Effects of psychological stress on small intestinal motility and bacteria and mucosa in mice

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AIM: To investigate the effects of psychological stress on small intestinal motility and bacteria and mucosa in mice, and to explore the relationship between small intestinal dysfunction and small intestinal motility and bacteria and mucosa under psychological stress.

METHODS: Sixty mice were randomly divided into psychological stress group and control group. Each group were subdivided into small intestinal motility group (n = 10), bacteria group (n = 10), and D-xylose administered to stomach group (n = 10). An animal model with psychological stress was established housing the mice with a hungry cat in separate layers of a two-layer cage. A semi-solid colored marker (carbon-ink) was used for monitoring small intestinal transit. The proximal small intestine was harvested under sterile condition and processed for quantitation for aerobes (Escherichia coli) and anaerobes (Lactobacilli). The quantitation of bacteria was expressed as log_{10}(colony forming units/g). D-xylose levels in plasma were measured for estimating the damage of small intestinal mucosa.

RESULTS: Small intestinal transit was inhibited (39.80±9.50% vs 58.79±11.47%, P<0.01) in mice after psychological stress, compared with the controls. Psychological stress resulted in quantitative alterations in the aerobes (E. coli) in the proximal small intestinal flora (1.78±0.30 log_{10}(CFU/g) vs 1.37±0.21 log_{10}(CFU/g), P<0.01), and there was decrease in relative proportion of Lactobacilli and E. coli of stressed mice (0.53±0.63 vs 1.14±1.07, P<0.05), while there was no significant difference in the anaerobes (Lactobacilli) between the two groups (2.31±0.70 log_{10} (CFU/g) vs 2.44±0.37 log_{10}(CFU/g), P>0.05). D-xylose concentrations in plasma in psychological stress mice were significantly higher than those in the control group (2.90±0.89 mmol/L vs 0.97±0.33 mmol/L, P<0.01).

CONCLUSION: Small intestinal dysfunction under psychological stress may be related to the small intestinal motility disorder and dysbacteriosis and the damage of mucosa probably caused by psychological stress.
was purchased from NanJing Jiancheng Bioengineering Institute (NJBI).

Methods

Establishment of animal model Sixty mice were randomly divided into psychological stress group and control group. Each group were subdivided into small intestinal motility group (n = 10) and bacteria group (n = 10) and D-xylose administered into stomach group (n = 10). Mice in psychological stress group were housed in the bottom of the two-layer cage, with a hungry cat being housed in the proximal layer of the cage for 10 min each day for 15 d, but mice and the cat had no physical contact. Procedure of the control group mice was as same as psychological stress group except for no contact with the cat.

Measurement of small intestinal transit The carbon-ink transit test was modified as described. Mice were deprived of food for 24 h and water for 12 h prior to experiment, and 0.3 mL carbon-ink (10% gum acacia, 5% activated charcoal) was administered into stomach by orogastric gavage. Twenty-five minutes later, the mice were killed, abdomen was opened and small intestine was dissected. The total length of the small intestine (pylorus-cecum) and the distance traveled by carbon-ink were measured. Results were expressed as ratio (%) of the distance traveled by carbon-ink to the total length of the small intestine.

Measurement of small intestinal bacteria Mice were deprived of food for 24 h and water for 12 h prior to experiment. The mice were killed, abdomen was opened and the proximal small intestine was harvested under sterile condition. The 2-cm-long small intestine, which was dissected at about a 10-cm point from pylorus was rinsed with sterile saline thrice, and then the leftover was sucked by sterile filter paper. After weighing, the small intestine with 2 mL sterile saline was placed in a sterile glass homogenizer and homogenized. Homogenate was diluted with sterile saline at different ratios and 100 μL dilution was plated on SS agar (Escherichia coli) and MRS agar (Lactobacilli). Quantity of E. coli was determined after 24 h of incubation at 37 ℃. Quantity of Lactobacilli was determined after 48 h of incubation at 37 ℃. Colony forming units (CFU) of bacteria were quantified by counting CFU from agar. The quantity of bacteria was expressed as log_{10}(CFU/g). Quality of aseptic manipulation was evaluated by swab of abdominal cavity inoculating on sheep-blood agar.

Measurement of D-xylose concentrations in plasma Mice were deprived of food for 24 h and water for 12 h prior to experiment, and 0.4 mL 5% D-xylose solution was administered into stomach by orogastric gavage. One hour later, blood samples were collected into chilled tubes containing 100 U heparin immediately after the mice were killed. The blood was centrifuged at 3 000 r/min at 4 ℃ for 10 min. The plasma was stored at -70 ℃ until assayed. Levels of D-xylose in plasma were measured with D-xylose kit.

