**Name of Journal: *World Journal of Gastroenterology***

**Manuscript NO: 32465**

**Manuscript Type: Evidence-Based Medicine**

**Anti-apoptotic effect of banhasasim-tang on chronic acid reflux esophagitis**

Shin MR *et al*. Banhasasim-tang and chronic acid reflux esophagitis

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**Author contributions:** Shin MR performed the majority of experiments and wrote the paper; Seo BI analyzed the data; An HJ and Roh SS designed and coordinated the research; All authors read and approved the final version for publication.

**supported by** National Research Foundation of Korea; and Korean government (MSIP), No. 2017R1A2B2006858.

**Institutional animal care and use committee statement:** All procedures involving animals were reviewed and approved by the Institutional Animal Care and Use Committee of the Daegu Haany University (No. DHU2016-81).

**Conflict-of-interest statement:** The authors declare no competing financial interests related to this study or its publication.

**Data sharing statement :** No additional data are available.

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**Manuscript source:** Unsolicited manuscript

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**Received:** January 9, 2017

**Peer-review started:** January 10, 2017

**First decision:** February 23, 2017

**Revised:** March 8, 2017

**Accepted:** April 12, 2017

**Article in press:**

**Published online:**

**Abstract**

***AIM***

To evaluate the anti-apoptotic effect of banhasasim-tang (BHSST) on chronic acid reflux esophagitis (CARE) using a rat model.

***METHODS***

A surgically-induced CARE model was established in Sprague-Dawley rats. The modeled rats were divided into a treatment group or untreated group, and given BHSST (1 g/kg body weight per day) or water, respectively, for 15 consecutive days (*n* = 7 each group). Changes in expression of proteins related to nicotinamide adenine dinucleotide phosphate (NADPH) oxidase and apoptosis were assessed by western blotting. Changes in esophageal pathology were analyzed by gross and histological examinations.

***RESULTS***

The CARE exposure modeled rats showed increased levels of the NADPH oxidase subunit, NOX4 and p47phox in the esophagus. The BHSST treatment completely resolved these CARE-related increases. The CARE rats also showed markers of cytokine stress, including elevated levels of TNF-and reactive oxygen species as well as of the consequent increase in JNK activation, and subsequent decrease in pro-survival gene expressions, such as *Bcl-2*. BHSST treatment resolved the CARE-related changes. BHSST also exerted an anti-apoptotic effect, as evidenced by altered expression of the apoptosis-related genes for bax, cytochrome c, and caspase 3. Finally, the BHSST treatment markedly ameliorated the CARE-related esophageal mucosal ulcerations.

***CONCLUSION***

In the rat model of CARE, BHSST can suppress development of esophageal mucosal ulceration *via* regulation of reactive oxygen species-dependent apoptosis.

**Key words:** Banhasasim-tang; Chronic acid reflux esophagitis; Esophageal ulcer; Reactive oxygen species; Apoptosis

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**Core tip**: Banhasasim-tang (BHSST) has been used widely as an herbal prescription in East Asia for its therapeutic effects on symptoms associated with gastroesophageal reflux disease. In this study, BHSST is shown to play a protective role against chronic acid reflux esophagitis -induced esophageal mucosal ulcer in a rat model, and that this effect involves regulation of reactive oxygen species-dependent apoptosis.

Shin MR, An HJ, Seo BI, Roh SS. Anti-apoptotic effect of banhasasim-tang on chronic acid reflux esophagitis. *World J Gastroenterol* 2017; In press

**INTRODUCTION**

Gastroesophageal reflux disease (GERD) is one of the most frequently encountered gastrointestinal disorders worldwide, occurring in children as well as adults and showing a trend towards increasing risk in the past few decades[1]. Various etiologies have been demonstrated as underlying the disease pathogenesis, ranging from diet, the increased social life stress of modern-day life, and diverse array of medical conditions[2]. As such, the symptoms of GERD are equally diverse, ranging from heartburn and regurgitation to severe erosive esophagitis and its associated complications. Patients with GERD can also suffer sleep disturbances, chest pains or respiratory symptoms[3].

