Glucagon-like peptide-2 enhances intestinal epithelial barrier function of both transcellular and paracellular pathways in the mouse

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Abstract
Background and aims—Glucagon-like peptide-2 (GLP-2) is a recently identified potent intestinotrophic factor. We have evaluated the effect of GLP-2 treatment on intestinal epithelial barrier function in mice.

Methods—CD-1 mice were injected subcutaneously with GLP-2 or a protease resistant analogue, h[GLy2]GLP-2, twice daily for up to 10 days. Saline injected mice served as controls. Jejunal segments were mounted in Ussing chambers. Tissue conductance was measured and unidirectional fluxes were determined for (i) Na⁺ and the small inert probe Cr-EDTA (both transported via the paracellular pathway) and (ii) the macromolecule horseradish peroxidase (HRP, transported via the transcellular pathway).

Results—Mice treated with GLP-2 or h[GLy2]GLP-2 for 10 days demonstrated significantly reduced intestinal conductance and fluxes of Na⁺, Cr-EDTA, and HRP. Electron microscopy confirmed that GLP-2 reduced endocytic uptake of HRP into enterocytes. Functional changes (evident by four hours) preceded morphological changes (evident by 48 hours).

Conclusions—GLP-2 enhances intestinal epithelial barrier function by affecting both paracellular and transcellular pathways and thus may be of therapeutic value in a number of gastrointestinal conditions.

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Keywords: intestinal permeability; macromolecular transport; growth factors

The gastrointestinal tract is lined by a single cell layer of epithelium in which adjacent cells are joined together at their apical poles by tight junctions. This epithelial lining performs a critical barrier function to limit uptake of luminal antigens and noxious substances into the body. Epithelial permeability is increased in a number of enteropathies, such as inflammatory bowel disease, coeliac disease, and food allergies (reviewed by Bjarnason and colleagues,1 Meddings,2 and Sartor3). Indeed, it has been suggested that a primary defect in the barrier function of the gastrointestinal epithelial layer may be a predisposing factor in the pathogenesis of Crohn’s disease.4 Although this hypothesis remains to be confirmed, it is likely that excessive uptake of luminal antigens and/or bacterial products into the mucosa may trigger inflammatory reactions with release of cytokines and bioactive mediators resulting in intestinal damage and dysfunction.

Animal models and epithelial cell culture studies have demonstrated that many factors are capable of increasing intestinal epithelial permeability. Such factors include the nutritional status of the host,5 certain medications,6 stress,7 bacterial infections and/or toxins8-10 and immune cell cytokines (for example interferon γ,11 tumour necrosis factor α,12 13 and interleukin 414 15). Although it is clear that a number of stimuli can elicit greater gastrointestinal permeability, less is known about factors that maintain or enhance the barrier function of the intestinal epithelium. Growth factors, including transforming growth factor β16 17 insulin-like growth factor-1,18 and epidermal growth factor19 20 are capable of maintaining or restoring intestinal mucosal integrity thereby fostering healing and restitution. However, most growth factors are not organ specific and induce additional non-specific changes that can affect homeostasis.

In 1971, Gleeson et al described a glucagon secreting endocrine tumour capable of inducing intestinal hyperplasia.21 More recent investigations have shown that a cleavage product of the molecule, designated glucagon-like peptide-2 (GLP-2), is a trophic factor specific for the bowel. Thus in mice, twice daily treatment with low doses of GLP-2 resulted in increased gut weight and villus height in the small intestine but did not affect the thickness of the smooth muscle layer.22 Concomitant with the change in gut architecture was an increase in disaccharidase levels and increased nutrient absorption.23

Despite the recognition that GLP-2 stimulates small intestinal growth, the relationship between this novel growth factor and intestinal barrier function has not been reported. Therefore, the primary aim of this study was to determine the ability of GLP-2 and a protease resistant analogue to regulate the barrier function of the murine small intestine. Our data illustrate that GLP-2 significantly reduces paracellular transport of ions and small molecules, and moreover dramatically inhibits endocytic uptake of macromolecules. These findings support the postulate that GLP-2 treatment may be an effective therapy for...
GLP-2 enhances barrier function

