Lipid peroxidation and electrogenic ion transport in the jejunum of the vitamin E deficient rat

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Abstract
Increased concentrations of reactive oxygen species in children with depleted antioxidant defences have been implicated in a cycle of malnutrition, malabsorption, and infection leading to protracted diarrhoea. A rat model of chronic vitamin E deficiency has been used to study the effects of antioxidant depletion on jejunal structure and function in vitro. Basal intestinal short circuit current (Isc), a measure of net electrogenic ion movement across the intestinal epithelium, was greater in chronically vitamin E deficient jejunum than controls, as was the electrogenic secretory response to aminophylline and Escherichia coli STa but not to bethanecol. The galactose stimulated current was also greater in vitamin E deficient jejunum. Indices of lipid peroxidation (concentrations of thiobarbituric acid reactive substances and malondialdehyde) were increased in the vitamin E deficient small bowel. Small intestinal brush border membranes from vitamin E deficient animals displayed changes in both static and dynamic components of membrane fluidity measured by steady state fluorescence polarography. The results of these studies support the hypothesis that oxidative stress in subjects with compromised antioxidant defences results in small intestinal hypersecretion, which could predispose to or perpetuate protracted diarrhoea.

Protracted diarrhoeal disease is an important cause of morbidity and mortality in young children worldwide. While this represents an enormous problem in the Third World, idiopathic protracted diarrhoea may account for a significant portion of paediatric gastroenterologists' workload. Despite the magnitude of this health problem, little is known about many features of its pathogenesis. Larcher et al. have implicated a vicious cycle of infection, malabsorption, and malnutrition in the pathogenesis of protracted diarrhoea, yet the mechanism(s) whereby an apparently minor episode of enteric infection progresses to a potentially life threatening protracted diarrhoeal disease is often unclear. Studies of malnourished infants and children in the West Indies have reported a decrease in blood concentrations of vitamins A and E, and of a number of micronutrients all of which have important roles in limiting cell damage after an oxidising stimulus. Infection, a common precipitant of protracted diarrhoeal disease, can produce a potent oxidising stimulus which, in the presence of impaired antioxidant defences, would be more likely to damage the lipid and proteins of cell membranes. This damage might be expected to have an effect upon the functions of receptors, enzymes, and transporter proteins within that membrane. Vitamin E is the most important chain breaking antioxidant in biological membranes, which is present in the small intestine in amounts comparable with those found in many other tissues. A deficiency of vitamin E may therefore render the enterocyte membrane more susceptible to oxidative damage. We have examined the hypothesis that vitamin E deficiency results in oxidative damage to enterocytes leading to a change in gastrointestinal function. Cell damage sustained in this manner might therefore be important in the pathogenesis of protracted diarrhoeal disease. In this paper we report the effects of chronic vitamin E deficiency in a well validated rat model upon parameters of lipid peroxidation, membrane biophysical characteristics, and intestinal transport in vitro in rat jejunum.

Materials and methods

MATERIALS
Most chemicals were purchased from Sigma Chemical Company Ltd, Poole, England. 1,6-diphenyl-1,3,5-hexatriene (DPH) was obtained from Aldrich Chemical Co (Gillingham, UK) and DL-12-(9-anthroyloxy)-stearic acid (12-AS) from Molecular Probes Inc (Junction City, Oregon, USA).

ANIMALS AND DIET
Two groups of 21 day old weaning male Wistar rats (Charles Rivers) were placed on synthetic diets (Dyets Inc, Bethlehem, Pennsylvania, USA), based on tocopheryl stripped lard (10%), vitamin free casein (20%), and glucose (65%) with a vitamin E free mineral and vitamin mix (Machlin/Draper-HLR No 814). In one group (vitamin E sufficient (E⁺)) the diet was supplemented with α-tocopheryl acetate (40 mg/kg of feed). All animals were permitted continued free access to food and water. Animals were housed four to a cage in a room maintained at a temperature of 70°C and humidity of 70% on a 12 hour light/dark cycle. Balance studies were carried out in six animals from both groups (studied individually in metabolic cages) over a five day period in the week before the in vitro transport study.

