Pathological effects of *Phaseolus vulgaris* isolectins on pig jejunal mucosa in organ culture

M J L Kik, J F J G Koninkx, A van den Muysenberg, F Hendriksen

**Abstract**

The interaction of plant lectins with pig small intestinal epithelium in organ culture was studied. The binding of *Phaseolus vulgaris* (PHA) isolectins E4 and L4 to the microvilli and microvillous vesicles in the top area of the villi was shown by immunoelectron microscopy. Differences were observed in the distribution of the isolectins. In the explants cultured for five hours with the PHA isolectins, the enterocyte height and the villus length were decreased, and a lower villus: crypt ratio was calculated. Ultrastructurally, the microvilli were shorter and irregularly positioned. After incubation with both PHA E4 and PHA L4, clusters of small vesicles, tied off from the microvilli, were seen in higher numbers when compared with control explants. The activity of the brush border enzyme sucrase-isomaltase was reduced in the PHA E4 incubated explants but did not change in the PHA L4 incubated explants. This investigation shows that explants of pig jejunal mucosa in organ culture are suitable for study of the pathological effects of lectins on the small intestinal mucosa. This method may also be used in elucidating the mechanisms by which damage to mucosal structure occurs.

The ubiquitous presence of lectins – for example, seeds of legumes like pulses and beans – in plants eaten by man and his farm animals have been described in recent surveys. The biological activity of lectins can be abolished by proper heat treatment, although heat processing is not always very effective. Since most lectins are resistant to proteolytic breakdown in the gastrointestinal tract, they can reach the small intestine in a biological active form and bind to the glycoconjugates of the mucus and glycoalkx. Pusztai et al suggest that many of these lectins may account for the morphological and functional changes in the epithelium of the small intestine leading to reduced efficiency of nutrient utilisation. Despite several studies, however, the pathogenesis of lectin induced changes in the intestinal tract has not fully been explained. In vitro studies show promising results in investigation of the effects of lectins on the small intestinal mucosa without any interference from the luminal contents and the intestinal micro flora. Moreover, the experimental conditions of the mucosal explants can be controlled more precisely than those in vivo and therefore may allow investigators to single out the direct effect of any given damaging agent on the mucosa. Another advantage of in vitro studies is the fact that only small amounts of deleterious material are needed. We chose explants of pig jejunal mucosa in organ culture, and therefore closely related to the situation in vivo, to study the interaction between *Phaseolus vulgaris* (PHA) isolectins E4 (4 erythroagglutinating subunits) and L4 (4 leuco-agglutinating subunits) and the jejunal mucosa in vitro. We investigated the binding of PHA isolectins E4 and L4 to the explants and the subsequent lectin induced effects on morphology and brush border enzyme activity.

**Methods**

Approval for the experiments was given by the ethical and experimental committee of TNO-CIVO, Zeist, The Netherlands.

Four 7 week old, male SPF pigs (CD1, Leilystad, The Netherlands) were premedicated with Stresnil and atropine and anaesthetised with fluanthone and nitrous oxide in oxygen. The abdomen was opened and four jejunal segments 20 cm in length were taken 3 m proximal to the ileocaecal ligament. Explants were prepared according to Danielsen et al. The segments were cut open longitudinally at the mesenterial attachment and rinsed in Trowell's T8 medium at 21°C. With a scalpel, the mucosa was separated from the submucosa and small explants (9 mm²) were cut from the mucosa. The explants were cultured according to the method described by Browning and Trier. Briefly, the culture conditions are as follows. Explants were placed on triangular cut stainless steel grids, the villus side facing upwards (six explants per grid), in sterile plastic organ culture dishes (Falcon 3037). To the central well of these dishes 0.9 ml of Trowell's T8 medium was added, until a thin layer was drawn over the villous surface of the explants by capillary action. In the isolec tin experiments, 50 μg of the PHA E4 and PHA L4 isolectins respectively (E-Y Laboratories Inc, San Mateo, CA, USA) were added per ml of Trowell's T8 medium. The outer ring of the dish contained 3 ml of bidistilled water. The dishes were placed in an airtight container, gassed with a mixture of 95% O₂ and CO₂ for 15 minutes, and placed in an incubator (37°C). Explants cultured for five hours in Trowell's T8 medium and explants cultured for five hours in Trowell's T8 medium in the presence of 50 μg per ml of PHA E4 and L4 respectively, were processed for histomorphometric, electron microscopic, and biochemical examinations.

