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**Effects of Lizhong Tang on gastrointestinal motility in mice**

Lee MC*et al.*Lizhong Tang and GI motility

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**Abstract**

**AIM:** To investigate the effects of Lizhong Tang, a traditional Chinese medicine formula, on gastrointestinal motility in mice.

**METHODS:** The *in vivo* effects of Lizhong Tang on GI motility were investigated by measuring the intestinal transit rates (ITRs) and gastric emptying (GE) values in normal mice and in mice with experimentally induced GI motility dysfunction (GMD).

**RESULTS:** In normal ICR mice, the ITR and GE values were significantly and dose-dependently increased by Lizhong Tang (ITR values: 54.4 ± 1.9% *vs* 65.2 ± 1.8%, *P* < 0.01 with 0.1 g/kg Lizhong Tang and 54.4 ± 1.9% *vs* 83.8 ± 1.9%, *P* < 0.01 with 1 g/kg Lizhong Tang; GE values: 60.7 ± 1.9% *vs* 66.8 ± 2.1%, *P* < 0.05 with 0.1 g/kg Lizhong Tang and 60.7 ± 1.9% *vs* 72.5 ± 1.7%, *P* < 0.01 with 1 g/kg Lizhong Tang). The ITRs of the GMD mice were significantly reduced compared with those of the normal mice, which were significantly and dose-dependently reversed by Lizhong Tang. Additionally, in loperamide- and cisplatin-induced models of GE delay, Lizhong Tang administration reversed the GE deficits.

**CONCLUSION:** These results suggest that Lizhong Tang may be a novel candidate for development as a prokinetic treatment for the GI tract.

**Key words:** Lizhong Tang; Gastrointestinal disorders; Motility; Intestinal transit rate; Gastric emptying

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**Core tip:** Lizhong Tang, a traditional Chinese medicinal formula, has been widely used in China, Japan, and Korea for many years to ameliorate gastrointestinal (GI) disorders. Our data suggest that Lizhong Tang is a novel candidate for development as a prokinetic agent for treatment of GI motility dysfunctions in man.

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**INTRODUCTION**

Lizhong Tang, also known as Yijung-tang or Richu-to, is a traditional Chinese medicine (TCM) formula[1] and is composed of Radix Ginseng (*Panax ginseng* C.A. Meyer), Rhizoma Zingiberis (*Zingiber officinale* Roscoe), Rhizoma Atractylodis Macrocephalae (*Atractylodes macrocephala* Koidz.), and Radix Glycyrrhizae (*Glycyrrhiza uralensis* Fisch)[2]. Lizhong Tang is widely used in traditional medicine to treat gastrointestinal (GI) disorders, such as vomiting, diarrhea, stomach pain, chronic gastritis, stomach bleeding, and GI ulceration, in China, Japan, and Korea[1-3]. However, no studies have been conducted to evaluate the effect of Lizhong Tang on GI motility.

Prokinetic agents are medications that enhance coordinated GI motility and the transit of content in the GI tract mainly by amplifying and coordinating the GI muscular contractions. In addition, prokinetic therapy should be considered as a means to improve gastric emptying and symptoms of gastroparesis, balancing the benefits and risks of treatment[4]. Recently, prokinetic therapy has been shown to improve the symptoms and quality of life in patients with GI motility disorders[5]. Therefore, there has been an increasing need to develop safer and more effective gastroprokinetic agents.

In our previous report, we investigated the effects of Lizhong Tang on mouse small intestine interstitial cells of Cajal (ICC)[6]. These cells are the pacemaker cells of GI muscles and generate rhythmic oscillations in membrane potentials known as slow waves[7-9] by activating Ca2+ entry through L-type Ca2+ channels in smooth muscles to initiate GI contractions[10,11]. In this report, we found that Lizhong Tang affected GI motility by modulating pacemaker activity in ICC through internal Ca2+- and phospholipase (PLC)-dependent pathways[6]. However, despite the widespread use of Lizhong Tang to treat GI disorders, little is known about its regulatory effects on GI motility. Therefore, we performed this study to investigate the effects of Lizhong Tang on the mouse GI tract *in vivo*.