Economic analyses have indicated that the increased, incidence of GERD is accompanied by increased direct and indirect costs related to its diagnosis, treatment and supervision, as well as of costs related to its and its complications. The currently prescribed medications for GERD are pharmaceutical drugs-primarily H2 receptor antagonists (H2Ras) and proton pump inhibitors (PPIs)[4]. However, these drugs are associated with multiple adverse effects, such as headaches, diarrhea, nausea (H2RAs and PPIs) and constipation (PPIs)[5,6]. Herbal medicines have been proposed as a useful alternative, and the field of GERD research has begun to search for an herbal therapeutic useful for the treatment and/or prevention of GERD and to define the molecular mechanisms underlying any such beneficial activities.

Recently, [Wei](http://www.sciencedirect.com/science/article/pii/S0952327816300242) *et al*[7] uncovered a close relationship between chronic acid reflux esophagitis (CARE) and an altered profile of markers of oxidative stress (OS). An exaggerated or unregulated prolonged inflammatory process, in response to various exogenous and endogenous stimuli, can induce tissue damage and has been implicated in the development and progression of many chronic diseases, include CARE[8]. Continuous reflux of gastric contents causes ulceration and destruction of the normal squamous epithelium of the esophagus. Over time, the esophagus adapts to the continuous reflux stimuli with metaplastic convertsion to columnar epithelium[9,10]. These series of processes led to Barrett's esophagus, a common premalignant lesion of esophageal adenocarcinoma. Accordingly, management of Barrett's esophagus (*via* the suppression of the gastroesophageal reflux) is recommended, at the earliest stage possible, to prevent the development of gastroesophageal carcinoma in an early stage[11].

Banhasasim-tang (BHSST) is a classic herbal formulation in Traditional Chinese Medicine (TCM). Its first recorded description is in the Shang-Han Lun (transl. *Treatise on Cold Damage and Miscellaneous Diseases*) written by the Chinese physician Zhang Zhong-jing (A.D. 150 - 219). He suggests the treatment of disease and a proper mixture of medicines in Shang-Han Lun after he synthesized clinical experiences and therapeutic principles based on the existing medical books[12,13]. It continues to be in use throughout Asia, including in Korea, for treating the symptoms associated with GERD[14-16].

However, the molecular mechanisms underlying its observed protective effects against GERD remain unknown. The research study described herein was designed to begin to investigate the molecular mechanisms of BHSST protection against esophageal mucosal ulcer by using a rat model of CARE.

**METERIALS AND METHODS**

***Materials***

Protease inhibitor mixture cocktail, 4,6-dihydroxy-2-mercaptopyrimidine (2-thiobarbituric acid, TBA), and ethylenediaminetetraacetic acid (EDTA) were purchased from Wako Pure Chemical Industries, Ltd. (Osaka, Japan). 2′,7′-Dichlorofluorescein diacetate (DCF-DA) was obtained from Molecular Probes (Eugene, OR, United States). Phenylmethylsulfonyl fluoride (PMSF) was purchased from Sigma-Aldrich Chemical Co (St. Louis, MO, United States). Pierce bicinchoninic acid protein assay kit was obtained from Thermo Fisher Scientific (Waltham, MA, United States). Enhanced chemiluminescence (ECL) western blotting detection reagents and nitrocellulose membranes were supplied by GE Healthcare (Chicago, IL, United States). Rabbit polyclonal antibodies against bax (1:1000; SC-7480), p47phox (1:1000; SC-14015) and Cytochrome c (1:1000; SC-13156); goat polyclonal antibodies against tumor necrosis factor- (TNF-)(1:1000, SC-1351); mouse monoclonal antibodies against phosphor-c-Jun NH2-terminal kinase (p-JNK) (1:1000; SC-6254), histone (1:1000; SC-8030) and -actin (1:1000; SC-4778) were purchased from Santa Cruz Biotechnology, Inc. (Dallas, TX, United States). Mouse monoclonal antibody against activator protein-1 (AP-1) subunit c-Jun (1:1000; #2315) was obtained from Cell Signaling Technology, Inc. (Danvers, MA, United States). Mouse monoclonal anti-caspase-3 (1:1000; 3004-100) was purchased from BioVision Inc. (Mountain View, CA, United States). Rabbit polyclonal anti-reduced nicotinamide adenine dinucleotide phosphate oxidase 4 (NOX4) was purchased from LifeSpan BioSciences (Seattle, WA, United States). Rabbit anti-goat (1:3000; SC-2774), goat anti-rabbit (1:3000; SC-2004) and goat anti-mouse (1:3000; SC-2005) immunoglobulin G (IgG) horseradish peroxidase (HRP)-conjugated secondary antibodies were acquired from Santa Cruz Biotechnology, Inc. All other chemicals and reagents used were of the analytical grade purchased from Sigma-Aldrich Co.