Statistical analysis Throughout this report, data were expressed as mean±SD. Experimental results were analyzed by t test. P<0.05 was considered statistically significant.

RESULTS

Small intestinal transit

Figure 1 presents data for the overall mean ratio of small intestinal transit (percentage of the distance traveled by intragastric carbon-ink to the total length of the small intestine after 25 min). The overall mean ratio of small intestinal transit under psychological stress was lower than that of the control (39.80±9.50% compared with 58.79±11.47%, P<0.01), indicating that psychological stress could inhibit small intestinal transit.

Measurement of D-xylose concentrations in plasma

D-xylose concentrations in plasma in psychologically stressed mice were significantly higher than those in the control group (2.90±0.89 mmol/L compared with 0.97±0.33 mmol/L, P<0.01), indicating that small intestinal mucosa was damaged (Figure 2).
stress on the stomach and the small intestine, but had no demonstrable effect on the microflora. Dulas et al[7], demonstrated a distinct decrease in the numbers of Bifidobacterium in gnotobiotic mice suppressed by restraint stress. In the present study, we demonstrated that restraint stress but not by footshock stress. And footshock stress increased small intestinal motility both during fasting and after food. But restraint stress completely inhibited the small intestinal transit was significantly inhibited by restraint stress but not by footshock stress. And footshock stimulation may cancel the inhibition of small intestinal motility by restraint stress. In the present study, we demonstrated that the ratio of small intestinal transit was significantly decreased by psychological stress.

Host and intestinal normal flora are in a state of balance. Varied microorganisms of intestinal normal flora interact to keep themselves in a state of balance too. Changes of host and external environment could lead to disequilibrium. Itoh and Freter demonstrated that the Lactobacilli in gnotobiotic mice suppressed E. coli multiplication in the stomach and the small intestine, but had no demonstrable effect on E. coli multiplication in the large intestine. Stress can cause significant change of intestinal flora. Lizko[8,9] revealed the factors that are neuroemotional tension, hypokinesia, increased physical load, isolation under conditions of altered gaseous environment and microclimate participated in development of dysbacteriosis under extreme conditions (space flights of various duration). Intestinal microflora in man responded by decreased counts of bifidobacteria and Lactobacilli participating in maintenance of the intact ecological barrier and colonization resistance. Furthermore, the neuroemotional stress played the main role in development of dysbacteriosis in man under extreme conditions, which was supported by the following study. During the preparation phase of the flight and the period immediately before the take-off and after the flight, there was a distinct decrease in the numbers of Bifidobacterium and Lactobacilli as well as a substantial increase in the numbers of E. coli. This seems to be due to nervous-emotional stress effects. Logan et al[11], reported that patients with chronic fatigue syndrome (CFS) had marked alterations in microbial flora, including lowered levels of bifidobacteria and small intestinal bacterial overgrowth (SIBO). Bailey and Coe[12] reported that maternal separation disrupts the integrity of the intestinal microflora in infant rhesus monkeys. Fecal bacteria decrease significantly, especially Lactobacilli. The drop in the microflora was correlated with the display of stress-indicative behaviors, but not with cortisol secretion. In addition, infants who displayed numerous stress-indicative behaviors were more susceptible to opportunistic bacterial infection. Gritsenko et al[13], revealed that 6-h immobilization stress initiates the increase of the concentration of E. coli in the proximal sections of the digestive tract (the duodenum and the jejunum). Ringo et al[14], reported that the total culturable bacterial numbers or population level of the lactic acid bacteria associated with the digestive tract of Atlantic salmon (Salmo salar L.) had no significant alteration under excessive handling stress and starvation. In the present study, the results were similar to the above research. After psychological stress, the number of E. coli of the proximal small intestine in mice increased significantly compared to the control. The number of Lactobacilli was not statistically significant and different from the control. The ratio of Lactobacilli and E. coli was lower than that of the control. Glunder[15] demonstrated that stressful situations such as overcrowding in small cages coincident with increased noise and low light levels can enhance the colonization of the gut with E. coli. The present study was involved in stressful situations because all of the mice were placed in the cage when they were under stress. Lenzen et al[16], reported that the lactoflora of cosmonauts showed distinct changes due to the emotional stress before the take off. Compared to long-time flights, short-time flights caused even stronger alterations of lactoflora. The reason was that the disturbance that took place before the take-off as to the short time could not be balanced by the macroorganism. Additionally, after a certain adaptive period the factors of the space travel began to act. The reason may be the explanation that there was no difference in the number of Lactobacilli of small intestine between the stressed mice and the control in the present study.