**Tissue Processing for Morphometric Analysis**

For morphometric analysis one explant from each dish was fixed in 0·1 mmol/l phosphate...
buffered 4% formalin pH 7-3, dehydrated, and embedded in paraffin. Serial sections (5 μm) were cut and stained with haematoxylin and eosin (HE). Using these sections, 10 well orientated villi and crypts were measured by means of a TEA Image Manager system (DIFA, Breda, The Netherlands). The villus height was represented by the distance from the crypt opening to the tip of the villus, and the crypt depth from the base of the crypt to the level of the crypt opening. The villus:crypt ratio was calculated to indicate the extent of morphological abnormality. The same sections were used to measure the enterocyte height from at least 25 epithelial cells with basally centrally situated nuclei distributed randomly over the villi.Crypt enterocytes as well as enterocytes in the extrusion zone were excluded from these measurements.

TISSUE PROCESSING FOR SCANNING ELECTRON MICROSCOPY
One explant per dish was fixed in 2-5% glutaraldehyde in 0-1 mmol/l cacodylate buffer (pH 7-35; 440 mOsm) for 24 hours at 4°C, and then rinsed in 0-1 mmol/l cacodylate buffer (pH 7-35). The fixed explants subsequently were treated according to the OTOTO method (alternating osmium tetroxide and thio-carbohydrazide immersion),34 dehydrated through graded concentrations of ethanol, transferred to amylacetate, critical point dried with CO₂, and mounted, villus side upwards, on aluminium stubs with silver paint. Specimens were examined using a Cambridge Camscan scanning electron microscope at 10 kV.

TISSUE PROCESSING FOR TRANSMISSION ELECTRON MICROSCOPY
One explant per dish was fixed in 0-1 mmol/l cacodylate buffered 2-5% glutaraldehyde (pH 7-35, 440 mOsm) for 24 hours at 4°C, and subsequently rinsed in 0-1 mmol/l cacodylate buffer (pH 7-35). After postfixation with 1% OsO₄ in 0-1 mmol/l cacodylate buffer (pH 7-35) for 16 hours at 4°C, the explants were dehydrated in graded water-aceton mixtures and embedded in an EponAraldite mixture. Semi-thin sections were stained with toluidine blue. Ultra-thin sections were stained with uranyl magnesium acetate and lead citrate and examined with a Philips EM410LS electron microscope at 60 kV.

TISSUE PROCESSING FOR IMMUNOELECTRON MICROSCOPY
One explant per dish was fixed in 2% paraformaldehyde and 0-5% glutaraldehyde in 0-1 mmol/l cacodylate buffer (pH 7-35) for seven days at 4°C and subsequently embedded in Lowicryl (polymerised at 20°C with 360 nm wavelength UV light). Ultra-thin sections were cut with a glass knife on a Reichert Ultracut E ultramicrotome and collected on uncoated 150 mesh copper grids. The primary antibodies used in this study were rabbit anti-PHA E4 and rabbit anti-PHA L4 (E-Y Laboratories Inc, San Mateo, CA, USA). Section labelling was performed by floating grids, sections down, on droplets placed on a strip of parafilm. All incubations were carried out at room temperature. The immunolabelling procedure is outlined as follows. Sections were incubated and pretreated on 20 μl droplets of 1% bovine serum albumine (BSA) (Organon, Oss, The Netherlands) in 0-1 mmol/l phosphate buffer saline, pH 7-4 (1% BSA/PBS) for 15 minutes. Excess buffer was removed with pieces of filtering paper before the grids were incubated on 20 μl droplets consisting of antibody diluted in 1% BSA/PBS for 30 minutes. The final dilution of both primary antibodies was 1:160 with 1% BSA/PBS. After incubation the grids were washed three times for five minutes by floating, section down, on fresh 20 μl droplets of 1% BSA/PBS. Excess buffer was removed before incubation for 30 minutes with colloidal gold labelled secondary antibody. In this study goat anti-rabbit conjugated to 15 nm colloidal gold (EM grade) (Janssen Pharmaceutica, Berse, Belgium) was used in a final dilution of 1:20. Before counterstaining with uranyl magnesium acetate and lead citrate, the grids were washed three times for five minutes with droplets of bidistilled water. Grids were air dried and examined with a Philips EM410LS electron microscope at 60 kV. Control sections were treated as above, omitting the primary antibody incubation step.35