ELECTROGENIC SECRETION AND ABSORPTION IN VITRO
Electrogenic secretion and absorption in vitro was studied after 52 weeks on the respective diet.
After an 18 hour fast, during which time free access to water was permitted, animals were anaesthetised with intraperitoneal sodium pentobarbitone (Sagatal, 60 mg/kg bodyweight). A 3 cm segment of jejunum, beginning 15 cm distal to the ligament of Treitz was removed by a midline incision, irrigated with ice cold Ringer bicarbonate solution (containing (mmol 1⁻¹) NaCl 107, KCl 4.5, NaHCO₃ 25, Na₂HPO₄ 0.2, NaH₂PO₄ 0.2, CaCl₂ 1.25, MgSO₄ 1.0, pH 7.4, preoxygenated with 95% O₂/5% CO₂), and was transported in this same cold sugar free solution before mounting in a modified Ussing chamber. This segment of jejunum was opened along its mesenteric border and mounted unstripped across a number of pins between perspex half chambers, exposing an area of 2 cm². The maximum time between removal from the animal and mounting in the Ussing chamber was 10 minutes. Each side was bathed with 15 ml of the Ringer bicarbonate solution at 37°C oxygenated by a gas lift buffer with 5% CO₂/95% O₂. Glucose (final concentration 10 mM) was added to the serosal buffer and the same amount of mannitol to the mucosal buffer after mounting the tissues. Transepithelial potential difference was measured with calomel electrodes in saturated KCl by agar salt bridges (3% agar wt/vol in saturated KCl washed in Ringer solution), the tips of which were positioned near the tissue surface. Tissues were continuously short circuited at 0 mV with an automatic voltage clamp (World Precision Instruments, New Hampshire, USA) by Ag/AgCl electrodes connected to the bathing solutions with salt agar bridges. An adjustable offset with the clamp circuit compensated automatically for the fluid resistance between the tissue and the tips of the potential sensing electrodes. Tissues were unclamped every three minutes to allow measurement of open circuit potential difference and calculation of tissue resistance (Rt) using Ohm's law. A single dose of secretagogue, or mucosal galactose coupled with the simultaneous addition of an equal amount of serosal mannitol, was added 15 minutes after mounting when basal short current circuit (Isc) had reached a stable plateau. No tissue was exposed to more than one of these agents.

The adjacent (aboral) 20 cm of small bowel was flushed with ice cold Ringer solution, opened along its mesenteric border, and mucosal scrapings were made with a microscope slide. These were snap frozen for biochemical analyses.

**BRUSH BORDER MEMBRANES**

Brush border membranes were prepared by MgCl₂ precipitation and differential centrifugation and purified on a Percoll density gradient as described by Yakymyshyn et al.⁴. Enrichment of mucosal disaccharidases and alkaline phosphatase was checked by the methods of Dalquhist⁵ and Wilkinson.⁴

**PROTEIN AND LIPID ASSAYS**

Protein was measured with a commercial kit using bicinchoninic acid (Pierce, Illinois). The total lipid content of mucosal scrapings was measured by the hydroxamic acid method.⁹

**VITAMIN E**

Vitamin E was measured by high performance liquid chromatography (HPLC) by a modification of the procedure described by Metcalfe et al.¹⁰ as described previously.³

**LIPID PEROXIDATION**

Lipid peroxidation was estimated in mucosal scrapings by two methods.

**Thiobarbituric acid reactive substances** – these were measured according to the method of Yagi.¹¹ The resulting fluorescence was measured in a Perkin Elmer LS-3 fluorimeter (Ex 515 nm, Em 553 nm) with appropriate blanks and assessed against a standard calibration curve of malondialdehyde (0–2 nmol/l) prepared by acid hydrolysis of tetramethoxypropane.

**Free and total malondialdehyde** – free malondialdehyde was measured as the 2,4-dinitrophenyldihydrazine (DNPH) derivative by a modification of the methods of Ekstrom¹² and Kawai.¹³ In summary 100 µl of a 25% (wt/vol) homogenate of mucosal scrapings in 0.1 M phosphate buffer pH 7.4 was mixed thoroughly with an equal volume of acetonitrile, spun at 16 000 rpm in an Eppendorf centrifuge for two minutes, and 125 µl of the supernatant decanted. To this was added 12.5 µl 2 M HCl, 5 µl DNPH (0.5 mg/ml in 1 M HCl) and 5 µl 62.5 mM nitoresocoril (internal standard). The mixture was allowed to stand at room temperature for 60 minutes and a 20 µl aliquot analysed by HPLC. HPLC conditions were mobile phase 45% acetonitrile/55% 0.1 M HCl, flow rate 1 ml/min, column Spherisorb 5-ODS2 (250×4.6 mm ID), reverse phase (orb men maintained at room temperature and fitted with a Spherisorb 5-ODS column. Malondialdehyde peaks were detected by ultraviolet absorption at 310 nm and the peak areas compared with malondialdehyde standard after correcting for the amount of internal standard. Total malondialdehyde was measured after subjecting the sample to alkaline hydrolysis in 0.1% sodium hydroxide at about 60°C for 30 minutes in the presence of 0.05% (wt/vol) promethazine as an antioxidant. In validating this method of measuring malondialdehyde we found it to have an intrabatch coefficient of variation of 3-3%. Recoveries of malondialdehyde were about 96.5%.