**MATERIALS AND METHODS**

***Preparation of the standard solutions and sample extracts***

Liquiritin, 6-gingerol, ononin, glycyrrhizin, ginsenoside Rg1, isoliquiritin, and atractylenolide III were accurately weighed and dissolved in methanol (all at 100 μg/mL) to prepare standard solutions. Lizhong Tang powder was dissolved in the water and then filtered through a 0.2 μm syringe filter (BioFACTTM, Korea) prior to injection in the HPLC.

***Chromatographic conditions***

An Agilent 1200(Agilent Technologies, Palo Alto, CA, United States) equipped with an autosampler, degasser, quaternary solvent pump, and diode array detector (DAD) was used for the analysis. The data were acquired using ChemStation software (Agilent Technologies, Palo Alto, CA, United States). Separation was performed on a Capcell Pak Mg II C18 column (4.6 mm × 250 mm, 5 μm; Shiseido, Tokyo, Japan) at 35 °C. The mobile phase consisted of water containing 0.1% trifluoroacetic acid (A) and acetonitrile (B), and gradient elution was conducted as follows: 5% (B) for 0–1 min; 5%–10% (B) for 1–5 min and held for 5 min; 10%–15% (B) for 10–12 min and held for 4 min; 15%–20% (B) for 16–18 min and held for 4 min; 20%–23% (B) for 22–25 min and held for 6 min; 23%–28% (B) for 31–32 min and held for 8 min; and 28–65% (B) for 40–80 min and held for 2 min. The column was then re–equilibrated using 5% (B) for the subsequent analyses. The flow rate was set at 1.0 ml/min, and the detection wavelengths were 205, 230, 250, 280, and 360 nm.

***Animals***

Male ICR mice (Samtako BioKorea Co., Ltd., Osan, Republic of Korea) weighing 23–30 g were used to investigate the effects of the Lizhong Tang extract on the GI tract *in vivo*. The animals were maintained under controlled conditions (21 ± 3°C, relative humidity 50 ± 6%, lights on 6 a.m.–6 p.m.). The mice were allowed free access to a commercial diet and tap water, but were fasted for 24 h before the experiments. All experiments were conducted between 10 a.m. and 6 p.m.

***Measurement of ITR using Evans blue staining***

We used Evans blue solution (5%, w/v, in distilled water (DW)) to determine the ITRs of the Lizhong Tang extract *in vivo*. The Evans blue solution was administered (0.1 mL/kg of body weight; i.g.) through an orogastric tube 30 min after the Lizhong Tang extract was intragastrically (i.g.) administered to the normal ICR mice. The animals were sacrificed 30 min after Evans blue administration, and the intestinal transit distances of the dye were determined by measuring the distance the Evans blue dye had migrated in the intestine from the pylorus to its most distal point. Intestinal transit was quantified using the intestinal transit rates (ITR) (%), which were calculated by expressing the distance the Evans blue dye traveled in 30 min as a percentage of the total small intestine length (from the pylorus to the ileal terminus).

***Induction of GI motility dysfunction in mice***

Two experimental GI motility dysfunction models were used: an acetic acid (AA)-induced peritoneal irritation mouse model and a STZ-induced diabetic mouse model. For the AA model, peritoneal irritation was induced by administering AA to ICR mice 30 min after the i.g. administration of the Lizhong Tang extract (or DW as vehicle) by intraperitoneally (i.p.) injecting 10 ml/kg AA (0.6%, w/v, in saline) as previously described[12-14]. After injecting AA, the mice were placed in individual cages and allowed to recover for 30 min. Male ICR mice (aged 5 wk) were used for the STZ-induced diabetic mouse model. The mice were randomly allocated to two groups: a control group or a diabetic group. The mice were fasted overnight and an STZ (Sigma-Aldrich, St. Louis, MO) solution was administered i.p. on the following day to produce diabetes. Fresh STZ was prepared in 0.1 mol/L ice-cold citrate buffer (pH 4.0) and administered at 200 mg/kg body weight[15]. The control mice were i.p. administered the same volume of 0.1 mol/L citrate buffer. The animals had free access to food and water and were maintained under standard conditions (24–27 °C, RH 60%–65%) under a 12 h light/dark cycle. Two months after the STZ injection, blood was withdrawn from a tail vein after an 8 h fast and the blood glucose concentrations were measured using a ONE-TOUCH Select Simple kit (Johnson and Johnson Medical Company). Diabetes was defined as a blood glucose level of > 16 mmol/L. No mortality occurred during the study period, and no mouse recovered from STZ-induced diabetes.

