Increased antigen and bacterial uptake in follicle associated epithelium induced by chronic psychological stress in rats

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Background: Chronic stress affects the course of inflammatory bowel disease and experimental colitis, and may also initiate intestinal inflammation in rats. Aim: To investigate the effects of stress on the M cell containing follicle associated epithelium, specialised in antigen uptake.

Subjects and methods: Wistar rats were submitted to acute water avoidance stress for 1 hour or chronic water avoidance stress for 1 hour/day for 10 consecutive days. Permeability to $^{51}$Cr-EDTA, horseradish peroxidase, and chemically killed Escherichia coli K-12 was studied in both villus and follicle associated epithelium in Ussing chambers. Segments were further examined by light, electron, and confocal microscopy.

Results: Acute stress increased horseradish peroxidase flux in villus as well as in follicle associated epithelium. Chronic stress further increased permeability to horseradish peroxidase in villus and follicle associated epithelium, in the latter by almost fourfold. Moreover, chronic stress induced over 30 times increased E coli passage in follicle associated epithelium whereas there was no significant increase in villus epithelium. Bacterial uptake was confirmed by confocal microscopy showing fluorescent bacteria penetrating and passing through the epithelial surface.

Conclusions: These results show that the barrier function of follicle associated epithelium can be modulated, and that chronic stress enhances the uptake of luminal antigens and bacteria via the follicle associated epithelium. This can increase antigen exposure in Peyer’s patches thereby having implications in the initiation of proinflammatory immune responses within the intestinal mucosa.

Materials and methods

Animals

The study was approved by the local committee of animal ethics. Thirty six male Wistar rats (B&K Universal AB, Sollentuna, Sweden), body weight 150–200 g at arrival, were used. After arrival, rats were acclimatised for one week and then handled daily for two weeks before the stress protocols were initiated. During these three weeks, and the rest of the experimental period, animals were kept under pathogen free conditions with a 12:12 hour light:dark cycle and fed ad libitum on a diet of standard rat pellets (R70; Lactamin AB, Vadstena, Sweden) and tap water.

Acute stress

Six rats were subjected to acute stress, with one hour of water avoidance stress (WAS), and six other rats were subjected to sham stress (SS) and used as controls. WAS rats were placed on a platform (height 8 cm, depth 6 cm) in a plastic container (height 56 cm, depth 50 cm) with 25 degrees C water, 1 cm below the platform. SS rats were treated in exactly the same manner except for the platform.

Abbreviations: CD, Crohn’s disease; FAE, follicle associated epithelium; HRP, horseradish peroxidase, IBDD, inflammatory bowel disease; M cell, membranous cell; SS, sham stress; VE, villus epithelium; WAS, water avoidance stress; PBS, phosphate buffered saline
same way as WAS rats except that during the stress session SS rats were placed on a similar platform but in a waterless container. After one hour, the number of faecal pellets were counted as an indirect index of changes in colonic propulsive activity.\(^2\) \(^3\) Rats were anaesthetised by isoflurane inhalation and 15 cm segments from the distal ileum (starting 5 cm proximal to the ileocecal valve) were taken out and immediately transported to the laboratory in ice cold oxygenated Krebs buffer (115 mM NaCl, 1.25 mM CaCl\(_2\), 1.2 mM MgCl\(_2\), 2 mM KH\(_2\)PO\(_4\), and 25 mM NaHCO\(_3\); pH 7.35).

**Chronic stress**

The results of the acute stress experiments confirmed stress mediated effects on FAE. As chronic stress is more interesting from a pathogenic point of view, 24 Wistar rats were subjected to WAS or SS (controls) for 10 consecutive days, 1 h/day. To minimise the effect of circadian rhythm, the procedures were started between 9am and 10am. Before each session, rats were weighed, while the number of faecal pellets were counted following the WAS or SS period. After the final period, rats were anaesthetised and intestinal segments were taken out as described above. Blood for corticosterone analysis was collected by cardiac puncture and centrifuged at 3000 \(g\) for 10 minutes. The collected serum was stored at \(-70^\circ\)C until further use for corticosterone assay (see below).

