Establishment and evaluation of an experimental rat model for high-altitude intestinal barrier injury

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Abstract. In the present study an experimental high-altitude intestinal barrier injury rat model was established by simulating an acute hypoxia environment, to provide an experimental basis to assess the pathogenesis, prevention and treatment of altitude sickness. A total of 70 healthy male Sprague-Dawley rats were divided into two groups: Control group (group C) and a high-altitude hypoxia group (group H). Following 2 days adaptation, the rats in group H were exposed to a simulated 4,000-m, high-altitude hypoxia environment for 3 days to establish the experimental model. To evaluate the model, bacterial translocation, serum lipopolysaccharide level, pathomorphology, ultrastructure and protein expression in rats were assessed. The results indicate that, compared with group C, the rate of bacterial translocation and the apoptotic index of intestinal epithelial cells were significantly higher in group H (P<0.01). Using a light microscope it was determined that the intestinal mucosa was thinner in group H, there were fewer epithelial cells present and the morphology was irregular. Observations with an electron microscope indicated that the intestinal epithelial cells in group H were injured, the spaces among intestinal villi were wider, the tight junctions among cells were open and lanthanum nitrate granules (from the fixing solution) had diffused into the intestinal mesenchyme. The expression of the tight junction protein occludin was also decreased in group H. Therefore, the methods applied in the present study enabled the establishment of a stable, high-altitude intestinal barrier injury model in rats.

Introduction

High altitude is a low-pressure and hypoxic environment and it has been demonstrated that acute hypoxia can cause injury to a number of organs (1-3). The specialized blood supply and anatomical structure of the intestine increases its sensitivity to hypoxia, which makes the intestine vulnerable to the effects of hypoxic stress (4-6). It has been demonstrated that there are varying degrees of human gastrointestinal tract injury that occur in high-altitude hypoxic environments (7-9). As the largest bacteria and endotoxin library in the body, once the barrier function of the intestine is damaged, a large number of bacteria and endotoxins may translocate to other tissues or organs via the blood stream (10,11). This induces a cascade of inflammatory mediators, the onset of systemic inflammatory response syndrome and multiple organ dysfunction syndrome (MODS) (12,13) that may severely impact the health of individuals at high altitudes. The pathogenesis of acute hypoxia is complicated and remains unclear. There are vast plateau areas in China, where >60 million people live. When the Qinghai-Tibet railroad opened, large numbers of plainsmen began to participate in local economic construction. The safeguarding of people's safety and health is closely associated with national economic development. Research into the pathogenesis of high-altitude gastrointestinal disease may improve the quality of life of people living in the plateau areas and may provide guidance for such people with regard to prolonged, healthy living.

There are no similar conditions for assessment, and a lack of stable and repeatable experimental animal models of high-altitude intestinal injury restricts the understanding of this condition and the evaluation of potential therapeutic strategies. Thus, establishing an animal model is particularly important, as it is the basis of studying this injury. In the present study, a simulated high-altitude environment chamber combined with exercise on a treadmill was used to establish an animal model, which may provide an experimental basis for future studies.

Materials and methods

Animals and grouping. A total of 70 male Sprague-Dawley rats (7 weeks old) were purchased from the Experimental
Animal Center of West China Hospital (Sichuan, China). The rats weighed 250±15 g and were randomly divided into two groups: Control group (group C) and the high-altitude acute hypoxia group (group H). The experiment was approved by the Chengdu Military General Hospital Ethics Committee (Chengdu, China).

Reagents and equipment. MacConkey agar culture medium and KTA Tachypleus amebocyte lysate were purchased from Zhanjiang Bokang Marine Biological Co., Ltd. (Guangzhou, China), terminal deoxynucleotidyl transferase-mediated dUTP nick end labeling (TUNEL) kit was purchased from Roche Diagnostics (cat. no. 11684817; Basel, Switzerland) and occludin rabbit monoclonal antibody (cat. no. 71-1500) was purchased from Thermo Fisher Scientific, Inc. (Waltham, MA, USA). A FLYDWC50-1A simulated high-altitude low-pressure chamber was purchased from Guizhou Fenglei Air Ordnance Ltd. Co. (Guizhou, China). A Tecnai 10 transmission electron microscope (Koninklijke Philips N.V., Amsterdam, Netherlands), S-3400 N scanning electron microscope (Hitachi, Ltd., Toyo, Japan) and ZH-PT experimental animal treadmill (Huaiabei Zhenghua Biological Instrument Equipment Co. Ltd., Anhui, China) were also used in the present study.

