was shown that
MTX rapidly inhibited glycolysis, depressed cellular ATP and impaired glucose uptake. The latter was
restored by correction of ATP depletion. There was no evidence that MTX affected the saturation
characteristics of glucose transport and the changes in uptake were observed before suppression of
protein synthesis by MTX became apparent.

Although high dose systemic MTX leads to reduced activity of Na⁺K⁺ ATPase in rat jejunal enterocytes within 48 hours the role of this enzyme in
the mediation of the currently reported changes is unknown.

In addition to effects on jejunal hexose transport topical MTX also induces early and marked changes in rabbit ileal electrolyte transport in vitro. These
changes are exaggerated, both quantitatively and qualitatively, in the presence of sub-physiological
conzentations of Mg. A similar enhancement has been observed in the secretory response of ileum to high Mg concentrations in vitro.

Although intestinal secretory has been previously described in association with radiation enteropathy, the changes in transport currently described occur in the
absence of gross mucosal morphological changes. Taminiau et al. have reported reduction in net sodium fluxes, in response to glucose, after systemic
MTX, but did not examine the effects on chloride transport in their experiments. An early effect upon
intracellular glycolysis has previously been reported but the relevance of this to our findings is
unclear. Thus the mechanisms by which transport is influenced remain speculative and although these
data show that functional disturbances occur they give few clues to the precise factors involved. Study of other cytotoxics which act at different sites on
cellular metabolism would be of value to elucidate the mechanisms of topical toxicity. All previous
studies of cytotoxic enteropathy have, however, considered only the functional derangement associated with gross villous damage.

What are the possible clinical implications of this local toxic effect? In addition to direct exposure from oral doses up to 10% of a single dose of MTX
is recycled within the enterohepatic system irrespective of the route of administration, and this
provides a mechanism by which the jejunal mucosa might be exposed for long periods to potentially
toxic drug concentrations.

The precise concentration of MTX in the small gut after oral or parenteral doses is unclear. The
only published data showed that in a single case 24 hours after a small dose (5 mg) the jejunal
concentrations were 8×10⁻⁶M and 17×10⁻⁶M after oral and intramuscular administration respectively.
At five days the concentration was 62×10⁻⁶M showing considerable enterohepatic recycling. It is
likely that higher concentrations are found in the small gut during the early absorptive phase after an
oral dose or within a few hours of a parenteral dose. With current intermediate dose (500 mg/m²) and
high dose (up to 13 g/m²) schedules these concentrations probably exceed 10⁻⁶M. Moreover
after oral doses the duration of exposure of the small gut to MTX is variable. A group of 'slow absorbers'
has been defined in whom serum drug profiles indicate prolonged residence in the small gut. In
the present study the concentration of 10⁻⁶M was chosen as this is generally accepted as the
concentration at which the drug exerts a cytotoxic effect on the intestinal mucosa.

Protracted diarrhoea might therefore occur where there is an unusually high amount of MTX excreted in the bile and topical toxicity could also explain the
prolonged xylose malabsorption associated with intermittent MTX administration in children with
leukaemia. Malabsorption is evident for up to three weeks after a single dose of MTX, at which time the
jejunal mucosa would be expected to have recovered from any systemic effects. Furthermore
jejunal morphology at this time has been shown to be normal. In conclusion these data show that
MTX alters small intestinal transport by a previously unrecognised local action which may have an
important role in the pathogenesis of malabsorption and diarrhoea seen in some patients on cytotoxic
therapy.

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