***Evaluation of gastric emptying***

As previously described by Scarpignato *et al*[16], the mice were fasted for 24 h with free access to water[14]. Gastric emptying (GE) was performed by administering a 0.05% (w/v) phenol red solution (0.5 mL/mouse) 30 min after treatment with the Lizhong Tang extract. Twenty min later, the mice were sacrificed and the stomachs were immediately removed, cut into several pieces, placed into 5 mL of 0.01 N NaOH, and homogenized. The homogenates were treated with 0.2 mL of 20% trichloroacetic acid per mL of homogenate. The mixtures were centrifuged for 10 min at 1050 × *g*, and the supernatants (0.05 mL) were added to 0.5 N NaOH (0.2 mL). The absorbances of these mixtures were measured using a spectrometer at 560 nm. The GE value (%) was calculated as 100-(A/B) × 100, where A is the test stomach absorbance (560 nm) and B is the control stomach absorbance (560 nm) immediately after phenol red administration.

***Drugs***

All drugs were purchased from Sigma-Aldrich (Sigma-Aldrich, St. Louis, MO). In addition, an aqueous extract of the dried immature fruit of *Poncirus trifoliate* Raf. (PF) was prepared as previously described[17,18] and its prokinetic activities were compared with the Lizhong Tang extract. PF is one of the most popular traditional folk medicines used in Korea and is obtained from Rutaceae fruits. PF has been shown to possess unique, potent prokinetic activities in normal rodents and rodents with GI motility dysfunction (GMD)[13,17].

***Statistical analysis***

The results are expressed as the means ± SE. Statistical analysis was performed using Student's t test or analysis of variance followed by Tukey's multiple comparison test, as appropriate. Statistical significance was accepted for *P* values < 0.05.

**RESULTS**

***Identification of standard compounds in the Lizhong Tang extract***

The following components of the Lizhong Tang extract were detected by HPLC using commercial standards (retention time): liquiritin (25.1 min); ononin (35.9 min); isoliquiritin (36.2 min); ginsenoside Rg1 (36.7 min); glycyrrhizin (58.2 min); 6-gingerol (63.0 min); and atractylenolide III (65.8 min) (Figure 1).

***Effects of the Lizhong Tang extract on ITR in normal mice***

After 30 min, the mean ITR (%) for Evans blue in normal mice was 54.4 ± 1.9% (Figure 2). PF (1g/kg), which has been shown to have prokinetic activity in the GI tract[17,18], significantly accelerated the ITR [79.4 ± 2.3% (*P* < 0.01)], similar to the Lizhong Tang extract, which dose-dependently increased the ITR (%) [ITR values at 0.01, 0.1 and 1 g/kg were 56.1 ± 2.1%, 65.2 ± 1.8% (*P* < 0.01) and 83.8 ± 1.9% (*P* < 0.01), respectively; Figure 2]. Loperamidedecreased the ITR (%), which is consistent with previous reports[19], and the Lizhong Tang extract inhibitedthis loperamide-induced decrease in ITR [ITR value for loperamide was 56.1 ± 2.1%; and ITR value for loperamide with the Lizhong Tang extract was 65.2 ± 1.8% (*P* < 0.01); Figure 2].