Methods

ANIMALS
Female CD-1 mice (eight weeks of age, mean weight 50 g) were obtained from Harlan Sprague-Dawley Inc. (Indianapolis, Indiana, USA). Mice were not restricted by diet or activity during the experiment. After one week of acclimatisation, mice were matched by weight and allocated to one of three groups. Mice were injected subcutaneously twice daily for 10 consecutive days with 5 µg of either native human GLP-2 or a synthetic protease resistant human analogue h[Gly-NH2]GLP-2 (donated by Allelix Biopharmaceuticals Inc., Mississauga, Ontario, Canada). This dose was chosen based on pilot studies with native GLP-2 that showed it produced consistent, significantly increased mucosal growth. Controls were injected with the vehicle (phosphate buffered saline (PBS)). Experiments were conducted on day 11 when mice were weighed and sacrificed by cervical dislocation. The small intestine was removed, flushed with cold saline, blot dried, and weighed. The jejunum was divided into segments for morphometric assessment, protein determination, and Ussing chamber studies. To obtain information on the timing of effects, additional experiments were conducted in mice injected once with h[Gly-NH2]GLP-2 and studied at four hours and in mice injected twice a day for two days and studied at 48 hours. The studies described in this paper were conducted with approval from the McMaster University Animal Care Committee.

MORPHOLOGY

Light microscopy
A 1 cm segment of intestine (~10 cm from the pylorus) was opened along the mesenteric border and immersion flat fixed in 10% neutral buffered formalin. After dehydration and wax embedding, tissue sections were stained with haematoxylin and eosin. Mucosal height, villus length, and crypt depth were measured in coded sections (8–10 intact villus-crypt units per mouse) by a single investigator (MAB), who was unaware of the mouse treatment, using a calibrated eyepiece. The villus:crypt ratio was subsequently calculated.

Electron microscopy
An additional 1 cm segment (distal to that taken for light microscopy) was immediately fixed with 2% glutaraldehyde in 0.1 mol/l sodium cacodylate buffer for two hours at room temperature, transferred to sodium cacodylate buffer, and stored at 4°C overnight. Tissues were subsequently processed for routine electron microscopy. Photomicrographs of well orientated epithelial cells from the mid villus region were prepared. Using a computer supported image analysis system (Mop Videoplan, Kontron, Germany) one investigator (PCY), who was unaware of the mouse treatment, analysed 12 coded photomicrographs per group and measuredenterocyte width and length, and microvillus length.

MUCOSAL PROTEIN

A 7 cm segment of small intestine (proximal to that used for morphology) was cut and the mucosal layer was scraped off between two glass slides. The mucosal scraping was weighed and buffer was added at a concentration of 10 ml/g tissue. The mucosal suspension was homogenised on ice for 30 seconds, snap frozen in liquid nitrogen, and stored at −20°C before determination of protein content.

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A 12 cm segment of small intestine (distal to that used for morphology) was opened along the mesenteric border and cut into four flat sheets. Full thickness segments (devoid of Peyers patches) were mounted in modified Ussing chambers and bathed with oxygenated Krebs buffer that contained (in mM): 115 NaCl, 1.25 CaCl2, 1.2 MgCl2, 2.0 KH2PO4, and 25 NaHCO3, pH 7.35±0.02, at 37°C. In addition, the serosal buffer contained 10 mM glucose as an energy source balanced by 10 mM mannitol in the mucosal buffer. Glucose was not added to the mucosal buffer as we were interested specifically in permeability rather than glucose linked ion transport. Tissues were short circuited at zero volts using a World Precision Instruments automated voltage clamp (Narco Scientific, Mississauga, Ontario, Canada). Conductance (G) was calculated according to Ohm’s law using potential difference and short circuit current (Isc) values. Tissues with abnormally high initial conductance values (>40 nS/cm2) or whose conductance increased during the course of the experiment (calculated every 15 minutes) were considered damaged and were excluded from subsequent analysis.

(a) Flux of Na+

The unidirectional serosal-to-mucosal flux of Na+ was determined as an indication of passive ion permeability via the paracellular pathway. Flux was measured under voltage clamped conditions by adding 10 µCi of 22Na (New England Nuclear, Lachine, Quebec, Canada) to the serosal buffer. Following a 20 minute equilibration period, 1 ml samples from the “cold” buffer and 50 µl samples from the “hot” buffer were collected and replaced with the appropriate volume of buffer every 15 minutes. Radioactivity was measured in a gamma counter and flux (in µmol/cm2/h) was calculated.