Establishing the animal model. The animals were raised in the respiratory department of Chengdu Military General Hospital (Chengdu, China) at an altitude of ~500 m and a temperature of 24°C for 1 week to allow them to adapt to the environment. Food and water were provided ad libitum and a natural light cycle (12 h light/12 h dark) was adopted.

Adaptive training on the plain. The adaptive training began at 8:00 a.m. on day 8 and was conducted as follows: A total of 10 tracks of the ZH-PT experimental animal treadmill were set to a slope degree of 0° and a rotating speed of 8 m/min. Rats exercised for 30 min, followed by a 30 min rest. During the training, rats were free to drink and had access to food ad libitum. The training was repeated 5 times in the same day, with a total training time of 2.5 h.

Intervention training stage on the plain. The present model was established based on the high-altitude pulmonary edema (HAPE) animal model described by Bai et al (14), which did not aim to exhaust the animals. In order to induce fatigue stress, the treadmill settings were adjusted. At 8:00 a.m. on day 9, the animals were put on the treadmill, the slope of the conveyor belt was set to 0° and rotation speed was set to 8 m/min. The rats ran for 15 min to adapt to the treadmill. Following this, the slope was adjusted to 10° and the rotation speed was increased to 15 m/min. The rats exercised for 4 h, followed by a 30 min rest. Rats had ad libitum access food and water during the training. The training was repeated until 8:00 a.m. on day 10, after 24 h training. If the rats stopped, they were transported to the end of treadmill where there were some electrified pillars. Thus, appropriate electrical stimulation was applied to maintain the running state. The aim was to make the exercise as consistent as possible for all animals.

Stimulated hypoxia stage. In order to stimulate acute hypoxia in group H rats, at 8:00 a.m. on day 10, rats in groups C and H were moved into separate chambers of the FLYDWC50-1A simulated high-altitude low-pressure chamber. A compression resistant door separated the chambers, with an adapter chamber in the middle and a precise control system regulated the two separate environments. Rats in group C were kept at a temperature 23-25°C, humidity of 60%, wind speed of 0.3 m/sec and a 12-h light/dark cycle. A total of 25 g/day rat food was provided for each rat (12.5 g every 12 h) and the rats were free to get water. Hypobaric hypoxia was not applied to the group C rats. For group H, the access to food and water were same to group C, but the altitude was set to 4,000 m, the chamber pressure was 40.4 kPa, the air capacity was 40 m3/h, the temperature was 25°C, the humidity was 60%, the O2 content was 18.0%, the oxygen partial pressure was 7.8 kPa, the ascending rate was 3 m/sec and the light/dark cycle was 12-h.

Sample collection. Following 72 h in the separate environments, blood and abdominal visceral tissue samples from rats in the two groups were collected. Following the induction of anesthesia (10% ethyl carbamate, 1 ml/kg body weight by intraperitoneal injection; Chengdu Kelong Chemical Co., Ltd. (Chengdu, China), samples from group H were collected within 10 min by surgical resection at a simulated altitude of 4,000 m and all operations were strictly aseptic. From each group (n=35), 20 rats were selected for sacrifice by surgical resection as described above, and the blood, liver, spleen, mesenteric lymph nodes and jejunum were collected and preserved in liquid nitrogen (-195°C). The remaining 15 rats from each group were anesthetized (10% ethyl carbamate, 1 ml/kg body weight by intraperitoneal injection) and 2 cm jejunum was removed, 10 cm from the pylorus. The jejunum was sliced into 1-mm samples, which were immersed in fixing solution for pathological observation. For each group, 5 samples were immersed in 4% paraformaldehyde fixation fluid, 5 samples were immersed in 2.5% glutaraldehyde fixation fluid, and the rest were immersed in lanthanum aldehyde fixation solution (2% triformol, 2.5% glutaraldehyde, 2% lanthanum nitrate, 0.1 M sodium cacadylate trihydrate). All of the fixing solutions were at 4°C. Samples were fixed for 24 h.