***Effects of the Lizhong Tang extract on ITR in mice with GMD***

We used the AA and STZ-induced diabetic mouse models of experimental GMD to examine the effect of the Lizhong Tang extract on GI motility. As mentioned above, the AA mouse model showed a significant retardation of ITR (%) [23.2 ± 1.5% (*P* < 0.01 *vs* normal); Figure 3]. However, a significant inhibition of this retardation was observed when the mice were intragastrically administered 0.01, 0.1, or 1 g/kg of the Lizhong Tang extract [25.3 ± 2.4%, 34.5 ± 2.1% (*P <* 0.01) and 51.8 ± 5.7% (*P <* 0.01), respectively; Figure 3]. No abnormal clinical signs or changes were observed in the AA mice after administration of the Lizhong Tang extract. In addition, loperamide decreased the ITR (%) in the AA mice [13.5 ± 2.4% (*P <* 0.01)], and the Lizhong Tang extractincreased this value [26.7 ± 2.1% (*P* < 0.01); Figure 3]. Furthermore, the STZ-induced diabetic mice also showed a significant ITR (%) retardation (44.1 ± 3.5%; Figure 4), which was also significantly inhibited by treatment with the Lizhong Tang extract at 0.01, 0.1 or 1 g/kg [53.8 ± 1.5% (*P <* 0.01), 57.7 ± 1.4% (*P <* 0.01) and 71.5 ± 3.0% (*P <* 0.01), respectively; Figure 4]. No abnormal clinical signs or changes were observed in the STZ-induced diabetic mice after the administration of the Lizhong Tang extract. In addition, loperamide decreased the ITR in the STZ-induced diabetic mice [20.6 ± 1.8% (*P <* 0.01)], and the Lizhong Tang extract increased this value [40.6 ± 2.2% (*P* < 0.01); Figure 4]. These results indicate that the Lizhong Tang extract increased the ITR in mice with GMD.

***Effect of the Lizhong Tang extract on accelerating GE***

In normal mice, the groups treated with the Lizhong Tang extract (0.01, 0.1 and 1 g/kg) showed significantly enhanced GE (%) values compared to that of the normal group [the GE values with 0.01, 0.1 and 1 g/kg of the Lizhong Tang extract were 61.7 ± 1.6%, 66.8 ± 2.1% (*P* < 0.05) and 72.5 ± 1.7% (*P* < 0.01), respectively; Figure 5]. Its effects were dose-dependent in the dosage range from 0.01 g/kg to 1 g/kg, and 1 g/kg of the Lizhong Tang extract displayed effects similar to those of 5 mg/kg mosapride [74.4 ± 3.3% (*P <* 0.01)] and 5 mg/kg domperidone [72.9 ± 1.9% (*P <* 0.01)] (Figure 5). Next, we examined loperamide-induced and cisplatin-induced models of GE delay to determine whether the Lizhong Tang extract could increase GE in abnormally depressed GE models. In the loperamide-induced model of GE delay, the GE value was lower than normal [40.9 ± 1.6% (*P <* 0.01); Figure 6], and this decrease was recoveredby treatment with the Lizhong Tang extract at doses from 0.01 to 1 g/kg [the GE values for the Lizhong Tang extract at 0.01, 0.1 and 1 g/kg were 41.8 ± 2.2%, 47.8 ± 1.2% (*P* < 0.01) and 59.4 ± 1.5% (*P* < 0.01), respectively; Figure 6]. The maximal effect was obtained at 1 g/kg, and at this dose, the effect of the Lizhong Tang extract was comparable to that of 5 mg/kg mosapride [61.4 ± 2.3% (*P <* 0.01)] or 5 mg/kg domperidone [61.5 ± 1.7% (*P <* 0.01)] (Figure 6). In addition, in the cisplatin-induced model of GE delay, the decreased GE was recovered by treatment with the Lizhong Tang extract (0.01, 0.1 and 1 g/kg) [GE values of the Lizhong Tang extract at 0.01, 0.1 and 1 g/kg were 31.7 ± 1.3%, 43.1 ± 2.1% (*P* < 0.01) and 60.8 ± 1.7% (*P* < 0.01), respectively; Figure 7]. The maximal effect was obtained at 1 g/kg, and at this level, the effect of the Lizhong Tang extract was comparable to that of 5 mg/kg mosapride [65.6 ± 1.4% (*P <* 0.01)] or 5 mg/kg domperidone [63.7 ± 1.2% (*P <* 0.01)] (Figure 7).