(b) Flux of Cr-EDTA

Cr-EDTA is generally considered to be a marker of paracellular permeability. Transepithelial flux of this small (360 Da) inert probe was determined as described previously using the same approach as that for the radiolabelled ion. Cr-EDTA (6 µCi/ml, Radiopharmacy, Chedoke-McMaster Hospital, Hamilton, Ontario, Canada) was added to the mucosal buffer bathing jejunal segments mounted in Ussing chambers. Non-radioactive Cr-EDTA was
added to the serosal buffer at an equal concentration. Samples were obtained as above, counted, and flux expressed in nmol/cm²/h.

(c) Flux of horseradish peroxidase (HRP)
The transcellular passage of HRP (44 000 Da), a model protein antigen, was determined by flux and electron microscopy (see below) as previously described. Type VI HRP (Sigma-Aldrich Canada Inc., Oakville, Ontario, Canada) was added to the mucosal buffer at a concentration of 10⁻⁵ M. Samples were obtained at intervals as above. The sample (appropriately diluted) was then mixed with a reaction solution containing hydrogen peroxide and 4-dianisidine (Sigma-Aldrich Canada Inc.), and the rate of appearance of the reaction product was determined at 460 nm. Flux was expressed as pmol/cm²/h.

(d) Quantification of HRP containing endosomes in enterocytes.
Following exposure of the intestine in Ussing chambers to luminal HRP for 60 minutes, the segment was removed and immersed in 2% glutaraldehyde in 0.1 mol/l sodium cacodylate buffer for two hours at room temperature, transferred to sodium cacodylate buffer, and stored at 4°C overnight. The tissue was washed three times in 0.05 mol/l Tris buffer and then incubated for 30 minutes in 5 mg of 3,3′-diaminobenzidine tetrahydrochloro (Sigma-Aldrich Canada Inc.) in 10 ml of 0.05 mol/l Tris buffer and 0.1% hydrogen peroxide. The tissue was subsequently processed for electron microscopy and photomicrographs prepared of well oriented epithelial cells. The area occupied by HRP product containing endosomes located within a 112 µm² window in the apical region of the enterocyte above the nucleus was measured in coded photomicrographs by one of the investigators (PCY) using a computer supported image analysis system. A total of 12 windows for each of four mice per group (48 enterocytes) were analysed.

(e) Effect of in vitro addition of GLP-2.
GLP-2 (5–1000 nM) was added to the luminal or serosal side of intestinal segments from naïve mice and GLP-2 treated mice. As a consequence of villus elongation, there was a shift in the villus:crypt ratio that was statistically significant (p<0.05), from 5.12 (0.37) (control) to 6.18 (0.31) or 7.23 (0.41) for treatment with GLP-2 or h[Gly²]GLP-2, respectively (n=7–8 mice/group). Electron microscopy demonstrated that GLP-2 or h[Gly²]GLP-2 treatment changed the shape of the epithelial cells causing them to become narrower and significantly (p<0.05) more effective than GLP-2.

Figure 1B illustrates that the weight of mucosal scrapings was increased by 33% or 50% following treatment with GLP-2 or h[Gly²]GLP-2, respectively. In addition, GLP-2 or h[Gly²]GLP-2 resulted in a significant (p<0.001) increase in mucosal protein compared with control values (from 26.5 (2.2) to 42.4 (1.7) or 43.9 (2.4) mg, respectively; n=7–8 mice/group). Morphometric analysis of histological sections revealed that GLP-2 treatment of mice for 10 days increased both the total mucosal thickness and villus height by approximately 25% compared with control values (table 1). Neither experimental treatment caused any consistent or significant change in crypt depth. As a consequence of villus elongation, there was a shift in the villus:crypt ratio that was statistically significant (p<0.05), from 5.12 (0.37) (control) to 6.18 (0.31) or 7.23 (0.41) for treatment with GLP-2 or h[Gly²]GLP-2, respectively (n=7–8 mice/group). Electron microscopy demonstrated that GLP-2 or h[Gly²]GLP-2 treatment changed the shape of the epithelial cells causing them to become narrower and
longer than control enterocytes (table 1; fig 2A, B). Microvilli were also longer (table 1; fig 2C, D). Again, h[Gly2]GLP-2 was more effective than GLP-2 in altering enterocyte morphology.