***BHSST formulation***

Light brown granules of BHSST were purchased from Hankook Shinyak Corp (Nonsan-si, Chungcheongnam-do, South Korea) and had been produced according to the Korean Good Manufacturing Practice under permit granted by regulation of the Korean Food & Drug Administration (Seoul, South Korea). For experimentaion, 1.5 g BHSST was dissolved in distilled water. The specific composition of the BHSST formulation is presented in Table 1.

***Animal experiments***

All procedures involving animals were carried out in full accordance to the “Guidelines for Animal Experimentation” with preapproveal given by the Ethics Committee of the Daegu Haany University. Male Sprague-Dawley rats (Nara Biotec Co, Pyeongtaek, South Korea) weighing 160–170 g were housed in cages with constant temperature of 24 ℃± 2 ℃, relative humidity of 60% automated 12:12 h light/dark cycle (light on at 7:00 AM), and free access to food and water.

CARE was surgically induced as described by Omura *et al*[17]. Briefly, a midline laparotomy was performed to expose the stomach and transitional region (*i.e.*, limiting ridge) between the fundus and the glandular portion of the stomach was ligated with 2–0 silk thread in order to restrict the compliance of the stomach, which led to the reflux of gastric contents into the esophagus. Additionally, a latex ring (2 mm in thickness; ID, 4 mm, made from 18-Fr Nelaton catheter) was placed around the pyloric sphincter so as to restrict the emptying of gastric contents. Rats were then injected with gentamicin sulfate (antibiotic, subcutaneous injection) and dexamethasone (anti-inflammatory agent, subcutaneous injection) to prevent infection.

After surgery, the rats fasted for a further 48 h but water was provided 24 h after surgery. All animals had an operational adjustment for 7 d post surgery. Body weight was recorded every day over a course of 22 d from the surgery day (during the “operational adjustment period” of the first 7 d post-surgery and then during the “BHSST treatment period” of the next 15 d. Food intake was recorded every day during the 15-d (BHSST treatment period). For the 15-d treatment course, the CARE modeled rats were divided into two groups (*n* = 7 each) and given an oral administration (*via* stomach tube) of either water (CARE group) or BHSST at 1 g/kg body weight (BHSST-treated CARE group). Blood samples were collected by vena cava puncture from anesthetized rats. At day 22 post-surgery, all rats were sacrificed and the entire esophagus was removed and examined rapidly for gross mucosal injury and frozen in liquid nitrogen. Both the esophageal tissues and serum samples were kept at -80 ℃ until analysis.

***Serum reactive oxygen species and TBA-reactive substance measurement***

Serum reactive oxygen species (ROS) level was measured as described by Ali *et al*[18]. Briefly, 25 mmol/L DCF-DA was added to the serum sample and allowed to incubate for 30 min. The DCF-DA-induced change in fluorescence value was measured at an excitation wavelength of 486 nm and emission wavelength of 530 nm. TBA-reactive substance (TBARS) level was estimated according to the method of Mihara and Uchiyama[19].