**DISCUSSION**

GI motility results from the coordinated contractions of the *tunica muscularis*, which forms the outer wall of the alimentary canal from the distal esophagus to the external anal sphincter[20]. The pathogeneses of primary intestinal motility disorders is probably multifactorial and includes structural and biochemical abnormalities, forms of intestinal pseudo-obstruction, and mucosal inflammation[21].

The GI tract exhibits spontaneous mechanical contractions and electrical pacemaker potentials[7,8], and these pacemaker potentials are the basic determinant of GI smooth muscle activity[7,8]. Recent studies have shown that ICC act as the pacemakers and conductors of electrical slow waves in the GI tract[7,8].

In our previous report, we investigated the effects of Lizhong Tang on ICC in the GI tract[6]. Lizhong Tang induced pacemaker potential depolarizations that were mediated by non-selective cationic channels, intracellular calcium release, and PLC-dependent pathways. Therefore, we suggested that Lizhong Tang might have gastroprokinetic effects on ICC[6]. In addition, many scientists have studied the effects of Lizhong Tang. For example, Lizhong Tang has been shown to have therapeutic effects, such as immunomodulatory, anticancer, antitoxic, and antioxidant effects, and to modulategastric acid secretion[22-24]. Furthermore, in traditional medicine, it is used to treat the spleen or kidney deficiencies associated with the common symptoms of many diseases, such as pasty loose stools, heavy menstrual bleeding, soreness, and weakness of the lower back and knees[2,25,26]. Moreover, Lizhong Tang possesses potent anti-osteoporotic activity and has been suggested for use in the treatment of postmenopausal osteoporosis[3]. In addition, Lizhong Tang protects the gastric mucosa from acute ethanol-induced gastric injury and has been suggested as a treatment for acute gastric injury[1]. However, despite the considerable use of Lizhong Tang, little is known about its regulatory effects on GI motility *in vivo*.

In this study, the Lizhong Tang extract significantly and dose-dependently accelerated ITR (Figure 2). In experimental GMD (AA mouse and STZ-induced diabetic mouse) models, the Lizhong Tang extract significantly inhibited GMD-induced retardation (Figures 3 and 4), and Lizhong Tang extract-treated mice had significantly greater GE values than normal mice; 1 g/kg of the Lizhong Tang extract displayed effects similar to those of mosapride and domperidone(Figure 5). Furthermore, in abnormally depressed (loperamide- and cisplatin-induced) GE models, the Lizhong Tang extract increased GE (Figure 6 and 7).

Prokinetic drugs, which enhance GI motor function by acting on a variety of neurotransmitter receptors, have been used to treat patients with GI motility disorders[27] and are regarded as one of the most efficacious therapeutics for this disorder[28,29]. Cholinergic agonists, the original promotility agents, stimulated muscarinic M2-type receptors on the smooth muscle cells, but their effectiveness in motility disorders is inconsistent[30]. Metoclopramide and domperidone, dopamine antagonists, have been the most widely used as prokinetic agents[31], but their long-term use has been complicated by a trend toward tolerance and a significant incidence of central nervous system (CNS) side effects[32]. Cisapride was shown to promote esophageal peristalsis, augment lower esophageal sphincter pressure, and accelerate gastric emptying[33]. However, the use of this drug is now restricted due to serious cardiac arrhythmias related to a prolonged QT interval[34]. Mosapride, a selective 5-HT4 agonist, is available as a prokinetic agent in a number of Asian countries, but the efficacy data are contradictory[35]. Itopride is a dopamine D2 antagonist with prokinetic effects that is devoid of CNS or cardiovascular side effects and causes minimal elevations of prolactin levels[36]. In this study, we did not directly compare the GE and intestine motility rates with these prokinetics agents. However, in a previous study, we showed that Lizhong Tang depolarized the pacemaker potentials through G-protein-, PLC- and Ca2+-dependent pathways. Moreover, the nonselective cationic cation channel was involved in these effects[6]. Therefore, we believe that Lizhong Tang might mimic the major excitatory neurotransmitters of the GI tract and act as a gastroprokinetic agent. Additionally, herbal products may be an attractive alternative based on the perception of their ‘natural’ approach and their low risk of side effects[37]. Therefore, we believe that Lizhong Tang may be a good gastroprokinetic agent, and in the future, we should compare the experimental results with those of known prokinetics and analyze the side effects.