**TOTAL FATTY ACID ANALYSIS**

A chloroform/methanol (2:1) lipid extract of brush border membranes was made according to the method of Folch³⁴ after the addition of non-adecanoic acid internal standard. This total lipid extract was dried down under nitrogen, taken up into dry 3 M methanolic HCl (Superco) for saponification at 65°C for four hours, and later extracted into hexane for analysis by gas chromatography. Gas chromatography-mass spectrometry was performed on a Hewlett Packard 5890 gas chromatograph coupled to a 5970 series mass selective detector. The fatty
acid methyl esters were separated on a fused silica capillary column (30 m × 0.25 mm inside diameter) with a 0.25 μm chemically bonded BDI stationary phase (J and W Scientific, Folsom, CA95630 USA). The initial column temperature was 50°C rising at 50°C/min to 190°C, 2.5°C/min to 213°C, 5°C/min to 260°C, and 60°C/min to 300°C. Aliquots of 1 μl were injected in splitless mode with an injector temperature of 250°C. The detector temperature was 300°C. Fatty acids were identified by their retention time and by comparison of their total ion chromatogram with that of known standards. Quantitative measurement was achieved by peak area integration with reference to a known concentration of standards. The coefficients of variation for this method varied from 1.5% for the saturated and monounsaturated fatty acids to about 8% for arachidonic acid and docosahexaenoic acid. These are comparable with those found in published works.10

FLUORESCENCE ANISOTROPY

Steady state fluorescence polarisation studies were performed using a Perkin Elmer LS-3 fluorescence spectrophotometer according to standard methodology.11 Two hundred μl of brush border membrane suspension (about 200 μg of protein) was incubated for 30 minutes at 30°C with 1 ml of 5 μM DPH or 12-AS in hydroxyethylpiperazine-ethanesulfonic acid (HEPES) buffer. At the end of this period the brush border membrane was pelleted and resuspended in fresh HEPES buffer. The content of fluorophore in the preparations was estimated fluorometrically as described by Cogan.12 Corrections for light scattering and for fluorescence in the ambient medium were routinely made. Excitation and emission parameters were as follows: for DPH Ex 360 nm, Em 430 nm; for 12-AS Ex 360 nm, Em 471 nm.

Anistropy was calculated from the polarised fluorescence intensities using the standard equation anisotropy, R0 = (Iv - In)/(Iv - 2In).13

HISTOLOGICAL STUDIES

Small pieces of jejunum, immediately adjacent to that used in the Ussing chamber, were either fixed in 4% paraformaldehyde in 0.01 M phosphate buffered saline, 2.5% (wt/vol) glutaraldehyde in 0.1 M phosphate buffer, or snap frozen to allow processing for light and electron microscopy.

STATISTICS

Unless otherwise stated, comparisons were made using Student’s unpaired t test. Non-parametric data were analysed using the Kruskal-Wallis one way analysis of variance. Figures in parenthesis after the means or medians are the 95% confidence intervals for this parameter unless otherwise stated. n Refers to the number of animals in each group.

Results

Figure 1 shows the longitudinal growth data for the two groups of animals. The weight of the E− group lagged behind that of the E+ group from 20 weeks after weaning onto the diets. After 52 weeks the E− animals were significantly lighter than E+ (E+ mean 733 g (40), E− 519 g (44), n=40 p<0.01). Animals on vitamin E free (E−) diets developed signs of a hind limb ataxia after about six months on the diet, as previously described.13 An important element in the difference in weight between groups resulted from muscle wasting – a characteristic of the neurological syndrome accompanying chronic vitamin E deficiency in this rat model.14

E− rats ate significantly less diet than E+ when balances were performed in the week before the in vitro transport studies (E+ 23.5 g/kg−24 h−1 (1.8) v E− 21.5 g/kg−24 h−1 (1.2), n=12, p<0.05). Neither group of animals developed diarrhoea, and stool output (g/24 h) was the same in both groups.