**Ussing chamber experiments**

While immersed in ice cold oxygenated Krebs, the external muscle and myenteric plexus were stripped of distal ileal specimens. Sections of VE and FAE, identified macroscopically, were cut into appropriate sizes and mounted in modified Ussing chambers (Harvard apparatus Inc., Holliston, Massachusetts, USA).\(^2\) \(^3\) When mounting FAE, segments were carefully adjusted so that the patches covered the entire exposed tissue surface area of 0.096 cm\(^2\). Before placing the chambers in the 37°C system, the mucosal compartments were filled with 1.5 ml of cold 10 mM mannitol in Krebs buffer and the serosal compartments were filled with 10 mM glucose in Krebs buffer. Both sides were continuously oxygenated, 95% O\(_2\)/5% CO\(_2\), and circulated by gas flow. Each chamber had two pairs of Ag/AgCl electrodes with agar-salt bridges to monitor the transepithelial potential difference, short circuit current, and electrical resistance across the tissue. Tissue conductance, representing passive gas flow. Each chamber had two pairs of Ag/AgCl electrodes with agar-salt bridges to monitor the transepithelial potential difference, short circuit current, and electrical resistance across the tissue. Tissue conductance, representing passive gas flow.

**Macromolecular passage**

For acute stress, 12 VE and 12 FAE segments from six rats per group were mounted in Ussing chambers. For chronic stress, 18 VE and 18 FAE segments from six rats per group were mounted in Ussing chambers. After equilibration, chemically killed fluorescein conjugated *Escherichia coli* K-12 BioParticles (Molecular Probes, Leiden, the Netherlands) were added to the mucosal side of the reservoirs at a final concentration of 1 \(\times\) 10\(^5\) CFU/ml. Bacteria were killed with paraformaldehyde which stops their reproduction but retains antigenicity. After 45, 90, and 120 minutes, the entire volume of the serosal compartments were collected and analysed at 488 nm in a fluorimeter (Cary Eclipse, Varian, Victoria, Australia). Segments were fixed in situ in the chambers with 4% formaldehyde in phosphate buffered saline (PBS) for three hours and stored in the dark at 4°C until further use for confocal microscopy (see below).

**Bacterial passage**

For chronic stress, 18 VE and 18 FAE segments from six rats per group were mounted in Ussing chambers. After equilibration, chemically killed fluorescein conjugated *Escherichia coli* K-12 BioParticles (Molecular Probes, Leiden, the Netherlands) were added to the mucosal side of the reservoirs at a final concentration of 1 \(\times\) 10\(^5\) CFU/ml. Bacteria were killed with paraformaldehyde which stops their reproduction but retains antigenicity. After 45, 90, and 120 minutes, the entire volume of the serosal compartments were collected and analysed at 488 nm in a fluorimeter (Cary Eclipse, Varian, Victoria, Australia). Segments were fixed in situ in the chambers with 4% formaldehyde in phosphate buffered saline (PBS) for three hours and stored in the dark at 4°C until further use for confocal microscopy (see below).

**Corticosterone assay**

Total corticosterone in blood samples from chronic stressed rats was analysed by the IDS OCTEIA corticosterone kit (IDS, Boldon, U.K) according to the manufacturer’s instructions. All samples were run in duplicate and kit calibrators and controls were included in each analysis. Absorbances were measured at 450 nm, with a reference wavelength of 650 nm, in an ELISA microtitre plate reader (Anthos hTII; Anthos, Labtec Instruments, Salzburg, Germany). A calibration curve was prepared on semi-log graph paper and values for each unknown sample were read off the curve in ng/ml.

**HRP assay**

HRP was measured using the QuantaBlu Fluorogenic Peroxidase Substrate Kit (Pierce, Rockford, USA). Samples collected from the serosal side of the chambers were diluted in Krebs buffer with 0.02% bovine serum albumin and 50 \(\mu\)l was transferred to dark microtitre plates. To each well, 100 \(\mu\)l of QuantaBlu working solution (QuantaBlu Stable peroxidase solution diluted 1:10 in QuantaBlu substrate solution) was added and the plate was incubated in a shaker at 300 rpm at room temperature. After exactly 30 minutes, 100 \(\mu\)l of QuantaBlu stop solution was added and 10 minutes later absorbance of the coloured reaction product was measured at an excitation wavelength of 325 nm and emission wavelength of 420 nm using a fluorimeter (Cary Eclipse, Varian, Victoria, Australia). Blanks were included in each analysis and all samples were run in duplicate and measured against a standard curve. Intravariability coefficient was estimated as 8.6% and intervariability coefficient 9.3%.