Detection indices. To detect bacterial translocation, a homogenate was produced by adding 900 µl sterile saline solution to 100 mg samples of liver, spleen and mesenteric lymph node. A total of 0.3 ml homogenate was inoculated on MacConkey culture medium (17 g peptone, 5 g pig bile salts, 5 g NaCl, 17 g agar, 1,000 ml distilled water, 10 g lactose, 10 ml 0.01% crystal violet aqueous solution, 5 ml 0.5% neutral red aqueous solution). Following incubation at 37°C for 24 h, primary biochemical identification was applied. The bacterial translocation rate (%) = positive organ number in the same group/total number of cultured organs.

Dynamic turbidimetric analysis was used to detect the level of lipopolysaccharides (LPS) in the serum. A total of 3 ml serum was collected by apex puncture and centrifuged at 4°C at 960 x g for 15 min. From this, 100 µl serum was collected and 900 µl KTA Tachypleus amebocyte lysate was added. This was mixed evenly and kept at 65°C for 20 min. Following this, 200 µl mixture was added to the primary reagent of the enzyme reaction (from the kit) and LPS content was automatically detected following 60 min using a Bacterial
4% triformol at 4˚C, embedded and cut into 5-µm paraffin slices, totic epithelial cells. The samples were routinely fixed in 4% formaldehyde, 2.5% glutaraldehyde, 2% lanthanum nitrate, 0.1 M sodium cacodylate trihydrate, as lantham nitrate was used as the tracer. The samples were put into lanthanum aldehyde solution for fixation at 4˚C for 24 h, then, the samples were cut into 4-µm sections and stained with hematoxylin and eosin (H&E). The changes in the intestinal epithelium villi and mucous layer were observed using a CX23 optical microscope (Olympus Corporation, Tokyo, Japan); 10 high-power fields were selected in every section to detect the height and surface area of villi using the following formula: \[ S = \pi rh \], where \( r \) is the radius of villus and \( h \) is the height of villus. The mean value was noted.

Pathological observation by scanning electron microscopy. The perfusion and fixation solutions were changed to 2.5% glutaraldehyde solution in order to observe the samples using a scanning electron microscope, but other protocols were the same as previously stated for light microscopy. The samples were placed in 2.5% glutaraldehyde solution at 4˚C, and prepared as specimens according to the routine specimen preparation method (15), prior to observation under the scanning electron microscope.

Pathological observation by transmission electron microscopy. In order to observe samples using a transmission electron microscope, perfusion and the fixation solutions were changed to lanthanum aldehyde fixation solution (2% triformol, 2.5% glutaraldehyde, 2% lanthanum nitrate, 0.1 M sodium cacodylate trihydrate), as lanthanum nitrate was used as the tracer. The samples were put into lanthanum aldehyde solution for fixation at 4˚C for 24 h and prepared as specimens according to the routine specimen preparation (15), prior to observation under the transmission electron microscope.

Apoptosis of intestinal epithelial cells observed by histochemistry. A TUNEL assay was used to detect apoptotic epithelial cells. The samples were routinely fixed in 4% triformol at 4˚C, embedded and cut into 5-µm paraffin slices, according to the manufacturer’s protocol (Roche Diagnostics). A stained nucleus, in blue, was defined as positive for apoptosis. Every section was observed under a light microscope, at a magnification of x400, to count the cells (total cells of 4 fields/4), and the apoptotic index was calculated as follows: Apoptotic cell number/total cell number.