Table 1  Effect of GLP-2 or h[Gly2]GLP-2 treatment for 10 days on morphology

<table>
<thead>
<tr>
<th>Mouse treatment</th>
<th>Mucosal height (µm)</th>
<th>Villus length (µm)</th>
<th>Crypt depth (µm)</th>
<th>Microvillus length (µm)</th>
<th>Enterocyte width (µm)</th>
<th>Enterocyte length (µm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PBS</td>
<td>653 (33)</td>
<td>541 (31)</td>
<td>114 (5)</td>
<td>1.20 (0.08)</td>
<td>5.17 (0.20)</td>
<td>17.57 (0.47)</td>
</tr>
<tr>
<td>GLP-2</td>
<td>816 (26)*</td>
<td>688 (28)*</td>
<td>121 (6)</td>
<td>2.27 (0.07)*</td>
<td>4.83 (0.22)</td>
<td>29.00 (0.43)*</td>
</tr>
<tr>
<td>h[Gly2]GLP-2</td>
<td>880 (22)*</td>
<td>762 (22)*</td>
<td>118 (6)</td>
<td>2.31 (0.11)*†</td>
<td>4.19 (0.16)*†</td>
<td>32.75 (0.40)*†</td>
</tr>
</tbody>
</table>

Mice were injected daily with 5 µg of GLP-2 or its analogue (h[Gly2]GLP-2) for 10 days. All measurements are expressed in µm. Values are mean (SEM); n=7-8 mice per group. EM measurements were made on 12 photomicrographs per group.

*p<0.05 compared with phosphate buffered saline (PBS) treated mice (control); †p<0.05 compared with GLP-2 treated mice.

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Paracellular permeability

Treatment of mice with either GLP-2 or h[Gly2]GLP-2 for 10 days significantly decreased paracellular permeability.

Figure 2  Representative photomicrographs of murine intestinal enterocytes. (A) and (C) are from phosphate buffered saline (PBS) treated mice (control); (B) and (D) are from mice after 10 days of treatment with h[Gly2]GLP-2 (5 µg). Note the longer microvilli in h[Gly2]GLP-2 treated mice (B) compared with controls (A) (bar=200 µm). The arrowhead in (B) indicates an endosome containing HRP. Note the increased cell length in h[Gly2]GLP-2 treated mice (D) compared with controls (C) (bar=100 µm).
creased the conductance of jejunal tissues by approximately 25% (p<0.05) or 45% (p<0.01), respectively (fig 3A).

Isc also tended to be lower in treated mice although values were not significantly different (38.2 (2.0) in tissues from controls v 34.1 (2.6) in tissues from GLP-2 treated mice and 32.7 (2.1) in tissues from h[Gly2]GLP-2 treated mice; n=14–31 tissues/group). The flux of the small inert probe, Cr-EDTA, was significantly reduced by approximately 35% (p<0.05) across the intestine in mice treated with GLP-2 or h[Gly2]GLP-2 compared with intestinal tissue from control mice (fig 3B). In addition, h[Gly2]GLP-2 peptide treatment reduced the serosal-to-mucosal flux of Na+ by approximately 35% (p<0.05) from 11.5 (0.8) to 7.7 (0.6) µmol/h/cm² (n=8 tissues/group). Taken together, these findings indicate a reduced ability of ions and small molecules to permeate through the paracellular pathway.

Acute addition of GLP-2 (5–1000 nM) to tissues in chambers produced no significant change in conductance or Isc within 90 minutes compared with control tissues to which vehicle was added (G values ranged from 20.2 (1.7) to 24.2 (1.9) mS/cm² and Isc from 34.5 (2.1) to 37.7 (2.2) µA/cm²; n=8–10 tissues/group).

Transcellular permeability

The flux of HRP across mouse intestine was significantly (p<0.05) reduced by approximately 50% from control values (fig 4A). An even more dramatic decrease (p<0.001) in HRP transport (∼80%) was induced by treating mice with h[Gly2]GLP-2. Electron microscopy demonstrated that the pathway for HRP transport was transcellular in that HRP was evident within endosomes but not in the paracellular regions (fig 2B). There was a significant diminution (>50%) in the area of HRP containing endosomes in jejunal enterocytes of both GLP-2 and h[Gly2]GLP-2 treated mice compared with control values (fig 4B). Again, h[Gly2]GLP-2 was significantly (p<0.05) more effective than GLP-2.

