Increased intestinal permeability in rats with graft versus host disease

W A Koltun, M M Bloomer, P Colony, G L Kauffman

Abstract

Background/Aims—The study of graft versus host disease of the intestine has significant clinical relevance and may also be a model for other immune mediated intestinal diseases. There presently is no simple non-invasive test that can be used to evaluate graft versus host disease induced intestinal injury in humans or animal models. This study tested the hypothesis that graft versus host disease leads to an increase in host bowel permeability as assessed by the relative urinary excretion of orally administered lactulose and rhamnose.

Methods—The urinary excretion ratio of orally administered lactulose and rhamnose was determined daily for two weeks in (Lewis x Brown-Norway) F1 rats with graft versus host disease caused by either the transplantation of parental (Lewis) small bowel or the intraperitoneal injection of parental (Lewis) splenic lymphocytes.

Results—Significant twofold to fourfold increases in the lactulose to rhamnose ratio were seen in both small bowel transplant and splenic lymphocyte transfer animals suffering from graft versus host disease during the second postoperative week. This effect occurred sooner in small bowel transplant than in splenic lymphocyte transfer animals (postoperative day 7 versus 11, respectively). The signs of graft versus host disease, including splenomegaly and altered intestinal mucosal architecture, as well as the increased lactulose to rhamnose ratio were significantly attenuated in small bowel transplant animals treated with cyclosporine A (10 mg/kg/day).

Conclusions—Graft versus host disease is associated with an increase in the lactulose to rhamnose clearance ratio reflecting an increase in host bowel permeability. This increase, along with the signs of systemic graft versus host disease, can be significantly ameliorated by cyclosporine A. The lactulose to rhamnose clearance ratio is a non-invasive technique that can be used to assess the intestinal effects of graft versus host disease and the associated increase in intestinal permeability.

Keywords: graft versus host disease, intestinal permeability, small bowel transplantation, cyclosporine A.
the possibility of rejection and the native, non-operated intestine was evaluated, so minimizing the effects of surgical injury to the intestine. To characterise and confirm the presence and progression of GVHD in these models, food consumption, animal and splenic weights, and intestinal histology at the time of death was performed.

Methods

Animal models and groups

Male Lewis (Lew) and Lewis x Brown Norway (LBN) F1 virus antibody free rats (250–300 g) were obtained from a commercial breeder (Harlan Sprague Dawley, Indianapolis, IN) and handled in accordance with the guidelines of the American Association for Accreditation of Laboratory Animal Care. Animals were quarantined upon arrival from the supplier and monitored for one week for illness prior to use. Animals were housed in metabolic cages (0650-0100, Nalgene Co, Rochester, NY) during experimental protocol to permit easy collection of urine for bowel permeability assessment. Animals had free access to water and Purina Standard Rodent Chow 5001, although food was removed from cages during the hours of permeability measurements to avoid contamination of urine collections with food debris.

GVHD was produced by either the auxiliary transplantation of small bowel (Fig 1) or intraperitoneal injection of splenic lymphocytes from Lew (RT-1<sup>1,11</sup>) donors into LBN F1 (RT-1<sup>1,10</sup>) hosts. Such a cross from inbred parent to F1 progeny produces GVHD without rejection because the major histocompatibility (MHC) antigens present in the homozygous grafted tissue are also present in the F1 recipient. The heterozygous F1 host however, expresses MHC antigens that are recognised as foreign by the grafted parental lymphocytes, resulting in GVHD. All animals, whether receiving small bowel grafts or intraperitoneal splenic lymphocytes, had their native intestines left intact for subsequent permeability measurements. Five groups of animals were studied:

1. SBTx-GVHD (n=14): LBNF1 animals with auxiliary Lew small bowel transplants (SBTx) resulting in GVHD;
2. Sham-op controls (n=13): LBNF1 animals with the same operative dissection as SBTx-GVHD animals but without small bowel grafts;
3. Spl-GVHD (n=9): LBNF1 animals with an average of 3×10<sup>6</sup> range 3×10<sup>3</sup>–6×10<sup>6</sup> Lew splenocytes injected intraperitoneally, resulting in GVHD;
4. Un-op controls (n=8): LBNF1 animals without operative manipulation;
5. SBTx-CsA (n=8): LBNF1 animals with Lew small bowel transplants (SBTx) as in group 1 above who also received 10 mg/kg/day cyclosporine A (CsA) subcutaneously.