In summary, in normal ICR mice, both the ITR and GE values were significantly and dose-dependently increased by treatment with the Lizhong Tang extract. Furthermore, the ITRs of GMD mice were significantly reduced compared with those of the normal mice, and these reductions were significantly and dose-dependently reversed by treatment with the Lizhong Tang extract. In addition, in the loperamide-induced and cisplatin-induced model of GE delay, the Lizhong Tang extract prevented the observed GE delays. Taken together, our results suggest that Lizhong Tang is a good candidate for the development of a gastroprokinetic agent.

**COMMENTS**

***Background***

Lizhong Tang [composed of Radix Ginseng (*Panax ginseng* C.A. Meyer), Rhizoma Zingiberis (*Zingiber officinale* Roscoe), Rhizoma Atractylodis Macrocephalae (*Atractylodes macrocephala* Koidz.) and Radix Glycyrrhizae (*Glycyrrhiza uralensis* Fisch.)]is a traditional Chinese medicinal formula that has been widely used in China, Japan and Korea for many years to ameliorate the symptoms of gastrointestinal(GI) disorders. However, despite the considerable use of Lizhong Tang in traditional medicineto treat GIdysfunction, little was known of its regulatory effects on GI motility *in vivo*.

***Research frontiers***

Lizhong Tang is a good candidate for development as a gastroprokinetic agent.

***Innovations and breakthroughs***

In normal ICR mice, both the ITRs and GE values were significantly and dose-dependently increased by treatment with Lizhong Tang (0.1-1 g/kg). The ITRs of the GMD mice were significantly reduced compared with those of the normal mice, and the values were significantly and dose-dependently reversed by treatment with Lizhong Tang (0.1-1 g/kg). Moreover, in loperamide-induced and cisplatin-induced models of GE delay, Lizhong Tang prevented the observed GE delays.

***Applications***

Lizhong Tang may be a new target or a novel candidate prokinetic agent for the pharmacological treatment of GI motility disorders.

***Terminology***

GI motility: movements of the digestive system and the transit of contents within the digestive system; ITRs: rate of passage of food (sometimes in the form of a test meal) through the GI tract; Gastric emptying: to empty the stomach contents

***Peer-review***

This study is relevant, interesting, is written in suitable english, and have a correct methodological design. It is important to emphasize the contribution that this study provides for integration between western and eastern medicine, which is fundamental to the advancement of modern science.