SUCRASE-ISOMALTASE ENZYME ACTIVITY ASSAY
Three explants from each dish were frozen (liquid nitrogen) and stored at -70°C before analysis. The explants were pooled (three explants for each time point or PHA concentration) in 2 ml of 2 mmol/l TRIS-50 mmol/l mannitol buffer (pH 7-1). From here all procedures were performed at 4°C, unless mentioned otherwise.

The explants were sonified twice for 10 seconds in 1 ml of bidistilled water. After sonification an aliquot of the suspension was used for protein determination.36 Sucrase-isomaltase (EC 3.2.1.48) activity was determined according to Messer and Dahlqvist,37 using sucrose (Serva, Heidelberg, Germany) as substrate. The specific enzyme activity was calculated as U/g of protein and expressed as the mean relative specific enzyme activity (SEM).

STATISTICAL ANALYSIS
The statistical significance of differences between means was analysed using the paired Student’s t test. In the Tables the values are expressed as percentages of the corresponding control values.

Results
MORPHOLOGIC CHARACTERISTICS OF PIG JEJUNAL EXPLANTS
The histology of the control explants and PHA E4 cultured explants are shown in Figure 1A and B respectively. The results of the morphometric analysis of pig jejunal explants from PHA E4 and PHA L4 incubated and control explants are summarised in Table I. The morphology of the
control explants, cultured for five hours in Trowell’s T8 medium looked normal by light microscopy. In explants cultured in PHA isolectins E4 and L4 there was a considerable reduction in the villus:crypt ratio. (Incubation of the explants with the isolectins resulted in reductions of 30-1% for PHA E4 and 36-7% for PHA L4.) As shown in Table I, the PHA induced reduction in the villus:crypt ratio of the explants was due to a change in the villus length only, whereas the PHA L4 induced decrease was accomplished by both a decrease of the villus length and an increase of the crypt depth. After incubation of the explants for five hours with PHA isolectin E4, the mean enterocyte height decreased by 18.4%, whereas incubation with PHA isolectin L4 resulted in a decrease of 10.6%. Each of these reductions were statistically significant (p<0.01).

### ULTRASTRUCTURAL CHANGES IN THE EXPLANTS AFTER INCUBATION WITH PHA ISOLECTINS

The scanning electron microscopic studies showed that culturing explants for five hours in Trowell’s T8 medium did not influence the morphology of villi and microvilli (Figs 2A and B). However, as shown in Figure 3 A to D, exposure of the explants to 50 μg/ml of PHA E4 and PHA L4 respectively, resulted in changes in the morphology of the villi and microvilli. In the PHA E4 cultured explants a pock pitted villi surface and fusion of microvilli were seen (Figs 3A and B). The villi of the PHA L4 cultured explants were covered with dots of material (Figs 3C and D), which on cross section seemed to be derived from the microvilli.

The transmission electron microscopic study of control explants cultured in Trowell’s T8 medium, showed microvilli that were even in

![Figure 1: Effects of Phaseolus vulgaris isolectin E4 (PHA E4) on pig mucosal explants in organ culture (A). Control explant cultured for five hours in Trowell’s T8 medium (B). Explant cultured in Trowell’s T8 supplemented with 50 μg of PHA E4/ml. The depth of the crypts remains unchanged.](image)

![Figure 2: Scanning electron micrograph of a pig mucosal explant cultured in Trowell’s T8 medium (A). The microvilli are regularly shaped; some mucus is expelled on the surface (B).](image)

**TABLE I** Effects of Phaseolus vulgaris (PHA) isolectins on the length of villi, the depth of crypts, the villus:crypt ratio, and the height of enterocytes in pig jejunal explants (Values mean (SEM))

