Effects of psychological stress on small intestinal motility and expression of cholecystokinin and vasoactive intestinal polypeptide in plasma and small intestine in mice

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Abstract

AIM: To investigate the effects of psychological stress on small intestinal motility and expression of cholecystokinin (CCK) and vasoactive intestinal polypeptide (VIP) in plasma and small intestine, and to explore the relationship between small intestinal motor disorders and gastrointestinal hormones under psychological stress.

METHODS: Thirty-six mice were randomly divided into psychological stress group and control group. A mouse model with psychological stress was established by 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. CCK and VIP levels in plasma and small intestine in mice were measured by radioimmunoassay (RIA).

RESULTS: Small intestinal transit was inhibited (52.18±19.15% vs 70.19±17.79%, P<0.01) in mice after psychological stress, compared to the controls. Small intestinal CCK levels in psychological stress mice were significantly lower than those in the control group (0.75±0.53 μg/g vs 1.98±1.17 μg/g, P<0.01), whereas plasma CCK concentrations were not different between the groups. VIP levels in small intestine were significantly higher in psychological stress mice than those in the control group (8.45±1.09 μg/g vs 7.03±2.36 μg/g, P<0.01), while there was no significant difference in plasma VIP levels between the two groups.

CONCLUSION: Psychological stress inhibits the small intestinal transit, probably by down-regulating CCK and up-regulating VIP expression in small intestine.

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Key words: Small intestine; Psychological stress; Cholecystokinin; Vasoactive intestinal polypeptide; Intestinal motility

INTRODUCTION

In recent years, with rapid development of society and increasing competition pressure, psychological stress, which originates in society has imposed impacts on human health, and has become an important stressor. Psychological stress is widely believed to play a major role in gastrointestinal motor disorders, especially in irritable bowel syndrome (IBS) and functional dyspepsia (FD), by precipitating exacerbation of symptoms[12]. Available data clearly demonstrate that in experimental animals the most consistent patterns of gastrointestinal motor alternations induced by psychological stress are delayed gastric emptying[3,4] and accelerated colon transit[5,6]. However, almost no valid data are available on small intestinal motility disorders. Previous studies of psychological stress[7-9] (cold stress, restraint stress, foot-shock stress and swim stress), were all focused on somatic stress. In common, both central and peripheral nervous pathways are involved in the release of gastrointestinal hormones due to psychological stress, thus modulating gastrointestinal motility[10]. A large body of evidence derived from experiments suggest that CCK could accelerate small intestinal transit[11,12], while VIP could inhibit it[13]. However, there are few studies on whether gut hormones are involved in modulating small intestinal motility under psychological stress.

In the present study, experimental animals were obtained to test the relationship between small intestinal motility and release of gastrointestinal hormones during psychological stress. Furthermore, small intestinal motility disorders in response to CCK and VIP due to psychological stress were studied.

MATERIALS AND METHODS

Experimental animals and materials

Thirty-six healthy male mice (provided by Qinglongshan Experimental Animal Center) weighing 20-30 g were used in this study. Mice were housed individually in cages at constant room temperature with a 12-h light/dark cycle and had free access to laboratory chow and water. CCK kit was purchased from Neurobiological Department of Second Military Medical University. VIP kit was purchased from Beijing Haikerui Biological Technology Center.

Establishment of animal models

Thirty-six mice were randomly divided into psychological stress group and control group. Each contained 18 mice. Mice in psychological stress group were housed in the bottom of two-layer cage, with a hungry cat housed in the upper layer of the cage for 10 min each day for 10 d, but mice and the cat had no physical contact[14]. Procedure for the control group mice were the 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 0.25 mL carbon-ink was administered into their stomachs 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 (percentage) of the distance traveled by carbon-ink to the total length of the small intestine. Small intestine was washed with normal saline after measuring. Water was absorbed by filter paper and small intestine was kept in a dry bottle.

**Plasma and tissue preparation**

Immediately after the mice were killed, blood samples were collected into chilled tubes containing 0.3 μL ethylenediamine tetraacetic acid (EDTA) and 1000 KIU aprotinin. The blood was centrifuged at 3 000 r/min at 4 °C for 10 min. The plasma was stored at -70 °C until assayed.