All animals had the permeability of their native (in situ) bowel assessed daily for two weeks after transplantation using the lactulose/rhamnose (L/R) clearance technique (see below). Food cups were weighed daily to assess food consumption. Animals were weighed and inspected daily for signs of GVHD (cutaneous erythema, alopecia, weight loss, and dermatitis). Fourteen days after grafting or sham operation, animals were killed by exsanguination under ether anaesthesia. Spleens were weighed and segments of bowel harvested for histology to confirm the presence of intestinal GVHD.

Technique of small bowel transplantation

The technique of small bowel transplantation has been previously described. Briefly, small bowel from proximal jejunum to terminal ileum was harvested from the donor and subsequently anastomosed in an auxiliary fashion to the recipient’s aorta and vena cava. The proximal and distal ends of the graft were brought out as stomas, creating an auxiliary Thiry-Villa loop. The animal’s native bowel was left intact for subsequent bowel permeability assessment (Fig 1).

Splenic lymphocyte transplantation

Spleens were harvested from Lew rats using aseptic technique and placed into sterile Dulbecco’s phosphate buffered saline (D-PBS, GIBCO BRL, Life Technologies, Grand Island, NY). Tissue was gently minced using forceps and the resultant suspension passed through a 70 μm nylon mesh filter (Falcon, Becton Dickinson, Franklin Lakes, NJ). Red cells were lysed by the addition of 15 ml of red cell lysing buffer (Sigma, St Louis, MO). Spleen cells were suspended with D-PBS in a 50 cc conical tube and centrifuged for eight minutes at 200 g in a refrigerated centrifuge. The supernatant was discarded, the pellet washed two more times, and finally brought up to 10 cc with D-PBS. A cell count was done using a haemocytometer and trypsin blue exclusion to determine viability. An average of 3×10<sup>6</sup>–6×10<sup>6</sup> viable cells (range 3×10<sup>3</sup>–6×10<sup>6</sup>) cells were injected intraperitoneally into recipient LBN F1 animals using an 18 gauge needle. It has been shown that a minimum of 1×10<sup>8</sup> splenocytes are necessary to cause GVHD in this rat model.
Lactulose/rhamnose clearance

Animals were gavaged daily each evening with a solution containing 400 mg lactulose and 100 mg rhamnose (Pfanstiehl Laboratories, Waukegan, IL) in 5 cc of distilled water. The urinary clearance of orally administered lactulose and rhamnose has been shown to be largely complete in rats by eight hours.15 Urine was collected for at least eight hours overnight in receptacles containing 200 μl of chlorhexidine (Sigma, St Louis, MO) as a preservative. Urine samples were centrifuged at 200 g for 10 minutes and 1-0 ml aliquots of the supernatant were frozen at −70°C for later analysis by high pressure liquid chromatography (HPLC). Urine samples before gastric gavage were confirmed as containing unmeasurable amounts of both the test sugars and internal standard sugars (see below).

Stored urine specimens were defrosted on ice and centrifuged at 200 g for five minutes. The supernatant was diluted 10-fold to 200-fold with deionised water to yield final test sugar concentrations that fell within the linear range of standard curves for each sugar. Internal standards, arabinose, and cellobiose (Pfanstiehl Labs, Waukegan, IL) were added to obtain a final concentration of 20 mg/l. Samples were filtered through 0.2 μm pore size syringe tip filters (Supelco, Bellefonte, PA) and 20 μl aliquots analysed by HPLC using an anion exchange column (Carbopac PA1, Dionex, Marlton, NJ) and pulsed amperometric electrochemical detection (Coulochem II, ESA, Nieuw Vennep, Bedford, MA).17 Standard curves were run for test and internal standard sugars and confirmed to be linear by least squares analysis (r²>0.95). Peak height analysis of the chromatograms (Spectra Physics SP 4270) was used to determine concentrations of the sugars. Test sugar results were expressed as the ratio: L/R=percent urinary recovery of orally administered lactulose/percent urinary recovery of orally administered rhamnose.

Absolute and relative spleen weights

Both absolute and relative splenomegaly have been used as quantifiable indicators of GVHD. At the time of death (14 days postoperation) the absolute wet weight of each animal’s spleen was determined to be the nearest 0.1 mg. One animal in the Spl-GVHD group died on day 13 so only eight animals in this group had spleen weights determined. Relative spleen weight was expressed as absolute spleen weight (g)/animal body weight at the time of death (kg).