**Microscopic examination**

**Electron microscopy**

Glutaraldehyde fixed segments were washed in Tris buffer and incubated with 0.6 mg/ml DAB-3% H\(_2\)O\(_2\) for 30 minutes. Tissues were transferred to 0.1 M c Vademy buffer, pH 7.4, with 1% osmium (1:1), and after one hour washed in c Vademy buffer, dehydrated, and embedded in Epon (TAAB 812; Analytical Standards, Gothenburg, Sweden). Sections were stained with lead citrate and examined by transmission electron microscopy (JEM-1230; JEOL Sollentuna, Sweden).
Confocal microscopy

Tissues were rinsed in PBS and labelled with 5 µg/ml wheat germ agglutinin conjugated to Alexa Flour 594 (Molecular Probes) to visualise the epithelial cell surface. After repeated washing in PBS, tissues were mounted in glycerol and examined in a Nikon Eclipse E600W confocal laser scanning microscope (Nikon, NY, USA) with a 60× oil immersion objective. An ion laser permitted simultaneous excitation wavelengths of 488 nm for fluorescein labelled E. coli and 594 nm for Alexa labelled wheat germ agglutinin.

Light microscopy

Segments of VE from three randomly selected rats per group were frozen in liquid nitrogen, embedded in mounting medium, and cryosectioned at 3 µm. Coded slides were fixed in acetone for 10 minutes and stained with May-Grünwald/Giemsa. Cells identified as mast cells were counted in three tissues per rat using Easy Image software (Tekno Optik AB, Stockholm, Sweden) connected to a Nikon Eclipse 800 light microscope with a 60× oil immersion objective. For each tissue, eight non-overlapping areas (37 mm²) above the muscularis mucosa were examined.

Statistics

Parametric values between two groups were performed using the Student’s t test, and between more than two groups using one way ANOVA. Non-parametric values are given as median (SEM). Comparisons between two groups were done using the Mann-Whitney U test and between more than two groups with the Kruskal-Wallis one way ANOVA. Differences with p < 0.05 were considered significant.

RESULTS

FAE is responsive to acute stress

Acute stress significantly increased the frequency of defecation (12.3 (2.3) v controls 1.8 (0.8) pellets/h; p<0.0005). Moreover, acute stress exposure significantly increased the short circuit current and electrical conductance in both VE and FAE (fig 1A). There were no changes in 51Cr-EDTA permeability after stress exposure in VE (4.3 (0.7) v controls 4.5 (0.9) 10⁻⁶ cm/s) or FAE (8.0 (1.6) v 7.0 (0.8) cm/s). However, HRP flux was increased in both VE (15.1 (3.4) v controls 11.1 (6.6) pmol/h/cm²) and FAE (36.9 (10.7) v 21.8 (6.7) pmol/h/cm²). FAE was significantly more permeable than VE to both probes.

Changes in systemic parameters and electrophysiology by chronic stress

During the experimental period, stressed rats stagnated in weight (mean weight was 293 ± 6 g on day 1 and 298 ± 9 g on day 10) whereas control rats continued to gain weight (mean weight was 287 ± 6 g on day 1 and 324 ± 7 g on day 10). The difference in weight gain between controls and stressed rats was significant from day 4 of the stress period (p<0.05). Chronic stress significantly increased the defecation rate at each stress session, and the rate was consistently increased for each daily session over the 10 day period. The mean number of faecal pellets expelled daily over the period was 9.9 (0.3) pellets/h (range 7–15) in stressed rats and 2.4 (0.2) (range 0–4) in controls (p<0.0005). Serum corticosterone levels were elevated after 10 days of stress (215.4 (20.0) v controls 123.0 (9.8) ng/ml; p<0.0005). Furthermore, chronic stress exposure almost tripled the number of ileal mucosal mast cells (68 (3.0) v controls 23 (1.7) cells/mm²; p<0.0005), and electron microscopy showed signs of mast cell degranulation in stressed rats. Rats exposed to chronic stress showed...
a significant increase in short circuit current in both VE and FAE whereas electrical conductance was significantly increased in VE but not in FAE (fig 1B).

**Pronounced chronic stress induced increase in FAE uptake of HRP**
Permeability to $^{51}$Cr-EDTA was increased in VE after stress exposure ($8.1 \pm 0.8$ vs controls $6.1 \pm 0.7\times10^{-6}$ cm/s; $p<0.05$) whereas there was no significant change in FAE ($7.8 \pm 1.0$ vs $7.2 \pm 0.8$; $p = 0.08$) (fig 2A). Chronic stress enhanced HRP flux in VE (24.3 (4.3) vs controls 13.6 (2.5) pmol/h/cm$^2$; $p<0.05$) and caused a pronounced increase in HRP flux in FAE (92.5 (20.2) vs 25.5 (4.8); $p<0.005$) (fig 2B).