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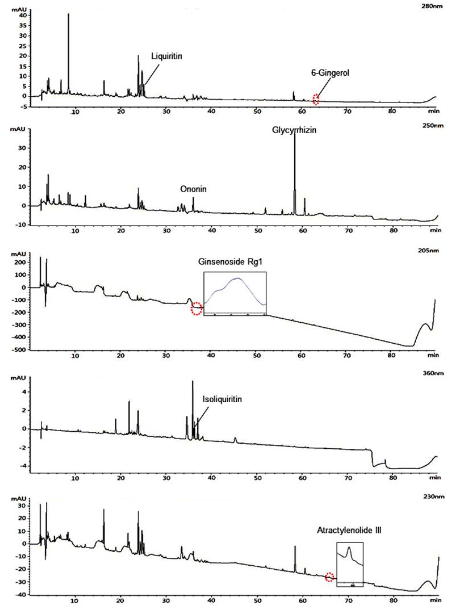
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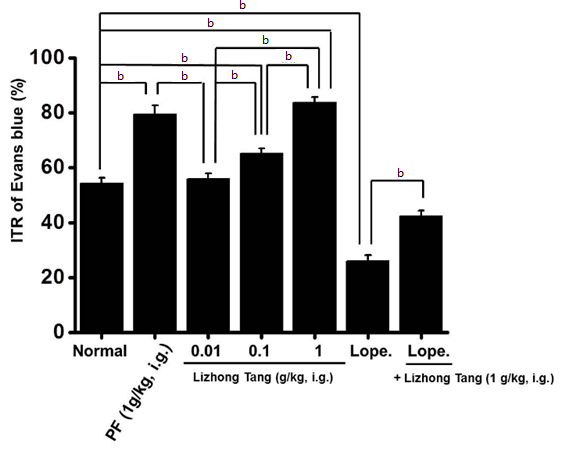
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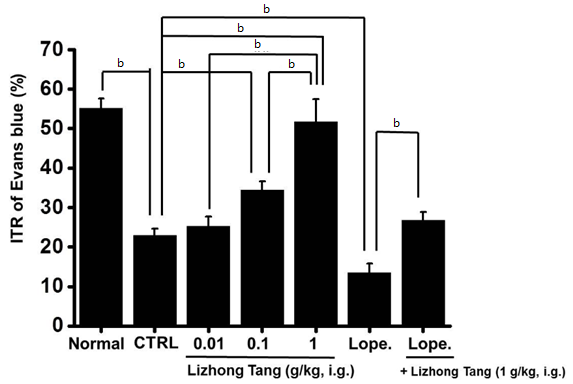
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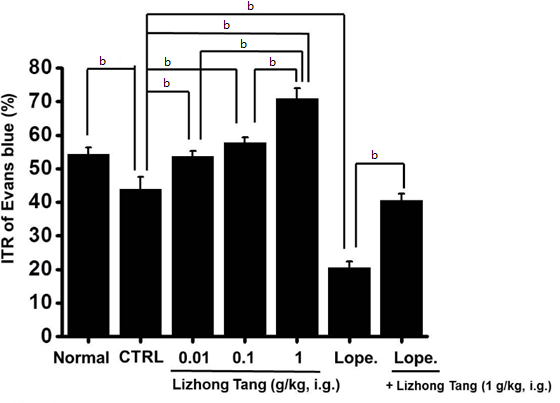
**Figure 1 Chromatograms of liquiritin, 6-gingerol, ononin, glycyrrhizin, ginsenoside Rg1, isoliquiritin, and atractylenolide III in Lizhong Tang extract.**



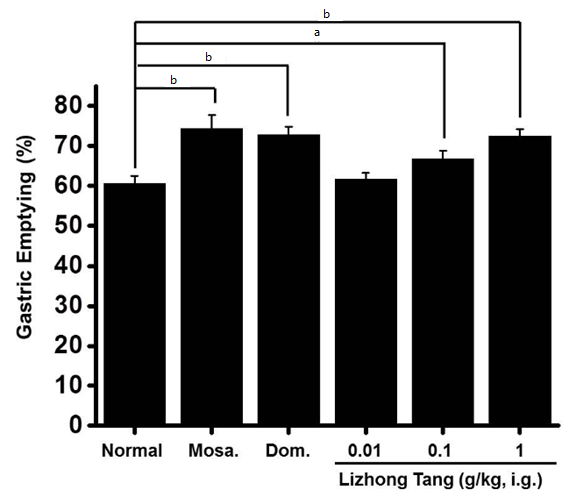
**Figure 2 Effects of the Lizhong Tang extract on the intestinal transit rates (%) in normal mice.** Intestinal transit rates (ITRs) (%) values of normal mice that were pretreated with the Lizhong Tang extract prior to Evans blue administration (*n* = 15 for each bar). The bars represent mean values ± SE. b*P* < 0.01; Significantly different from the normal controls. PF: *Poncirus trifoliate* Raf; Lope: Loperamide.