Intestinal histology

To confirm the effects of GVHD on the host intestine, samples of small bowel were harvested at the time of death from eight Sham-op, eight SBTx-CsA animals, six SBTx-GVHD animals (and three additional animals done subsequently for a total of nine SBTx-GVHD animals), and six Spl-GVHD animals and evaluated histologically for crypt hyperplasia and villus atrophy. These two features have been shown to be indicative of intestinal GVHD.18-20 Full thickness segments of proximal jejunum, mid (50 cm) and distal (90-100 cm) small bowel were harvested from each animal and stained in standard fashion with haematoxylin and eosin. An average villus and crypt length per segment of bowel studied was generated by a blinded histologist by measuring 20 well oriented villi and crypts with an eyepiece micrometer.
implemented in SAS PROC GLM (SAS Institute, Cary, NC) and p values reported.

Results

Clinical appearance
Both the SBTx-GVHD and Spl-GVHD animals recovered and appeared normal until the seventh or eighth postoperative day when the first signs of GVHD disease became manifest as mild cutaneous erythema of the paws, ears and snout, with some hair loss. These features gradually progressed to dramatic cutaneous hyperemia and dermatitis, alopecia, nasal and ocular discharge, and phimotic appearing genitalia. There was no diarrhea. Stools became viscid, but were always in pellet form, a fortuitous situation as urine collections were not contaminated by liquid faecal material. Animals became less active and assumed a hunched posture with apparent difficulty in fully extending their limbs by the 13th to 14th day. Sham operated animals recuperated and returned to normal activity and appearance promptly without clinical deterioration during the experimental period. Unoperated animals exhibited normal activity and behaviour. SBTx-CsA animals appeared generally healthy even during the second postoperative week. Mild cutaneous erythema was observed in some,
however, there were no hair or mucous membrane changes.

**Food consumption/weight changes**

Figure 2 shows the mean weight change of the animal groups over the course of the experiment. The Un-op group gained an insignificant amount of weight during the two week period, while the Sham-op group had a temporary period of postoperative weight loss before returning to a weight comparable to the preoperative level. Animals with either form of GVHD, as well as the CsA treated animals all had a similar pattern of weight loss over the experimental period that resulted in these three groups having a final weight at death significantly less than Sham-op or Un-op controls.

Figure 3 shows the mean daily food consumption for each of the groups during the first and second 7 day postoperative periods of the experiment. Food consumption during the first week was very similar, but significantly depressed in those animal groups that underwent surgery compared with those animal groups that did not have surgery. Food consumption during the second week returned to control levels in all animal groups, except the Spl-GVHD group in which there was a significant decline. This decrease in food intake may have accounted, in part, for the concurrent rapid weight loss seen in this animal group. The weight loss seen in the SBTx-GVHD and SBTx-CsA groups during the second week, however, could not be attributed to decreased dietary intake.

**Intestinal histology**

There were significant reductions in villus height and increases in crypt depth in SBTx-GVHD animals compared with Sham-op controls in the proximal, mid, and distal regions of the small bowel confirming the effects of GVHD on the host intestine (Fig 5). In SBTx-GVHD animals, villus blunting appeared to be most severe in the distal bowel where mean villus height was 38% less than Sham-op controls whereas it was only 12% and 20% less in the proximal and mid small bowel areas, respectively. Though there was also significant villus blunting diffusely in Spl-GVHD animals, crypt elongation was less prominent, suggesting less severe intestinal injury compared with SBTx-GVHD animals. It is noteworthy that epithelial integrity was maintained in all segments of bowel studied in both GVHD animal groups with no loss of surface epithelium or ulcerations noted (Fig 6). When SBTx animals were treated with CyA (SBTx-CyA group), there were significant improvements in villus and crypt lengths compared with SBTx-GVHD animals, approaching though not completely returning to Sham-op control values in all segments of intestine studied.