Western blot analysis to detect occludin expression. Total proteins from the jejunum tissue were extracted from a 50-mg sample and protein concentration was detected using a bicinchoninic acid (BCA) protein assay kit (Beyotime Institute of Biotechnology, Haimen, China). The protein was diluted to 5 µg/µl and 5X SDS loading buffer was added prior to heating to 100˚C for 3 min to degenarate the proteins. A total of 20 µg/µl protein solution was loaded in each well; this then underwent electrophoresis on a 5% SDS stacking gel and a 10% SDS separating gel at 90 V for 100 min. This was transferred to PVDF membrane by 100 V for 50 min, and put into 5% skimmed milk powder solution to block for 1 h at 25˚C. Then, occludin primary antibodies (71-1500; dilution 1:1,000) and β-actin primary antibodies (sc47778; dilution 1:3,000; Santa Cruz Biotechnology, Inc., Dallas, TX, USA) were added and the samples were incubated at 4˚C overnight. Following washing with Tris-buffered saline and Tween-20, the corresponding secondary antibodies (ZB-2301 and ZB-2305; Beijing Zhongshan Golden Bridge Biotechnology Co., Ltd., Beijing, China) for occludin and β-actin respectively, were added and the membrane was incubated at 25˚C for 1 h. The proteins were then developed using a BeyoEcl Plus kit (P0018; Beyotime Institute of Biotechnology Beijing, China). The relative grey value was used to quantify occludin expression, using ImageJ software, v2.1.4.7 (National Institutes of Health, Bethesda, MD, USA) according to the following formula: Relative grey value = grey value of occludin band/grey value of β-actin band.

Statistical analysis. SPSS, version 18.0 (SPSS, Inc., Chicago, IL, USA) was used for statistical analysis. Enumeration data were presented as percentages and the measurement data were presented as mean ± standard deviation. The comparison among groups was completed by independent-sample t-test and the comparison of ratio was completed by \( \chi^2 \) analysis. P<0.05 was considered to represent a statistically significant difference.

Results

Translocation of bacteria. There was no bacterial translocation detected in group C and the bacterial translocation was significantly higher in group H (P<0.01; Table I).

Table I. Change of bacterial translocation in the two groups.

| Group | Organ number (n) | Blood | Liver | Spleen | Lymph node | Translocation rate (%) |
|-------|-----------------|-------|-------|--------|------------|------------------------|
| C     | 80              | 0     | 0     | 0      | 0          | 0                      |
| H     | 80              | 3     | 6     | 5      | 7          | 26.25*                 |

\( ^* P<0.01 \) vs. group C. C, control group; H, high-altitude acute hypoxia group.

Endotoxin Dynamic Testing system (EDS-99; Tianjin Wireless Electronic Co., Ltd., Tianjin, China).

Pathological observation by light microscopy. Perfusion in situ combined with post-fixation was used to prepare the light microscope specimens. Animals were anesthetized (10% ethyl carbamate, 1 ml/kg body weight, by intraperitoneal injection), the right auricle was opened and the left ventricle was intubated. For perfusion, 200 ml normal saline at 4˚C and 100 ml 4% triformol were used. When there was clear effluent from the right auricle and the liver was hard, the perfusion was deemed successful. The 2 cm of jejunum was cut into 1-mm sections, following fixation using 4% triformol at 4˚C for 24 h, the specimens were dehydrated with 95% ethyl alcohol and embedded in paraffin. Then, the samples were cut into 4-µm sections and stained with hematoxylin and eosin (H&E). The pathological observation by scanning electron microscopy, perfusion and the fixation solutions were changed to 2.5% glutaraldehyde solution in order to observe the samples using a scanning electron microscope, but other protocols were the same as previously stated for light microscopy. The samples were placed in 2.5% glutaraldehyde solution at 4˚C, and prepared as specimens according to the routine specimen preparation method (15), prior to observation under the scanning electron microscope.

Pathological observation by transmission electron microscopy. In order to observe samples using a transmission electron microscope, perfusion and the fixation solutions were changed to lanthanum aldehyde fixation solution (2% triformol, 2.5% glutaraldehyde, 2% lanthanum nitrate, 0.1 M sodium cacodylate trihydrate), as lanthanum nitrate was used as the tracer. The samples were put into lanthanum aldehyde solution for fixation at 4˚C for 24 h and prepared as specimens according to the routine specimen preparation (15), prior to observation under the transmission electron microscope.
Table II. Change of height, surface area of villi and thickness of mucous layer in the two groups (mean ± standard deviation, n=5).

| Group | Height of villi (µm) | Surface area of villi (µm²) | Thickness of mucous layer (µm) |
|-------|---------------------|-----------------------------|-------------------------------|
| C     | 259±13              | 16,198±1085                 | 402±9                         |
| H     | 217±21              | 13,423±996                  | 336±7                         |

*P<0.05 vs. group C; C, control group; H, high-altitude acute hypoxia group.