Plasma vitamin E concentrations declined rapidly after weaning in the E− group and rose to a stable plateau in the E+ group (Fig 2). Sixteen weeks after starting the diets the mean plasma concentration was 34.5 (1.4) μmol/l in E+ animals, but undetectable in the E−.

After 52 weeks on the diet, vitamin E was undetectable in mucosal scrapings from E− jejunum, but abundant in E+ jejunum (mean 1.33 (0.09) μmol/g lipid, n=6).

Basal short circuit current was significantly higher (p<0.005) in the E− group than the E+ group (Table I). This is reflected in the higher transepithelial potential difference. Tissue resistances were comparable. Substitution of the bathing solutions with a chloride free medium on both the mucosal and serosal sides (gluconate replacing Cl−) led to a reversible 75–80% decrement in basal Isc in both groups. Application of 1-25 μM tetrodotoxin to the serosal surface of unstimulated tissues had no effect on Isc in either group. Table II shows the effect of added secretagogues. Ten mM mucosal and serosal amineophylline produced a greater increment in Isc in the E− animals as did mucosal E coli heat stable toxin (Sta) at a concentration of 60 mouse units ml−1. The increment in Isc seen with 1 mM serosal bethanecol was comparable in E+ and E− jejunum.

The maximal A Isc produced by the addition of 28 mM galactose to the mucosal surface was significantly higher in the E− group (114 μA cm−2 (29) v 75 μA cm−2 (18), n=8, p<0.05) as
were the maximal responses to 5, 10, and 20 mM galactose (Fig 3).

Mucosal thiobarbituric acid reactive substances were significantly greater in the E-animals (E+ 0.67 (0.09) vs E- 0.83 (0.09) nmol/mg protein, n=10, p<0.05).

Concentrations of both free and total malondialdehyde were significantly higher in the E-jejunum. (Free malondialdehyde E+ 100 (20) pmol mg⁻¹ protein v E- 307 (96) pmol mg⁻¹ protein, n=11, p<0.01; total malondialdehyde E+ 168 (30) nmol/mg protein v E- 397 (109), p<0.05, n=6). Higher concentrations of malondialdehyde seemed to be associated with higher short circuit currents (Fig 4). It would be unreasonable to expect a simple linear relation between Isc and malondialdehyde and hence regression data are not provided.

Steady state anisotropies with both probes in proximal intestinal brush border membranes with higher in E-animals. (DPH E+ 0.209 (0.008 – 99% CI) vs E- 0.231 (0.005), n=6, p<0.01; 12-AS E+ 0.0902 (4.6×10⁻³ – 99% CI) vs E- 0.0967 (6.2×10⁻³), n=6, p<0.01).

The total fatty acid composition of the two groups of brush border membrane (expressed as mol %) showed no statistically significant differences in fatty acid profile (Table III). The ratio of total lipid to protein in the brush border membrane (μmol/mg) was, however, significantly higher in the controls (E+ 0.62 (0.03), E- 0.54 (0.04), n=12, p<0.05), and so the absolute amount of each fatty acid present was less in the E-brush border membrane.

Histological studies of sections cut from jejunum of both groups showed no significant differences with either light or electron transmission microscopy (Fig 5).

Discussion

Protracted diarrhoeal disease remains an important world health problem accounting for roughly five million deaths worldwide each year. The pathogenesis of the progression from a seemingly minor infective diarrheal episode to this life threatening disorder remains ill understood. The intimate association of protracted diarrhoeal disease with malnutrition and malabsorption and the compelling arguments of Golden et al led us to explore the relation between vitamin E deficiency, free radical production, and small intestinal secretion and absorption.

This study shows an increase in both basal and
secretagogue induced electrogenic anion secretion and galactose coupled cation absorption in the small intestine in chronic vitamin E deficiency. These findings are associated with increased levels of lipid peroxidation within the small intestinal mucosa and a biophysical change in the enterocyte apical membrane.

We have used vitamin E deficiency as a means to increase free radical fluxes because this is the most important lipid soluble antioxidant in biological membranes. Vitamin E, however, has both antioxidant and structural roles in biological membranes, and so our model is one of both vitamin E depletion and increased free radical fluxes. The small intestine has an innate resistance to oxidative damage, which is independent of vitamin E, and this may account for the comparatively small changes seen in both lipid peroxidation and electrogenic secretion despite a chronic and severe deficiency of vitamin E. Increasing free radical fluxes within the small intestine by other means, for example with ionising radiation, has been shown to result in a decrease in net fluid absorption. Recently, an increase in electrogenic chloride secretion and a reduction in electroneutral sodium and chloride absorption in colon has been shown after exposure of the serosa to hydrogen peroxide.