***Preparation of cytosolic and nuclear fractions***

Protein extraction was performed according to the method described by Komatsu[20], with minor modifications. Briefly, esophageal tissue was homogenized in ice-cold lysis buffer A (250 mL) containing 10 mmol/L HEPES (pH 7.8), 10 mmol/L KCl, 2 mmol/L MgCl2, 1 mmol/L DTT, 0.1 mmol/L EDTA, 0.1 mmol/L PMSF, and 1250 μL protease inhibitor mixture cocktail. The homogenate was then incubated at 4 oC for 20 min, after which 10% NP-40 was added and mixed well. After centrifugation at 13400 ×g for 2 min at 4 ℃ (5415R centrifuge; Eppendorf, Hamburg, Germany), the supernatant liquid (cytosolic fraction) was sremoved to a new tube. The leftover pellet was then washed twice with buffer A, centrifuged and the supernatant discarded. The leftover pellet was suspended in lysis buffer C (20 mL) containing 50 mmol/L HEPES (pH 7.8), 50 mmol/L KCl, 300 mmol/L NaCl, 1 mmol/L DTT, 0.1 mmol/L EDTA, 0.1 mmol/L PMSF, 1% (v/v) glycerol, and 100 μL protease inhibitor mixture cocktail and incubated at 4 ℃ for 30 min. After centrifugation at 13400 ×g for 10 min at 4 ℃, the nuclear fraction was obtained. Both cytosolic and nuclear fractions were kept at -80 ℃ until use in analysis.

***Immunoblotting analyses***

For estimation of c-Jun and histone, 13.6 μg of protein from each nuclear fraction was resolved by electrophoresis through a 10% sodium dodecylsulfate polyacrylamide gel (SDS-PAGE). The separated proteins were transferred to a nitrocellulose membrane, blocked by incubating with 5% (w/v) skim milk solution for 1 h, and then incubated with primary antibodies (c-Jun and histone) for overnight at 4 ℃. After the blots were washed, they were incubated with anti-rabbit or anti-mouse IgG HRP-conjugated secondary antibody for 1 h at room temperature. In addition, 10-16 μg protein of each cytosolic fraction was eletrophoresed 10%-14% SDS-PAGE for immunodetection of TNF- p-JNK, NOX4, p47phox, bax, cytochrome c, caspase 3, and -actin. Each antigen-antibody complex was first processed with the ECL regents and then detected by the Sensi-Q 2000 Chemidoc instrument (Lugen Sci Co., Ltd., Gyeonggi-do, South Korea). Band densities were measured using ATTO Densitograph Software (ATTO Corporation, Tokyo, Japan) and quantified as the ratio to histone or -actin. Protein levels of the groups are expressed as relative to those of normal rat (set to 1.0).

***Statistical analysis***

The data are expressed as mean ± SE. Statistical analysis was performed using SPSS version 22.0 software (IBM SPSS Inc, Armonk, NY, United States). *P* values less than 0.05 were considered to indicate statistical significance.

**RESULTS**

***CARE-releated change in body weight and food intakeand effect of BHSST***

CARE modeled-rats had significantly lower body weight and food intake during the BHSST treatment period than the of normal rats **(**[**Table 2**](http://jn.nutrition.org/content/144/8/1150.long#T1)**).** However, the BHSST-treated CARE modeled-rats had significantly higher food intake than the untreated CARE. The BHSST treatment appeared to have no significant effect on the body weight of the CARE modeled-rats.

***BHSST protects against CARE-related effects on gross and histologic pathology***

Normal rats had no detectable esophageal mucosa lesions, whereas esophageal ulcers were readily apparent in the middle or distal esophagus of the untreated CARE modeled-rats (Figure 1). As compared to the normal rats, the untreated CARE modeled-rats exhibited basal layer thickening and inﬂammatory cell inﬁltration; the BHSST-treated CARE modeled-rats showed remarkably less extensive damage (Figure 2).