Leveau et al[17], demonstrated that impairment in intestinal motility probably played a pathophysiological role in the development of bacterial overgrowth. A delay in intestinal transit time appeared as an early event in acute pancreatitis, preceding intestinal bacterial overgrowth. Gangarosa[18] demonstrated that intestinal motility served as a normal cleansing mechanism of the intestine, and drugs that decreased this motility might facilitate replication of pathogens and their attachment to or infection of the intestinal tissue. Wang et al[19-21], demonstrated that delayed intestinal transit after hepatectomy might contribute to overgrowth of E. coli in small intestine. Administration of cisapride or CCK prevented overgrowth of E. coli by improving intestinal motility in rats.

Interdigestive small bowel motility has a regulatory function on the microflora of the upper small bowel. Nieuwenhuijs et al[22,23], clarified the role of the migrating motor complex (MMC) in the regulation of small intestinal motility in man under extreme conditions of altered gaseous environment and microclimate.
Chronic water avoidance stress further increased permeability of IBS. Madrid could explain occurrence and development of some symptoms seems to result in some normalization of motility. The above was still present had less phase III events than subjects with transit.

Intestinal bacteria promoted regular spike burst activity primitive fermenting metabolism (anaerobes) emerge as aboral migration of the migrating myoelectric complex (MMC) in proximal jejunum of germ-free rats was reduced. Intestinal bacteria promoted or suppressed the initiation and complex in proximal jejunum of germ-free rats was reduced. The growth of \( E. coli \) stimulated by the electric current was significantly inhibited after a period of intensive growth.

Intestinal microflora can modulate myoelectric activity of small intestine. Husebye et al\(^{[27]} \), demonstrated that after introduction of conventional intestinal microflora the interval between activity fronts of the migrating myoelectric complex in proximal jejunum of germ-free rats was reduced. Intestinal bacteria promoted or suppressed the initiation and aboral migration of the migrating myoelectric complex (MMC) depending on the species involved. Bacteria with primitive fermenting metabolism (anaerobes) emerge as important promoters of regular spike burst activity in small intestine. Lactobacillus promoted regular spike burst activity and reduced the MMC period and accelerated small intestinal transit. \( E. coli \) showed an inhibitory effect on MMC. Thorlacius et al\(^{[28]} \), reported that Lactobacillus could enhance intestinal transit. Sjogren et al\(^{[29]} \), demonstrated that bacterial adherence to the intestinal mucosa appeared to be important in eliciting the abnormal myoelectric responses. \( E. coli \) prolonged spike bursts. Tsafarov et al\(^{[30]} \), demonstrated that \( E. coli \) inhibited the movements of the intestine, which was manifested by a reduction of the frequency and amplitude of the intestinal contractions. Cuoco et al\(^{[31]} \), reported that bacterial overgrowth might contribute to the delay of intestinal transit. After eradication therapy, patients without bacterial overgrowth showed a significant reduction of oro-cecal transit time. Pimentel et al\(^{[32]} \), demonstrated that the duration and frequency of phase III was reduced in subjects with irritable bowel syndrome (IBS) and SIBO. Subjects whose SIBO was still present had less phase III events than subjects with eradicated overgrowth. Eradication of bacterial overgrowth seems to result in some normalization of motility. The above can explain occurrence and development of some symptoms of IBS. Madrid et al\(^{[33]} \), administrated patients with liver cirrhosis prokinetics (cisapride) or antibodies. After 6-mo treatment, both cisapride and antibodies significantly improved fasting cyclic activity, reduced the duration of oro-cecal transit time, and decreased SIBO. Cisapride administration was also followed by an increase in the amplitude of contractions.