Detection of serum endotoxin. The serum LPS concentration in group C was 0.051±0.011 EU/ml and the LPS concentration in group H was 0.803±0.115 EU/ml, a statistically significant difference (P<0.01; Fig. 1).

Pathological results. The results from light microscopy demonstrate that in group C, the epithelial mucous was smooth, the villi were arranged regularly and there was an even distribution of goblet cells among epithelial cells (Fig. 2A). In group H, the intestinal epithelial mucous was thinner, the villi reduced, the intestinal epithelium demonstrated that for group C, the villi were smooth, the epithelial mucous was smooth, the villi height, surface area and mucous thickness were significantly decreased in group H (P<0.05; Table II).

Scanning electron microscopy. Scanning electron microscopy demonstrated that for group C, the villi were smooth, arranged regularly and morphology was plump (Fig. 3A), however, in group H, the epithelial microvilli were shorter, areas had shrunk and broken. Furthermore, there was space widening (Fig. 3B).

Transmission electron microscopy. Transmission electron microscope results indicated that in group C, there was a high density of lanthanum granules attached to the capillary cavity and microvilli surfaces, the tight junction between epithelial cells was closed and there was no lanthanum granule deposition in the basement membrane or extracapillary mesenchyme (Fig. 4A). The cytoplasm of epithelial cells was complete and the morphology of endoplasmic reticulum and mitochondria was normal (Fig. 4B). In group H, the tight junction among epithelial cells was open and lanthanum granules permeated into the widened tight junction (Fig. 4C). There were lanthanum granules in the extracapillary space and basement membrane, indicating continuous linear distribution. Furthermore, parts of the organelles were swollen (Fig. 4D).

Apoptosis of epithelial cells. In group C, there were no clearly apoptotic cells in the intestinal epithelium (Fig. 5A), and the TUNEL assay indicated that the apoptotic rate was 3.4±1.1% (data not shown). In group H, there were a large number of apoptotic cells in the intestinal epithelium (Fig. 5B) and the apoptotic rate was 18.7±1.9% (data not shown). The difference in apoptotic rates was statistically significant (P<0.01).

Expression of the tight junction protein, occludin. Western blot analysis was used to investigate the expression of occludin (Fig. 6A), and the expression levels were quantified by densitometric analysis (Fig. 6B). Occludin expression was significantly downregulated in group H (0.42±0.011) compared with group C (1.31±0.084; P<0.05) in the tight junction of the epithelium in rats.

Discussion

Increasing evidence suggests that gastrointestinal tract injury causes MODS and there is an increase in the incidence of gastrointestinal diseases in patients who have travelled to a high altitude. Therefore, the initiation mechanism for altitude sickness is being investigated (16-18). In this context, a stable and effective animal model of high-altitude intestinal barrier injury is required.

There are a number of methods for establishing an animal model of gastrointestinal tract injury; however, none produce a model similar to acute hypoxia-induced intestinal barrier injury. Reasons for this include the species of animal selected; animals such as rabbits or pigs (19-21) are genetically different from humans and thus the results of these studies are not reliable to humans. Another reason is the production methods used impact the models; although the physical or chemical methods including surgery, trauma and pharmaceutical treatments (22,23) exert significant effects, models established by invasive methods do not mimic the conventional mechanism of hypoxia induced acute injury. Finally, the practical application affects the model. It has been demonstrated that when a rat is kept at a simulated 7,000 m height, the gastrointestinal tract is clearly injured (8). However, this method for establishing a model is different to the practical situation. The height limit for human survival is 5,000 m and 90% of humans live within an altitude range of 3,000-4,000 m. Thus, whether the environment at an altitude of 7,000 m induces a similar injury to an altitude of 4,000 m remains unclear. Furthermore, when producing a model at a simulated altitude of 7,000 m, the samples must be collected at an altitude <4,000 m. This may cause reperfusion injury or oxidative stress, which
potentially differs from single acute hypoxia-induced intestinal injury (24). Based on previous preliminary studies (15,25), only acute hypoxia occurring at an altitude of 3,000-5,000 m does not cause injury. The present study selected rats as the experimental animal, used a high-altitude environment simulation chamber to create a specific altitude environment and increased the exercise load to establish an animal model for intestinal barrier injury following acute hypoxia.