***BHSST protects against CARE-related effects on serum ROS and TBARS***

As shown in Table 3, the levels of the OS-related biomarkers, ROS and TBARS, in untreated CARE modeled-rats were markedly higher than those of normal rats (*P* < 0.01). The BHSST-treated CARE modeled-rats showed a decrease, these CARE-induced elevations, to levels even lower than those in normal rats. In addition, the BHSST-treated led to significant decreases in serum ROS and TBARS, but only the ROS decrease was statistically significant (*P* < 0.05).

***BHSST protects against CARE-induced up-regulation of esophageal p47phox and NOX4 protein expression***

The expressions of both p47phox and NOX4proteins –the markers of NADPH oxidase activity in the esophageal tissues– were augmented in the CARE modeled-rats *(vs* normal rats*,* *P* < 0.01).The BHSST-treated CARE modeled-rats, however, had significantly down-regulated NADPH oxidase (*vs* untreated CARE modeled- rats, *P* < 0.01)(Figure 3).

***BHSST protects against CARE-induced up-regulation of esophageal TNF-* *p-JNK, and c-Jun protein expression***

The untreated CARE modeled-rats had increased protein expression for TNF-and p-JNK in the cytosolic fraction, and for c-Jun in the nuclear fraction. The up-regulation was reversed by the BHSST treatment (Figure 4).

***BHSST protects against CARE-induced up-regulation of apoptosis-related factors in the esophagus***

As shown in Figure 5, the apoptosis-related factors bax, cytochrome c and caspase 3 were examined in the cytosolic fraction. All three of these proteins were significantly higher in the untreated CARE modeled (*vs* normal rats, *P* < 0.05, < 0.001, and < 0.01, respectively). These elevated expressions were substantially enhanced by the BHSST treatment, with bax and caspase 3 expression being decreased to a level near that seen in the normal (*P* < 0.05 and < 0.01, respectively).

**DISCUSSION**

Several herbals from TCM have been shown to exert the therapeutic effect through the inhibition of apoptosis, particularly in the treatment of GERD[21]. However, the role of BHSST still lacks evidential data of its underlying molecular mechanism against CARE. The present study provided evidence that supplementation of BHSST significantly ameliorated CARE-induced esophageal ulcer.

Traditional herbal formulas have been generally used throughout East Asia for prevention and treatment of various inflammatory disorders[22]. BHSST is one such herbal medicine, and its formulation consists of seven herbs (in 5:3:3:3:3:3:1 propotions) : Pinelliae rhizoma, Scutellariae radix, Zingiberis rhizoma siccus, Ginseng radix alba, Glycyrrhizae radix, Zizyphi fructus, and Coptidis rhizoma. Studies of the various constituents have elucidated the antiemetic effect of Pinelliae rhizoma[23], the anti-inflammatory and antitumor effects of Scutellariae radix[24] and Ginseng radix alba[25], the antioxidant effect of Glycyrrhizae radix[26], Zingiberis rhizoma siccus[27], Coptidis Rhizoma[28] and Ziziphus jujuba[29] was widely used to ameoliorate the symptoms of gastrointestinal disorders in Eastern Asia. The 13 major bioactive components of BHSST are homogentisic acid, 3,4-dihydroxybenzaldehyde (from Pinelliae rhizoma), spinosin (Zizyphi fructus), liquiritin , liguiritigenin and glycyrrhizin (Glycyrrhizae radix), baicalin, baicalein, wogonoside and wogonin (Scutellariae radix), ginsenoside Rg1 and ginsenoside Rb1 (Ginseng radix alba), and 6-gingerol (Zingiberis rhizoma siccus)[30]. Especially, baicalin is composed of the most content. BHSST formulations including these bioactive components have been shown to exert gastroprotective effects and to do so through inhibition of inflammatory proteins[31,32]. Based on the previous studies of BHSST, we predicted that a formulation containing these bioactive herbs would allow for improvement of esophageal ulcer induced by CARE in a rat model. For this reason, the present study was conducted using the same CARE model system frequently used in the previous experimental studies[17]. Firstly, body weight gain during the experimental period and food intake during the BHSST treatment periods were confirmed. As shown in Table 2, although the normal rats and CARE modeled-rats started out at similar body weights CARE modeled-rats experienced weight loss until recovery from the surgery (4 d), during which time the normal rats experienced a gradual increase and this was due to low food intake. However, the significant increase in food intake upon BHSST administration led to some weight gain (but without statistical significance). Thus, the CARE-inducing surgery appears to have brought aboute the changes in food intake and body weight[33].