Meddings and Swain\(^{[34]} \) demonstrated that psychological stress might increase permeability of all regions of the gastrointestinal tract. This provides a potential mechanism for the observation of stress-induced disease recurrence in Crohn’s disease. Velin et al\(^{[35]} \), reported that the barrier function of follicle-associated epithelium could be modulated. Chronic water avoidance stress further increased permeability of villus and follicle-associated epithelium than acute water avoidance stress. Saunders et al\(^{[36]} \), reported that acute stress altered jejunal epithelial physiology. Both physical stress (2 h of cold-restraint stress) and psychological stress (1 h of water-avoidance stress) increased ionic and macromolecular permeability of intestine. Wilson and Baldwin\(^{[37]} \) demonstrated that mental stress (environmentally induced stress) caused pathological changes in the rat intestinal mucosa, which compromise the epithelial-endothelial exchange barrier, where intestinal villi were edematous and epithelial cells were detaching from the basement membrane at villus tips. Santos et al\(^{[38]} \), reported that chronic stress (water avoidance stress or sham stress (1 h/d) for 5 d) causes an epithelial barrier defect and epithelial mitochondrial damage. Soderholm et al\(^{[39]} \), demonstrated that chronic psychological stress could be an initiating factor in intestinal inflammation by impairing mucosal defenses against luminal bacteria. Chronic stress (water avoidance stress or sham stress as a model of ongoing life stress) induced barrier dysfunction in the ileum and colon (increased macromolecular permeability and depletion of mucus) and ultrastructural changes in epithelial cells (enlarged mitochondria and presence of autophagosomes) associated with bacterial adhesion and penetration into enterocytes. Shi et al\(^{[40]} \), reported that chronic restraint stress could cause damage on intestinal barrier function, increased intestinal permeability to D-xylene. The levels of D-xylene in plasma of stressed rats were higher than that of the control group. The result of our study showed that concentration of D-xylene in plasma of mice subjected to psychological stress was significantly higher than the control, which suggested that psychological stress caused damage of small intestinal mucosa and increased permeability to luminal substances.

Schiffrin et al\(^{[41]} \), suggested that the developmental condition of the host’s intestinal barrier might be an important regulator of the bacterial microenvironment of the newborn small intestinal mucosa. Garcia-Lafuente et al\(^{[42]} \), demonstrated that certain commensal bacteria could modify colonic wall permeability to luminal substances. \( E. coli \) significantly increased lumen to blood clearance. Colonization with Lactobacillus had the opposite effect and reduced permeability to mannitol. Isolauri et al\(^{[43]} \), reported that Lactobacillus counteracted permeability disorder of the mucosal barrier. Logan et al\(^{[44]} \), demonstrated that Lactobacillus might have a therapeutic role in the treatment of chronic fatigue syndrome (CFS). Lactobacillus are strong antioxidants, can enhance absorption of micronutrients by protecting the intestinal epithelial barrier, and have been used to treat SIBO. Bomba et al\(^{[45]} \), suggested that the Lactobacillli-produced organic acids might present an efficient barrier inhibiting the adherence of digestive tract pathogens to the intestinal mucosa. Lievin-Le Moal et al\(^{[46]} \), reported that Lactobacill isolated from the resident adult human gastrointestinal microflora, together with its antimicrobial activity, exerts a protective effect against the brush border lesions promoted by adhering \( E. coli \) in human intestinal cells. Ding et al\(^{[47]} \), reported that increased \( E. coli \) caused the magnitude of gut mucosal injury. Zarrie et al\(^{[48]} \), reported that gram-negative luminal bacteria could cause significant alterations in epithelial ion transport and barrier functions. McNamara
et al[40] clarified that E. coli disrupts intestinal barrier function, increased monolayer permeability and redistributed the tight junction-associated protein occludin. Perers et al [40], reported that E. coli damaged the brush border of the mucosal epithelium. Muza-Moons et al[41], reported that E. coli disrupted the structure and barrier function of host intestinal epithelial tight junctions (TJs). Michail et al[42], indicated that E. coli-induced neutrophil migration could occur without significant disruption of barrier function.

Gangarosa[53] demonstrated that intestinal motility served as a normal cleansing mechanism of the intestine, and drugs that decreased this motility might facilitate replication of pathogens and their attachment to or invasion of the intestinal tissue. Bacterial adherence to epithelia of intestinal mucosa plays an important role in the interaction of host and bacteria. Rocha et al[44], demonstrated that mucosa-associated E. coli played specific role on epithelial barrier dysfunction. Stress increased numbers of mucosa-associated E. coli in the cecum, which could increase epithelial permeability. E. coli of mice submitted to stress adhered to and altered the permeability of young adult mouse colon cells, whereas E. coli from the cecum of control mice were less adherent and had no effect on epithelial permeability.

Previous studies have shown that psychological stress can cause gastrointestinal dysfunction. For example, patients with gastrointestinal motility disorders, especially irritable bowel syndrome (IBS), symptoms of abdominal pain and bloating were precipitated after psychological stress (life events). Ford et al[45], demonstrated that anxiety resulting from psychological stress could enhance the colonic symptoms of gas. The defect of intestinal barrier and the above factors can make symptoms become more serious. These aspects of gas. The defect of intestinal barrier and the above factors can cause gastrointestinal dysfunction in humans.

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