**Lactulose/rhamnose bowel permeability**

Unoperated control animals (Un-op group) showed very consistent mean L/R ratios from day to day during the two week experimental periods (Fig 7). Sham-op L/R ratios were
transplant models and passive immunisation using anti-
Escherichia coli antiserum has been shown to improve animal survival.24-27 Similarly, human studies in bone marrow transplant recipients have suggested that gut decontamination with oral antibiotics may decrease the clinical signs and infectious complications of GVHD.28 A two stage model of intestinal injury in GVHD includes an immunological phase, in which grafted lymphocytes recognise alloantigen on the host tissues, and an effector or tissue damage phase, where the release of cytokines or other inflammatory mediators incur or aggravate injury to the bowel.25 The resulting increased permeability of the injured intestinal epithelium may represent a third phase, which might augment local injury or promote systemic sepsis by allowing the entry of noxious luminal microflora or toxins. The increase in the L/R ratio found in this study supports this concept of a compromised intestinal barrier in GVHD injury of the intestine. It is important to realise, however, that this increase in bowel permeability as identified by the raised L/R ratio need not necessarily correlate with increased bowel permeability to other compounds, including enteric organisms. Different mechanisms for uptake of various materials exist in the intestine. Translocation of bacteria and fungi has been shown to occur transcellularly, for example.29 The presently identified increase in L/R therefore, cannot be freely extrapolated to other materials, including much larger substances such as endotoxin or bacteria but does reinforce the notion that the intestinal defensive barrier is compromised in GVHD.

This study suggests increases in L/R are a consequence of immune mediated inflammation directed towards the intestine during GVHD. The exact mechanism is unclear but may entail the local release of potent cytokines.30 Two cytokines that have been implicated in GVHD injury include interferon γ (IFN γ) and tumour necrosis factor (TNF). Increased serum concentrations of TNF in GVHD after SBTx have been described and antibodies to both IFN γ and TNF have been shown to ameliorate some of the histopathological changes of GVHD injury in mice.14 18 31 Of significant interest, IFN γ has also been shown to play a part in intestinal barrier function. Using chamber studies using cultured monolayers of T-84 human intestinal cells have shown a decrease in transepithelial resistance at the level of the intercellular tight junction after IFN γ administration.32 33 As the L/R technique of bowel permeability measurement is felt to assess paracellular integrity, the alteration defined in this study may represent the in vivo correlate of this in vitro increase in epithelial permeability that occurs secondary to the local release of this cytokine during GVHD intestinal injury.

Weight loss during GVHD has been described before and may result from a number of factors, including intestinal dysfunction, compromised ability to utilise the absorbed nutrients or a hypermetabolic state.34

Discussion

The concept of the gut as a reservoir of bacteria or toxins that aggravates local injury or leads to systemic illness when the protective enteric permeability barrier is compromised, is one that has been proposed in numerous illnesses, including GVHD.23-26 Sterilisation or selective decontamination of the gut has attenuated GVHD in murine bone marrow

Figure 8: Effect of cyclosporine A (CsA, 10 mg/kg/day) on L/R in rats with graft versus host disease after small bowel transplantation. The L/R curves of the SBTx-GVHD and sham-op animals are reproduced from Figure 7 for comparison with the CsA treated animals (SBTx-CsA). The L/R ratio of the SBTx-CsA group for the 7–13 day period was significantly less than that in the SBTx-GVHD group (p<0.025), but still increased relative to the sham-op control group (p<0.04) by repeated measures analysis of variance.

statistically no different than those of the Un-op group during days 7–13, although there appeared to be a transient increase in L/R at days 5 and 6 as previously mentioned.

Compared with the Sham-op group, the SBTx-GVHD group exhibited a statistically significant increase in L/R ratios for the 7–13 day period (p<0.001), when the clinical signs of GVHD were manifest. Similarly, there was a statistically significant increase in the L/R ratio in the GVHD-Spl animals versus Un-op animals during the same 7–13 day period (p<0.02). Qualitatively, the increase in the L/R ratio in the SBTx-GVHD group occurred earlier than that seen in the Spl-GVHD group (postoperative day 7 versus postoperative day 11, respectively). When these two groups were statistically compared, this difference was found to be significant (p=0.05, GVHD-SBTx v GVHD-Spl).