Samples of small intestine were placed in saline and boiled for 10 min. Water was absorbed by filter paper. Small intestine was weighed by analytical balance and then dissolved in 1 mmol/L ice-cold acetic acid (0.3 mL/100 g), homogenized and then they were left at room temperature for 20 min, and centrifuged at 3 000 r/min at 4 °C for 10 min. The supernatants were stored at -70 °C until assayed.

**CCK radioimmunoassay**

Plasma and small intestine CCK levels were measured using radioimmunoassay (RIA). Briefly, samples and standards diluted in the perfusion medium were incubated with CCK antiserum in tubes for 24 h at 4 °C. After addition of 125I-CCK, all samples were further incubated for 24 h at 4 °C to reach equilibrium. Antibody-bound and free tracers were separated by addition of 0.4 mL of activated charcoal.

**VIP radioimmunoassay**

VIP was determined by radioimmunoassay. Samples and standards diluted in the perfusion medium were incubated with VIP antiserum in tubes for 48 h at 4 °C. After addition of 125I-labeled VIP, all samples were further incubated for 48 h at 4 °C. After incubation, antibody-bound and free tracers were separated by addition of 0.4 mL of activated charcoal.

**Statistical analysis**

Data were expressed as mean±SD. Experimental results were analyzed by t test, P<0.05 was considered statistically significant.

**RESULTS**

**Small intestinal transit**

After 25 min of intragastric carbon-ink administration, the overall mean ratio of small intestinal transit (P<0.01) under psychological stress was lower than that of the control (52.18±19.15% vs 70.19±17.79%), indicating that psychological stress could inhibit small intestinal transit. Figure 1 presents individual data for the ratio of small intestinal transit (percentage of the distance traveled by carbon-ink to the total length of the small intestine).

**CCK concentrations in plasma and small intestine**

CCK levels in small intestine of experimental psychological stress mice were significantly lower than those of the control (0.75±0.53 μg/g vs 1.98±1.17 μg/g, P<0.01). However, no significant changes in plasma CCK levels were observed in experimental psychological stress mice compared to the control (53.88±33.17 ng/L vs 52.70±20.10 ng/L, P>0.05) (Table 1).

**Table 1** Changes of CCK in plasma and small intestine (mean±SD)

| Group     | No. of animals | Small intestine (μg/g) | Plasma (ng/L) |
|-----------|----------------|------------------------|---------------|
| Control   | 18             | 1.98±1.17              | 52.70±20.10   |
| Stress    | 18             | 0.75±0.53<sup>b</sup> | 53.88±33.17   |

<sup>b</sup>P<0.01 vs the control.

**VIP concentrations in plasma and small intestine**

VIP levels in small intestine of experimental psychological stress mice were significantly higher than those of the control (8.45±1.09 μg/L vs 7.03±2.36 μg/L, P<0.01). However, plasma VIP levels showed no significant difference between the two groups (201.58±103.23 μg/L vs 190.05±90.08 μg/L, P>0.05).

**Table 2** Changes of VIP in small intestine plasma (mean±SD)

| Group     | No. of animals | Small intestine (μg/g) | Plasma (ng/L) |
|-----------|----------------|------------------------|---------------|
| Control   | 18             | 7.03±2.36              | 190.05±90.08  |
| Stress    | 18             | 8.45±1.09<sup>b</sup> | 201.58±103.23 |

<sup>b</sup>P<0.01 vs the control.

**DISCUSSION**

The association between psychological stress and small intestinal motility has been postulated for about twenty years. Some studies in experimental animals indicate contradictory results, and the mechanism by which psychological stress affects small intestinal motility is not well understood. Ditto et al<sup>[16]</sup> recently reported that stress enhanced small intestinal transit and produced a statistically significant reduction in the mean small intestinal transit time. However, Tsukada et al<sup>[6]</sup> demonstrated that psychological stress induced inhibition of small intestinal transit. In the present study, our results were similar to Tsukada’s. After 25 min of intragastric carbon-ink administration, the ratio of small intestinal transit significantly decreased during psychological stress. Previous studies show that patients with gastrointestinal motility disorders, especially irritable bowel syndrome (IBS), symptoms of abdominal pain and bloating are precipitated after psychological stress (life events or trauma). The mechanism may be that psychological stress induces decrease in small intestinal transit. Intestinal gas could not be evacuated effectively leading to gas retention, which evokes a series of symptoms, especially pain and bloating<sup>[17]</sup>.