<table>
<thead>
<tr>
<th>Explants</th>
<th>Villus</th>
<th>Crypt</th>
<th>Villus:Crypt ratio</th>
<th>No</th>
<th>Enterocytes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>100 (0.2-9)</td>
<td>100 (0.2-5)</td>
<td>100 (0.9-9)</td>
<td>70</td>
<td>100 (0.1-5)</td>
<td>278</td>
</tr>
<tr>
<td>PHA E4</td>
<td>74 (1.7)*</td>
<td>114 (5.4)</td>
<td>63 (2.5)*</td>
<td>30</td>
<td>89 (1.4)*</td>
<td>115</td>
</tr>
<tr>
<td>PHA L4</td>
<td>74 (1.7)*</td>
<td>114 (5.4)</td>
<td>63 (2.5)*</td>
<td>30</td>
<td>89 (1.4)*</td>
<td>115</td>
</tr>
</tbody>
</table>

*p<0.01 compared with control.
**Pathological effects of Phaseolus vulgaris isolectins on pig jejunal mucosa in organ culture**

villi and over the tied off microvillus vesicles. Staining was more evident in the top third of the villi but gold particles were also detected over the microvilli, along the length of the villi, and over the mucous contents of goblet cells, but not in the crypts. No gold particles could be detected in control explants.

**EFFECT OF PHA ISOLECTINS ON THE BRUSH BORDER ENZYME ACTIVITY**
The effect of PHA E4 and L4 on the specific activity of the brush border enzyme sucrase-isomaltase in pig jejunal explants is shown in Table II.

Incubation of the explants in PHA E4 50 μg/ml resulted in a significant decrease of the relative specific activity compared with control explants cultured in Trowell’s T8 medium (Table II). PHA L4 at a concentration of 50 μg/ml did not influence the relative specific activity of sucrase-isomaltase.

**Discussion**
The objective of this study was to investigate whether organ culture from pig small intestinal mucosal explants represents a suitable model for studying the harmful effects of PHA isolectins E4 and L4 on enterocyte morphology and function. If so, this model may be used to examine the mechanisms of damage to small intestinal mucosal structure.

When small intestinal mucosal explants are cultured for up to five hours in Trowell’s T8 medium, only minor changes occur in the histological as well as ultrastructural architecture (Figs 1A, 2, 4A, and 5A). In addition, other investigators have reported, based on light and electron microscopic criteria, that the mucosal morphology of pig small intestinal explants is well preserved during organ culture. The minor changes in the mucosal morphology that take place during culture are presumably caused by the in vitro environment. Exposure of the pig jejunal explants to PHA E4 and PHA L4 for five hours provokes numerous morphological changes (Table I, Figs 1B, 3B and C, and 5B). These changes include a significant reduction in mean enterocyte height, a decrease in the villus:crypt ratio because of the reduced villus length and increased crypt depth in PHA L4 treated explants, and the presence of shortened and irregularly positioned microvilli with vast numbers of tied off microvillus vesicles. Furthermore, in the PHA E4 treated explants the specific activity of the brush border membrane enzyme sucrase-isomaltase was decreased significantly (Table II). No significant difference could be detected in the PHA L4 treated explants.

By electron microscope immunocytochemistry, immunogold localisation of PHA E4 and PHA L4 was shown over the glycocalyx of the intestinal microvilli, the microvillus vesicles in the top one third area of the villi (Fig 6), and somewhat less over the microvilli of enterocytes in the lower parts of the villi. No differences were observed between the distribution of the isolectins on the cells, which is in agreement with

**Figure 3: Scanning electron micrograph of mucosal explants after exposure to 50 μg of Phaseolus vulgaris (PHA) E4/ml (A, B) or 50 μg of PHA L4/ml (C, D) of Trowell’s T8 medium (A).**

Note the pock-pitted surface (B). Note the irregularity of the microvilli (C). Note the dots on the surface of the microvilli, which on cross section seem to derive from the microvilli (D).

length and regularly positioned (Figs 4A and 5A). The core of the microvilli, consisting of bundled actin filaments, was clearly seen to protrude into the cytoplasm. No vesicles seemed to be tied off from the microvillar tops. However, explants cultured in PHA E4 (Figs 4B and 5B) and PHA L4 (Fig 4C) clearly showed shortening of the microvilli of enterocytes on the upper third of the villi. They were irregularly positioned and fusion of microvilli frequently occurred. In addition, vast numbers of microvillus vesicles had been tied off and areas could be observed where a complete loss of microvilli had taken place. The described changes, however, seem less abundant in the PHA L4 treated explants compared with the PHA E4 treated ones.