**Ultrastructural changes in FAE by chronic stress**
FAE from control rats demonstrated a normal morphology whereas stressed animals in some areas revealed abnormal epithelial cell morphology with vacuoles (fig 3A, B). Uptake of HRP was observed in all segments investigated, and occasionally bacteria penetrating into enterocytes were seen. Within the FAE, M cells with uptake of HRP in direct contact with lymphocytes were identified between enterocytes (fig 4A, B). M cells were identified according to definitions for identification previously established $^{12-4}$: irregularly shaped apical surface, short microvilli, attenuated cytoplasm, invaginated basolateral membrane with pockets containing lymphocytes and macrophages, and prominent vesicles.

**Chronic stress induced bacterial internalisation via the FAE**
Examination by electron microscopy illustrated signs of bacterial penetration into the enterocytes of the FAE of stressed rats. To further explore the effects of stress on bacterial passage, uptake was quantified by Ussing chamber experiments (table 1). After 45 minutes, there were no significant differences in *E. coli* passage between FAE from stressed rats and controls. After 90 minutes, passage was slightly elevated in stressed rats and after 120 minutes it was increased more than 30-fold. Bacterial uptake was confirmed by confocal microscopy where it was shown that after 45 minutes fluorescent bacteria were observed close to the epithelial surface, after 90 minutes bacteria were found penetrating the epithelium, and after 120 minutes bacteria had passed through the entire epithelial cell layer (fig 5A, B, C). In VE, *E. coli* flux was only slightly elevated after 120 minutes in stress exposed rats (table 1).

**DISCUSSION**
To our knowledge, this is the first report of a stress induced increase in transcellular permeability of FAE. There was a greater than threefold increase in permeability to the antigenic protein HRP in FAE after chronic stress. Moreover, macromolecular uptake was more than three times higher in FAE than in VE of the same stressed animal.

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**Figure 2**
Effects of chronic stress on permeability. Rats were subjected to sham stress or water avoidance stress for 10 consecutive days, 12 rats/group. Ileal segments from villus epithelium (VE) and follicle associated epithelium (FAE) were mounted in Ussing chambers. (A) $^{51}$Cr-EDTA flux in VE and FAE. (B) Horseradish peroxidase flux (HRP) in VE and FAE. Bars represent mean (SEM). Comparisons were done using ANOVA.

**Figure 3**
Effects of chronic stress on epithelial ultrastructure. (A) Follicle associated epithelium (FAE) from control rats demonstrated a normal morphology. (B) FAE from stressed rats revealed an abnormal morphology with numerous vacuoles. Arrows indicate endosomal uptake of horseradish peroxidase into epithelial cells.
Enhanced uptake in FAE was further emphasised by increased passage of chemically killed *E. coli* in rats exposed to chronic stress. Under normal conditions, the HRP molecule passes through epithelial cells via pinocytosis, although after chronic restraint stress and in sensitised animals HRP can also permeate between cells. In the present study, paracellular permeability to ions (conductance) was similarly affected in VE and FAE after stress. After chronic stress, permeability to the paracellular marker molecule 51Cr-EDTA was increased in VE but almost unaffected in FAE. In contrast, stress induced changes in permeability to proteins was significantly more pronounced in FAE, with an almost fourfold increase in HRP flux. Electron microscopy revealed that HRP was transported in endosomes via the transcellular pathway whereas no HRP staining was found in the paracellular spaces. Taken together, these findings suggest that chronic stress, in the FAE, mainly affects transcellular antigen uptake. Ultrastructural examination further revealed penetrating bacteria in FAE of stressed rats which hinted at increased uptake of bacteria. Subsequent Ussing chamber studies demonstrated that permeability to *E. coli* was more than 30 times higher in the FAE of stressed rats compared with controls, and six times higher in FAE than in VE. Although the *E. coli* strain used in our study was chemically killed, lost motility, and could not replicate, they preserved their antigenicity. These particles have been used for phagocytosis studies and should be a good model of bacterial invasion. With confocal microscopy, it was seen that after 45 minutes, bacteria were present on the epithelial surface, and after 120 minutes, bacteria had passed through the epithelium. Increased uptake of bacteria by the FAE reflects a region made permeable to normally non-invasive bacteria. It may be speculated that a significant part of the bacterial translocation that takes part in various stress reactions may be due to invasion via the FAE, as revealed by our studies. As the earliest signs of CD are aphthoid ulcers in the FAE, barrier dysfunction of the FAE may be involved in the early stages of this disease.