Table 2 Effect of h[Gly2]GLP-2 treatment for four or 48 hours on intestinal weight and morphology

<table>
<thead>
<tr>
<th>Mouse treatment</th>
<th>Gut weight (g)</th>
<th>Mucosal protein (mg)</th>
<th>Villus height (mm)</th>
<th>Enteroocyte width (mm)</th>
<th>Enteroocyte length (mm)</th>
<th>MV length (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PBS</td>
<td>1.81 (0.05)</td>
<td>26.4 (2.0)</td>
<td>521 (25)</td>
<td>5.6 (0.2)</td>
<td>20.6 (0.8)</td>
<td>1.15 (0.05)</td>
</tr>
<tr>
<td>h[Gly2]GLP-2 (4 h)</td>
<td>1.80 (0.03)</td>
<td>27.2 (1.7)</td>
<td>512 (14)</td>
<td>5.5 (0.2)</td>
<td>23.4 (0.6)</td>
<td>1.15 (0.06)</td>
</tr>
<tr>
<td>h[Gly2]GLP-2 (48 h)</td>
<td>1.76 (0.06)</td>
<td>28.4 (3.0)</td>
<td>530 (17)</td>
<td>5.0 (0.2)</td>
<td>31.6 (0.9)*</td>
<td>1.60 (0.06)*</td>
</tr>
</tbody>
</table>

Mice were injected either once (four hour group) or daily for two days (48 hour group) with 5 µg of h[Gly2]GLP-2. Values are mean (SEM); n=5 mice per group. EM measurements were made on 12 photomicrographs per group; MV, microvilli. *p<0.05 compared with phosphate buffered saline (PBS) treated mice (control).
Table 3  Effect of h[Gly2]GLP-2 treatment for four or 48 hours on intestinal permeability parameters

<table>
<thead>
<tr>
<th>Mouse treatment</th>
<th>G (mS/cm²)</th>
<th>n</th>
<th>Cr-EDTA flux (nmol/h/cm²)</th>
<th>n</th>
<th>HRP flux (pmol/h/cm²)</th>
<th>n</th>
</tr>
</thead>
<tbody>
<tr>
<td>PBS</td>
<td>29.8 (0.8)</td>
<td>81</td>
<td>15.5 (1.1)</td>
<td>31</td>
<td>3.8 (0.5)</td>
<td>33</td>
</tr>
<tr>
<td>h[Gly2]GLP-2 (4 h)</td>
<td>25.7 (1.7)*</td>
<td>23</td>
<td>10.2 (0.4)*</td>
<td>9</td>
<td>1.6 (0.2)**</td>
<td>13</td>
</tr>
<tr>
<td>h[Gly2]GLP-2 (48 h)</td>
<td>25.7 (1.4)*</td>
<td>20</td>
<td>11.7 (0.8)*</td>
<td>11</td>
<td>1.3 (0.4)**</td>
<td>12</td>
</tr>
</tbody>
</table>

Mice were injected either once (four hour group) or daily for two days (48 hour group) with 5 μg of h[Gly2]GLP-2 or 33 phosphate buffered saline (PBS) treated mice (control) (combined from all experiments).

*p<0.05, **p<0.01 compared with PBS treated control mice.

**Discussion**

Unimpeded entry of antigen into the mucosa can result in an inflammatory or immune cascade that culminates in major tissue damage and functional abnormalities. The epithelial lining of the intestine effectively acts as a regulated physiological sieve, maintaining the integrity of the barrier, and thereby acting as a first line of defence. In the present study we demonstrated for the first time that GLP-2 enhanced intestinal barrier function, as demonstrated by several parameters. Conductance, a parameter responsive than mice to GLP-2 because of degradation by dipeptidyl peptidase IV.37

Gut segments from mice treated with GLP-2 or h[Gly2]GLP-2 for 10 days displayed enhanced intestinal barrier function, as demonstrated by several parameters. Conductance, under voltage clamped conditions, indicates the passive movement of ions across the epithelium mainly via the paracellular pathway. Thus the reduced conductance of tissues from GLP-2 or h[Gly2]GLP-2 treated mice indicated that the passive flow of ions had been retarded, and this was confirmed by data illustrating reduced serosal-to-mucosal flux of Na⁺ of approximately 35%, a non-active process related mainly to the paracellular pathway.

Under the conditions of our experiment, Jsc was similar in both controls and GLP-2 treated mice. This result is not inconsistent with previous findings of enhanced expression of glucose transporters as in our study the mucosal buffer did not contain glucose to stimulate Na⁺ absorption via SGLT1 (which would increase Jsc).