**BINDING OF PHA E4 AND L4 TO PIG JEJUNAL EXPLANTS**

As seen by immunoelectron microscopy PHA E4 (Fig 6A) and PHA L4 (Fig 6B) bind to the microvilli of the cultured explants. No differences could be detected with regard to the distribution of gold particles. The gold particles were located almost exclusively over the micro-
the findings of King et al²⁴ in rats fed raw kidney bean protein.

Extensive changes in small intestinal enterocytes, decrease in villus length, irregularity of microvilli, clusters of vesicles associated with the brush border, and complete loss of microvilli are described in rats after intraluminal administration of concanavalin A or wheat germ agglutinin.¹¹ When rats (conventional as well as germ free) were fed purified PHA and adherence of PHA to the brush border membrane was obvious, none of these morphological changes of the small intestinal mucosa could be observed.⁴ However, rats fed a diet containing crude PHA beans display an extensive disruption and abnormal structure of microvilli in the small intestine.⁴⁻²⁵

Our findings clearly show that exposure of pig mucosal explants to PHA E4 or L4 for five

<table>
<thead>
<tr>
<th>Table II</th>
<th>Effect of Phaseolus vulgaris (PHA) isolectins on the specific activity of the brush border enzyme sucrase-isomaltase in pig jejunal explants (values, mean (SEM))</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Explant</strong></td>
<td><strong>Activity of sucrase-isomaltase (U/g protein)</strong></td>
</tr>
<tr>
<td>Control</td>
<td>100.0 (5.2)</td>
</tr>
<tr>
<td>PHA E4</td>
<td>82.9 (11.4)*</td>
</tr>
<tr>
<td>Control</td>
<td>100.0 (2.2)</td>
</tr>
<tr>
<td>PHA L4</td>
<td>98.7 (9.9)</td>
</tr>
</tbody>
</table>

* p<0.01 compared with controls.
hours results in binding of the isolecitins to the intestinal epithelium, brush border membrane changes, and diminished villus length; and in case of PHA E4 incubated explants, a decreased activity of sucrase-isomaltase.

Several mechanisms such as the globular (G):filamentous (F)-actin ratio, the cell actin content, the cytoskeletal protein turnover or the pool size of actin binding proteins, or both, might be involved in the regulation of the intestinal brush border villus length. In differentiated human colon carcinoma Caco-2 cells exposed to soybean agglutinin we have shown that the shortening of the microvillus length is accomplished only by a rapid shift in the G:F-actin ratio. After binding of the PHA isolecitins to the brush border membrane of the enterocytes, this mechanism (changes in the actin cytoskeleton of the cell) probably plays a role in the pathogenesis of microvillus abnormalities.

After exposure of rat intestinal epithelium to intraluminal dietary lectins clusters of vesicles associated with the brush border can be found. Also, in organ culture, blebbing and vesiculation of microvilli leading to increased numbers of vesicles nearby the brush borders of PHA isolecitins incubated explants were observed (Fig 5). The presence of these vesicles might reflect an increased turnover of microvillus membrane.

The loss of brush border membranes interferes at once with the activity of brush border membrane associated enzymes. Our results show a correlation between the amount of microvillus vesicles that have been tied off and the decrease in activity of sucrase-isomaltase (Table II). The similarity between the lectin induced lesions in vivo and in vitro responses of pig mucosal explants clearly shows that organ culture of pig mucosal explants represents a suitable...
model for further in vitro studies on the pathogenesis of lectin induced lesions in the pig small intestine.


Pathological effects of Phaseolus vulgaris isolectins on pig jejunal mucosa in organ culture.
M J Kik, J F Koninkx, A van den Muysenberg and F Hendriksen

Gut 1991 32: 886-892
doi: 10.1136/gut.32.8.886

Updated information and services can be found at:
http://gut.bmj.com/content/32/8/886

These include:
Email alerting service
Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

Notes

To request permissions go to:
http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/