In this study, we showed that CCK levels in small intestine of psychological stress mice were significantly lower than those of the controls, while they did not alter in plasma. CCK, a peptide secreted from I cells of gastrointestinal responds to the presence of nutrients in the small intestine. It is also synthesized in...
brain[28]. Two CCK receptor subtypes that differ markedly in their pharmacological profiles and anatomical distribution have been identified and characterized[29]. CCK-A receptors exist predominantly at the peripheral level where they are responsible for the digestive effects of CCK[30]. However, high densities of brain CCK-B receptors are present in cortical and limbic blood. Endogenous CCK appears to act in part by paracrine and endocrine mechanisms to affect gut motility[21]. It is well established that endogenous CCK regulates gastrointestinal motility, including delay of gastric emptying[22], acceleration of small intestinal transit[23] and enhancement of colon motor[24]. Various studies have provided evidence for the mechanism of CCK underlying small intestinal motility through involvement of CCK-A receptors in small intestine. The selective CCK-A receptor antagonists (devazepide and lorglumide) effectively attenuate this effect while CCK-B receptor antagonists have no effect on it[25]. Furthermore, both afferent and effenter nerves are implicated in CCK motor actions in the gastrointestinal tract. Patterson et al[27] performed immunohistochemistry using an antibody directed to the C-terminal region of the CCK-A receptor, and found that intestinal cells of Cajal (ICC) in the sphincter muscle and the circular muscle of proximal duodenum displayed strong CCK-A receptor immunoreactivity; at the same time, selection neurons in the myenteric ganglia and a few of single neurons in the submucosa near the proximal duodenum, also expressed strong CCK-A receptor immunoreactivity. In view of these results, we can conclude that the effect of psychological stress on intestinal motility and transit may be partially mediated by decreasing intraluminal concentrations of CCK, which leads to reduced migrating myoelectric complex (MMC) activity in the small intestine, prolonging MMC cycle length, reducing contraction frequency and decreasing propagation velocity.

VIP, which is released from D, cells of gut through exocrine into the lumen, has an established role in the regulation of gastrointestinal motor function. VIP is widely distributed in gut. In addition to GI mucosa[28], it is also found in submucous and myenteric plexus, central and peripheral nervous systems[29,30]. Furthermore, a large number of VIP receptors are expressed in those regions[31,32]. Sayadi et al[33] analyzed the distribution of VIP receptors using autoradiographic techniques, and found that the density of VIP receptors was greatest in duodenal mucosa but was lower in jejunal and ileal mucosa. Moreover, previous studies have provided evidence that VIP acts via two receptors (VIP1 and VIP2). In the intestine, VIP receptor is significantly higher than VIP2 receptor[34]. A great number of data demonstrate that VIP has various physiologic functions on small intestinal motility. First, VIP exerts direct stimulatory effect on intestinal smooth muscle, reduces the percentage motor index of pressure waves. This effect of VIP could be partially antagonized by VIP receptor antagonists[35]. Second, intraduodenal infusion of VIP in rats disrupts the MMC, prolongs MMC cycle length and decreases propagation velocity, while VIP receptor antagonists could reverse this effect[36,37]. Third, the effect of VIP on intestinal motility involves both myenteric and submucosal neurons, resulting in increased contraction of longitudinal smooth muscle[38]. Naslund et al[39] demonstrated that intraduodenal but not plasma concentrations of VIP showed an association with the MMC. In the present study, we showed that VIP levels in small intestine were significantly higher than those of the control, but had no difference in plasma. It indicates that small intestine VIP plays an important role in the inhibition of small intestinal transit caused by psychological stress.

The results of this study suggest that gastrointestinal motility disorders during psychological stress may be partially mediated by release of gut hormones from small intestine. But the following questions need to be answered in future: how does psychological stress modulate gut hormone release from small intestine and how are gastrointestinal motility disorders caused by gut hormones in small intestine.

In summary, psychological stress does induce changes in the small intestinal motility. Changes of CCK and VIP levels in the small intestine of mice may be closely related with the inhibition of small intestine transit. However, there is no relationship between gastrointestinal dysmotility and gut hormones in plasma during psychological stress.

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