Decreased ionomic permeability does not necessarily indicate similar permeability changes for larger molecules. Consequently, we examined the mucosal-to-serosal flux of Cr-EDTA to assess the effect of GLP-2 on the uptake of luminaly derived material. Because of its small size (molecular weight 360) and lipid insolubility, Cr-EDTA has been used extensively to assess paracellular permeability in patients with intestinal disorders.38 We observed a similar reduction in the flux of Cr-EDTA (about
35%) across Ussing chambered-intestinal segments from mice treated for 10 days with either GLP-2 or h[Gly²]GLP-2. Increased length of a tighter/narrower paracellular pathway is one possible explanation for the decreased permeability after GLP-2 treatment.

In another series of experiments we tested the ability of GLP-2 to affect transepithelial movement of macromolecules, choosing HRP as a model protein. HRP is an ideal macromolecular marker as: (1) it is a 44 kDa glycoprotein that under normal circumstances traverses the epithelium via transcytosis; and (2) intact HRP can be conveniently quantified. V-ect transepithelial flux of HRP across jejunum from GLP-2 or h[Gly²]GLP-2 treated mice compared with controls, with h[Gly²]GLP-2 being more effective than GLP-2. A 50% reduction in the area of HRP containing endosomes within enterocytes confirmed that peptide treatment decreased epithelial uptake of this macromolecule. It is possible that the increased length of the cells was responsible for a decreased rate of movement of this large molecule across the tissues. In addition, a longer period for transcytosis would provide increased opportunity for degradation of the protein by lysosomal enzymes.

To determine if the enhanced barrier function was due simply to an increased number of longer and more densely packed enterocytes, additional experiments were carried out in mice treated with h[Gly²]GLP-2 for four and 48 hours when, theoretically, epithelial cell growth would be negligible or minimal. Indeed, we found that there were no significant changes in any of the morphological features at four hours although both enterocyte length and microvillus length were significantly increased by 48 hours. In contrast, tissue conductance was significantly reduced at both four and 48 hours, and the flux of Cr-EDTA was also significantly decreased at both early times. These data indicate a discrepancy between the change in the length of the paracellular pathway and its permeability, and suggest that GLP-2 acts on elements involved in maintaining the integrity of epithelial tight junctions. Furthermore, the flux of HRP was more dramatically diminished (by >50% at four hours) indicating the ability of h[Gly²]GLP-2 to rapidly influence epithelial cell function. As four hours was the earliest time point we examined, we do not know if effects may occur even earlier. However, direct addition of GLP-2 to tissues in Ussing chambers did not result in any significant changes in Isc or conductance within 90 minutes. Collectively, our data imply that GLP-2 and h[Gly²]GLP-2 rapidly and effectively reduce paracellular permeability in the small intestine and so would be expected to retard the uptake of small, luminally derived proinflammatory molecules of comparable size into the mucosa. In addition, decreased uptake of HRP suggests that GLP-2 would reduce uptake of foreign protein antigens with immunogenic properties.

Thus our data have shown that GLP-2 and h[Gly²]GLP-2 are not only growth promoting agents for intestinal mucosa but also enhance gut epithelial barrier function. This latter finding has clear implications for innate immunity.

While the cellular and molecular mechanisms underlying these properties of GLP-2 remain to be defined, a number of scenarios can be postulated. Firstly, GLP-2 increases the rate of crypt cell proliferation and reduces apoptosis,28 and one group has reported29 that crypt cell apoptosis leads to increased permeability in murine epithelial cells. Modulation of tight junction form and function would affect paracellular permeability. Secondly, villus elongation could physically restrict access of the full thickness intestinal segments used in this study. Our data with Cr-EDTA and HRP suggest that GLP-2 treatment affects both paracellular and transcellular permeability and effectively re-rates the entry of bioactive immunogenic molecules, of a range of sizes, into the mucosa. Therefore, the recently reported beneficial effects of h[Gly²]GLP-2 in a mouse model of colitis30 may be due, at least in part, to enhanced barrier function.

In summary, we have provided clear evidence that GLP-2 and a protease resistant analogue enhance the barrier function of the intestinal epithelium. By virtue of its intestinotrophic properties, therapeutic uses for GLP-2 have been suggested for conditions that result in villus atrophy such as short bowel syndrome37 and consequent to total parenteral nutrition.38 Furthermore, the research findings presented here underscore the potential role of GLP-2 therapy in combating inflammatory disorders that are characterised by disruption of the normal barrier function of the epithelial lining of the small intestine.

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GLP-2 enhances barrier function