The animals were subjected to adaptive training at a normal altitude on an animal treadmill to avoid any coordination difficulty due to unfamiliar equipment. The rats received a sport load, to cause only slight fatigue stimulation and not cause pathophysiological changes associated with the gastrointestinal tract. Notably, the sport load is similar to the fatigue status of individuals who go to high-altitude regions by railway or road, making the model more applicable to humans.

Following training, the rats were kept in a simulated high-altitude environment chamber at an altitude of 7,000 m. Environmental factors were strictly controlled and food was supplied on ration to control the non-experimental factors that may alter the results. The altitude of 4,000 m is closer to the living altitude of humans. Therefore, after 3 days, the samples were collected at an altitude of 4,000 m, which ensured the safety of researchers and decreased the potential for secondary injury to rats from decreasing the altitude. This made the model more similar to the actual condition of hypoxia-induced intestinal injury.

MacConkey culture medium was used to identify the bacteria in each organ to judge whether it was from the intestine, following sample collection. As a selective culture medium, MacConkey culture medium contains the nutrient substances suitable for intestinal bacteria, which can partially inhibit gram-positive bacteria and completely inhibit the migrating growth of *Proteus bacillus vulgaris* (26). The addition of a pH indicator, neutral red and lactose, allows the pathogenic enterobacteria to be identified (27). The results indicated that there was bacterial translocation in group H but not in group C (P<0.05). This demonstrates that the modeling method may damage the intestinal mucous and cause intestinal bacterial translocation. The microscopy results indicated abnormalities in the intestinal epithelial ultrastructure, manifesting as epithelial villi falling off, shrinking and shortening, thinning of the epithelial mucosa, widening of inter-villi space and swelling and the disappearance of organelles in cytoplasm. Lanthanum nitrate-traced transmission microscopy was used to assess the degree of injury, as a high density of lanthanum...
granules was readily observed using electron microscopy. The diameter of the lanthanum granules was 3-4 nm and under normal conditions these would not enter the mesenchyme from the complete epithelium or microvascular tight junction. The results demonstrated that in group H, lanthanum granules were evenly distributed at epithelial cell tight junctions, basement membrane of vessels and intercellular space. This indicates that there was injury to the general structure and ultra micropathology of the intestinal barrier in the animal model. The lanthanum nitrate-traced transmission electron microscopy technique (28) observed that the lanthanum granules permeated into the deep layer from the opened epithelial cell tight junction. Thus, it is suggested that there is a correlation between the injury of epithelial barrier and damage to tight junction complexes. The tight junction of epithelial cells is a multi-functional complex consisting of many proteins (29,30), including occludin, claudins and actin. As a large protein family, there are a number of claudin subtypes with diverse functions (31,32). Therefore, monomeric occludin has become the preferred protein to detect (33-35). In the current study, western blot analysis performed to detect occludin in the rat intestinal epithelium following animal model establishment demonstrated that expression of occludin in tight junctions was downregulated. This indicates that the intestinal barrier function was weakened and permeability was increased, and verifies that the animal model was established successfully at a molecular level. However, it remains unclear how acute hypoxia induced damage to barrier function and this requires further analysis. In the future, occludin may be used to predict the...
permeability change of epithelial cells; however, further studies are required to confirm this.

In conclusion, the current study established an animal model for hypoxia-induced gastrointestinal injury in rats. The mechanism of inducement is similar to the pathophysiological process of the disease occurring under natural conditions. The establishment of this animal model may provide the basis for studies into possible treatments of intestinal barrier damage induced by hypoxia.

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