Figure 8 shows the L/R ratios of the SBTx-CsA group compared with the SBTx-GVHD and Sham-op groups. When animals receiving small bowel transplants were treated with CsA (SBTx-CsA group), L/R ratios for the 7–13 day period were significantly lower than that seen in the untreated SBTx-GVHD group (p<0.025). Cyclosporine A did not completely ameliorate the raised L/R ratios seen in the SBTx-GVHD group however, because ratios in the SBTx-CsA group were still significantly increased relative to the Sham-op group (p<0.04).
Increased intestinal permeability in rats with graft versus host disease

It is noteworthy that there was a similar pattern of weight loss in the SBTx-CsA animals who also had control levels of food intake but did not manifest overt signs of GVHD. This suggests that either CsA itself had cachexia inducing effects, or that CsA imperfectly treated the GVHD. The latter is probably the case as firstly, the L/R diuresis did not return completely to control values in the SBTx-CsA group and secondly, a subsequent group of six animals injected daily with CsA (10 mg/kg) had a mean weight after 14 days virtually identical to their mean starting weight (103 (3-5)%). The weight loss seen in the SBTx-CsA animals also disputes the possibility that the significant increases in the L/R ratios seen in the GVHD groups of animals were simply due to generalised nutritional deterioration causing intestinal atrophy and injury, because the SBTx-CsA group had a relatively low L/R ratio associated with significant weight loss.

The increased L/R ratio seen in this study was significant and presents the sugar clearance technique as a comparatively non-invasive method by which to follow GVHD injury in the controlled laboratory or clinical setting. Assessment of treatment directed at ameliorating intestinal GVHD injury could be accomplished, in part, by repetitive measurement of the L/R clearance ratio. Such a study in Crohn’s disease patients showed an increased lactulose/rhamnose ratio in those with active disease, which decreased when patients were treated and became clinically improved. It would be tempting to suggest that the lactulose/rhamnose measurement could provide a non-invasive method by which to diagnose intestinal GVHD in patients undergoing bone marrow or small bowel transplantation. Limitations on the applicability of the technique may exist, however, as many other factors have been associated with increased bowel permeability. These include surgery, chemotherapy, neoplasia, sepsis, and malnutrition, many of which could also be present in the patient with GVHD. Repetitive L/R measurements may none the less be clinically useful in the individual GVHD patient by providing an improving or deteriorating trend that could minimise the need for more morbid endoscopic biopsy procedures. In this regard, the L/R technique has significant advantages over other techniques of bowel permeability measurement, especially those using radioactive tracers, as it can be repeatedly applied in the clinical arena with minimal risk. In the controlled laboratory setting, the L/R ratio could similarly be effective. Therapeutic interventions aimed at decreasing intestinal GVHD injury, such as was done here with CsA, could be assessed in animals on a daily basis without the need for death.

The clinical manifestations of GVHD in the SBTx-GVHD and the Spl-GVHD groups were similar, but both the L/R and intestinal histology data suggest that a more severe intestinal injury occurred in the SBTx-GVHD animals. Crypt elongation occurred in proximal, mid, and distal small bowel in the SBTx-GVHD animals, but was present only in the proximal portions of the bowel in Spl-GVHD animals. The L/R increase appeared sooner in SBTx-GVHD animals compared with Spl-GVHD animals (day 7 v 11, respectively). The disparity in food consumption between the Spl-GVHD and SBTx-GVHD animals at the time of the measurement suggest that GVHD after SBTx may be different than that seen with splenocyte transfer. Such differences may be related to the finding that lymphocytes from the intestine tend to ‘home’ back to the gut. GVHD intestinal injury after SBTx may be greater than that caused by peripheral or splenic lymphocyte transfer because of this trafficking phenomenon, directing and preferentially localising the donor enteric lymphocytes and injury to the host bowel. This suggests that data regarding GVHD mediated intestinal injury obtained from clinical bone marrow transplantation patients may not be directly applicable to the increasing number of patients receiving small bowel transplantation. Further studies are warranted, evaluating this concept of lymphocyte homing and host intestinal injury after bowel transplantation.

In summary, using both splenocyte and small bowel transplant models of GVHD, increases in host bowel permeability, as measured by the lactulose/rhamnose urinary clearance technique were observed. This increase in the lactulose/rhamnose ratio was attenuated by treatment of small bowel transplant animals with cyclosporine A. These data suggest that the lactulose/rhamnose urinary clearance assay may provide a non-invasive method by which to assess intestinal injury and the effect of treatment in GVHD mediated injury of the bowel.

13 Kohlz WA, Diamantis T, Kirkman RL. Synergism between anti-interleukin-2 receptor monoclonal antibody
Increased intestinal permeability in rats with graft versus host disease.

W A Koltun, M M Bloomer, P Colony and G L Kauffman

Gut 1996 39: 291-298
doi: 10.1136/gut.39.2.291

Updated information and services can be found at:
http://gut.bmj.com/content/39/2/291

Email alerting service

These include:

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/