In VE, conductance was, as in previous studies, enhanced by both acute and chronic stress. In contrast, permeability to the paracellular probe 51Cr-EDTA was increased only by chronic stress. This finding suggests partially different mechanisms involved in the paracellular permeability defect induced by acute and chronic stress. Recently, Ferrier et al. found that in vivo permeability to 51Cr-EDTA in the mouse colon was unaffected by a single stress episode whereas repeated stress over 3–4 days induced a pronounced increase in paracellular permeability. This barrier defect was mediated by T cells and interferon γ. It could be speculated that this T cell activation, leading to interferon γ production and eventually increased paracellular permeability to macromolecules, is initiated by a WAS induced increase in transcellular permeability to antigens. This increase in transcellular permeability may involve mast cells. The importance of mast cells in stress induced mucosal abnormalities have been demonstrated previously. Direct contact between gastrointestinal mast cells and nerves has been widely discussed, and it is known that in response to stress, mast cells secrete mediators that affect nerves and immune cells. Direct contact between gastrointestinal mast cells and nerves has also been reported in the small intestine, but whether the altered FAE function seen in the present study involves nerve-mast cell interaction

### Table 1 Effects of chronic stress on bacterial passage

<table>
<thead>
<tr>
<th>Group, tissue</th>
<th>45 min passage</th>
<th>90 min passage</th>
<th>120 min passage</th>
</tr>
</thead>
<tbody>
<tr>
<td>WAS, FAE</td>
<td>0.6 (0.3–1.2)</td>
<td>4.3 (2.0–5.9)</td>
<td>47.9 (17.6–94.7)*†</td>
</tr>
<tr>
<td>SS, FAE</td>
<td>0.0 (0.0–0.0)</td>
<td>1.8 (1.2–2.2)</td>
<td>1.4 (0.9–5.1)</td>
</tr>
<tr>
<td>WAS, VE</td>
<td>0.4 (0.1–0.7)</td>
<td>2.2 (1.5–7.3)</td>
<td>7.8 (6.4–19.3)‡</td>
</tr>
<tr>
<td>SS, VE</td>
<td>0.6 (0.4–1.4)</td>
<td>3.9 (3.6–4.2)</td>
<td>5.0 (3.0–5.8)</td>
</tr>
</tbody>
</table>

Rats were subjected to water avoidance stress (WAS) or sham stress (SS) for 10 days, 1 h/day, 6 rats/group. Segments of villus epithelium (VE) and follicle associated epithelium (FAE) were mounted in Ussing chambers. Passage of chemically killed *E. coli* K-12 was studied over time. Values are presented as median (25–75th interquartile range). Comparisons between groups were performed using Kruskal-Wallis one way ANOVA: *p* < 0.005 versus SS FAE, †p < 0.05 versus WAS VE; ‡p < 0.05 versus SS VE.
Stress and antigen uptake

remains to be elucidated. Nevertheless, the responsiveness of FAE to acute stress suggests neurally mediated effects on FAE permeability. However, another possibility could be corticosterone mediated effects on barrier function which has been shown in vivo in severe acute stress.12

The present study clearly demonstrates the importance of FAE in the uptake of luminal antigens. Its superior uptake capacity compared with VE is partly due to M cells that constitute approximately 10% of rat FAE.33 M cells lack a developed microvillus border and glycocalyx coat which facilitates access for luminal pathogens.2 Previous studies showed that M cell numbers were altered in bacterial enteritis,14,15 indomethacin induced ileitis,15 and in patients with spondylarthropathy.15 These studies indicate that M cells may play a role in the development of intestinal inflammation. The results of the present study may suggest that stress alters the function of M cells in the FAE. M cells meeting the criteria for identification14 were found in the FAE by electron microscopy. The close interaction between antigen, M cells, and immune cells was visualised and most likely this interaction is of major importance in the alteration of FAE function. However, more analyses are needed to determine whether the alteration involves changes in M cell number or activity, a problem superseded by the lack of reliable rat M cell markers. Cytokeratin 8 (clone 4.1.18) has been suggested as a problem superseded by the lack of reliable rat M cell markers. Cytokeratin 8 (clone 4.1.18) has been suggested as a cell marker. Cytokeratin 8 (clone 4.1.18) has been suggested as a marker. Cytokeratin 8 (clone 4.1.18) has been suggested as a marker.


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