These effects were apparently largely because of the actions of released prostaglandins on nerves within the lamina propria, and almost certainly not to oxidative damage to the enterocyte membrane. In contrast, in this study the increase in short circuit current (Isc) seen in vitamin E deficient jejunum could not be abolished by neural blockade with tetrodotoxin. Our findings, therefore, suggest that oxidative damage to the enterocyte in vitamin E deficiency is associated with small intestinal hypersecretion.

The E− group of animals ate less food than E+, suffering in effect from chronic undernutrition. We did not pair feed the animals. If vitamin E deficiency induces anorexia or satiety then we felt this appropriate for the purposes of this study. Undernutrition (33% of normal intake for nine days) has no effect on basal Isc in stripped duodenum, jejunum, or ileum, but is associated with increased electrogenic secretory responses in jejunum to the Ca2+ mobilising agonist bethanecol, 5 hydroxtryptamine, and prostaglandin E2, and also to E coli STa. The secretory response to dibutyryl cAMP and forskolin, which cause secretion by cyclic AMP dependent mechanisms was not increased in these studies. These patterns of jejunal hypersecretion are quite different from those found in our vitamin E deficient 'chronically undernourished' rat.

Chronic vitamin E deficiency in humans and in our model is associated with a characteristic neuropathy of the peripheral, central, and autonomic nervous systems. The enteric nervous system is important in the control of intestinal ion transport. We were unable to show any histological or histochemical evidence of a neuro-pathy in E+ or E− jeuna, or any pharmacological evidence of an increase or decrease in neural tone that might have given rise to the noted changes in secretion and absorption.

We used two methods to detect oxidative damage within the intestinal mucosa. The thio-barbituric acid assay is a widely used measure of lipid peroxidation. This test is sensitive not specific and prone to interference. We therefore measured malondialdehyde by high performance liquid chromatography, which is both sensitive and specific. Malondialdehyde is one of a group of cytotoxic aldehydes formed as secondary products of lipid peroxidation after the metal ion dependent decomposition of lipid hydroperoxides. Our findings of increased concentrations of both thiobarbituric acid reactive substances and free malondialdehyde in E− jeuna are consistent with a greater degree of oxidative damage within these jeuna. The association of higher concentrations of mucosal malondialdehyde with higher values for basal Isc may provide indirect evidence that oxidative damage rather than vitamin E deficiency in itself gives rise to the increased secretion, though this matter requires further clarification.

The changes in membrane biophysical characteristics reported in this study could have arisen by a number of means including a variation in the lipid/protein ratios within the membranes, directly as a consequence of vitamin E deficiency, or as a consequence of peroxidation and crosslinking of membrane lipids or lipids
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with proteins. Vitamin E depletion alone without lipid peroxidation would, however, be expected to decrease the measured anisotropies with both probes – that is, 'fluidise' the membranes. A decrease in the lipid/protein ratio of the apical membrane, as has been noted to occur in the vitamin E deficient animals, would have the effect of increasing the measured anisotropy with both the probes used though given the magnitude of the change it is unlikely that this accounts for the increase in anisotropy on its own. The anisotropy data presented here pertain to a membrane 'average', and the use of probes that partition into different membrane lipid domains might provide further understanding of the changes we have seen. The relation between these changes in fluidity and enhanced intestinal secretion is unclear. There are other mechanisms whereby oxidising stress might predispose to secretion including direct actions upon membrane proteins and release of pro secretory substances from the subepithelium.

This study shows that chronic vitamin E deficiency is associated with oxidative damage within the small animal mucosa, a change in enterocyte apical membrane biophysical characteristics, and enhanced basal and secretagogue induced secretion and sodium coupled glucose absorption. The relation between these findings is worthy of further investigation.

There are clearly many reasons why malnourished children should be prone to develop protracted diarrhoea. Our data lend support to the hypothesis that malnourished children with depleted antioxidant defences are predisposed to upper small intestinal hypersecretion. We suggest that this is one mechanism whereby malnutrition might predispose to the perpetuation of the diarrhoeal state. This will have implications not only for subjects with idiopathic diarrhoea but also for malnourished subjects in whom a specific cause for the diarrhoea has been ascertained.

Parts of these data have been presented to the British Society of Gastroenterology, the European Society of Paediatric Gastroenterology and Nutrition, and the Physiological Society, and have appeared in their publications. The diarrhoea study was a pilot study and was not power calculated for malnourished subjects in whom a specific cause for the diarrhoea has been ascertained.

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