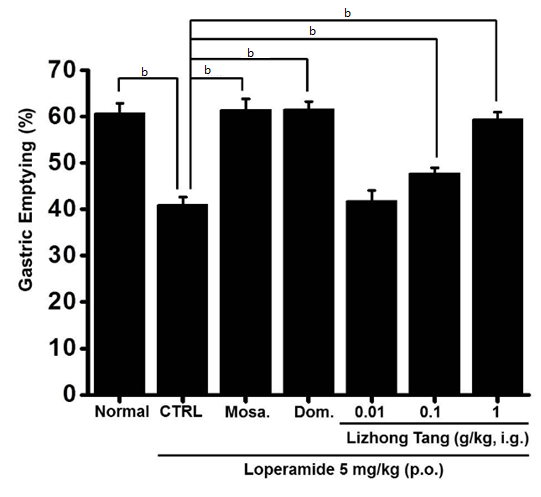
**Figure 3 Effects of the Lizhong Tang extract on the intestinal transit rates (%) in AA mice.** Intestinal transit rates (ITRs) (%) values of AA mice induced 30 min before the i.g. administration of Evans blue (*n* = 12 per bar). The bars represent mean values ± SE. b*P* < 0.01; significantly different from the AA controls. CTRL: Control; Lope: Loperamide.



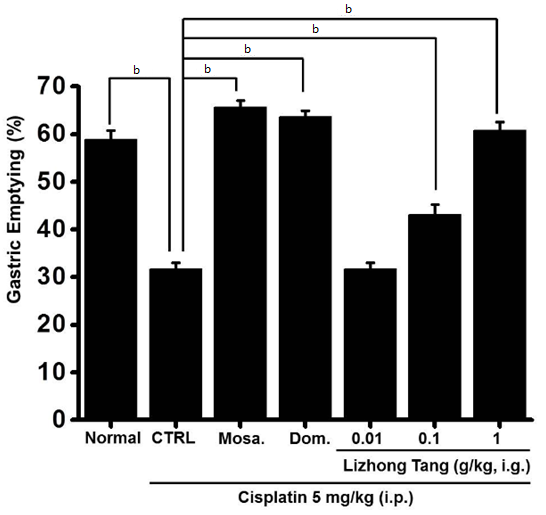
**Figure 4 Effects of the Lizhong Tang extract on the Intestinal transit rates** **(%) values in STZ-induced diabetic mice.** Intestinal transit rates (ITRs) (%) values of the STZ mice induced 2 months before the i.g. administration of Evans blue (*n* = 12 per bar). The bars represent mean values ± SE. b*P* < 0.01; significantly different from the STZ-induced diabetic controls. CTRL: Control. Lope: Loperamide.



**Figure 5 Effect of the Lizhong Tang extract on accelerating gastric emptying.** After a 24 h fast, the animals (*n* = 7/each group) were orally administered the indicated dosages of the Lizhong Tang extract, 5 mg/kg of a 5-HT4 receptor agonist (mosapride), 5 mg/kg of a dopamine receptor antagonist (domperidone), or distilled water (DW; controls). The GE percentages were calculated as described in the Materials and Methods. The bars represent means ± SE. a*P* < 0.05; b*P* < 0.01; significantly different from the normal controls. Mosa.: Mosapride. Dom.: Domperidone.



**Figure 6 Lizhong Tang extract ameliorates loperamide-induced delays in gastric emptying (GE).** After a 24 h fast, the animals (*n* = 6/each group) were orally administered the indicated dosages of the Lizhong Tang extract, 5 mg/kg of a 5-HT4 receptor agonist (mosapride), 5 mg/kg of a dopamine receptor antagonist (domperidone), or distilled water (DW; control). The loperamide-induced gastric emptying (GE) delay was inhibitedby the Lizhong Tang extract. The GE percentages were calculated as described in the Materials and Methods. The bars represent means ± SE. b*P* < 0.01; significantly different from the normal controls. Mosa: Mosapride; Dom: Domperidone; p.o.: per os.



**Figure 7 Lizhong Tang extract ameliorated cisplatin-induced delays in gastric emptying.** After a 24 h fast, the animals (*n* = 6/each group) were orally administered the indicated dosages of the Lizhong Tang extract, 5 mg/kg of a 5-HT4 receptor agonist (mosapride), 5 mg/kg of a dopamine receptor antagonist (domperidone), or distilled water (DW; control). The Lizhong Tang pretreatment prevented thecisplatin-induced GE delay. The GE percentages were calculated as described in the Materials and Methods. The bars represent mean values ± SE. b*P* < 0.01; significantly different from the normal controls. Mosa: Mosapride; Dom: Domperidone; i.p: intraperitoneally.