Next, gross morphological changes such as mucosal thickening and esophageal ulcer, both of which are associated with the metaplastic process of mucosal epithelial cells were observed in the CARE modeled-rats; and, this is in contrast to the features of hemorrhage, hyperemia and multiple erosions in acute reflux esophagitis[34]. The CARE-related tissue injuries were located in the lower part of esophagus, in particular. However, esophageal ulceration was completely absent in the normal rats. The normal esophagus exhibited a thin epithelial layer with squamous cells and few inﬂammatory cells in the submucosal layer, while the CARE esophagus exhibited the characteristic basal layer thickening and inﬂammatory cell inﬁltration[35]. Importantly, the CARE modeled-rats that were treated with BHSST showed remarkably less damage than their untreated CARE control counterparts.

Apoptosis, or programmed cell death, serves to remove superfluous, damaged, infected, or transformed cells, and is a key cellular process for maintenance of tissue homeostasis. Two main pathways lead to apoptosis: the extrinsic (or death receptor) pathway, and the intrinsic (or mitochondrial) pathway[36,37]. The extrinsic pathway requires activation of membrane “death” receptors, such as those for TNF. The intrinsic pathway, which functions independently of the death receptors, involves the mitochondria and is mediated through stimulation by stress molecules (*i.e.*, ROS)[38]. An important cellular source of ROS is the NADPH oxidase family. Previous studies have reported that pharmacological suppression of NADPH oxidase activity prevents tissue apoptosis[39]. These findings suggest that NADPH oxidases may contribute to apoptosis of esophageal tissue, a feature of GERD in humans and CARE in rats[40]. In the present study, CARE modeled rats showed up-regulation of both NADPH oxidase and TNF- in esophageal ulcer, these CARE-related changes were reversed by BHSST administration.

In the absence of OS, JNK can bind to glutathione S-transferase, resulting in inhibition of JNK activity. Under OS conditions which are characterized by ROS overproduction, glutathione S-transferase is dissociated from JNK, resulting in its activation[41]. Increased levels of ROS, which exceed the capacity of cellular antioxidant defense systems, lead to OS and have implicated in the pathology of several chronic diseases[42]. Numerous studies have shown that the JNK/AP-1 pathway can stimulate expression of pro-apoptotic genes, such as TNF. JNK is also known to decrease the expression of pro-survival genes, as has been shown for Bcl-2. Besides, the pro-apoptotic protein bax may promote cytochrome c release and the subsequent activation of effector caspases. In a healthy cell, Bcl-2 is expressed on the external surface of the mitochondrial membrane, where it is available for binding to Apaf-1, which is thereby kept inactivated. Any alteration in the internal equilibrium of the cell, such as ROS accumulation, then causes mitochondrial release of cytochrome c. In turn, Bcl-2 liberates Apaf-1, that then binds to the released cytochrome c[43]. Furthermore, the release of cytochrome c from mitochondria into the cytoplasm is an important regulatory step in caspase 3 activation.

In the current study, JNK activation was moderately enhanced in CARE modeled-rats, as compared to normal rats. BHSST treatment facilitated a substantial down-regulation of the CARE-related enhancement of JNK activation. Moreover, BHSST was found to increase transcription of the gene encoding the anti-apoptotic Bcl-2-like protein, which would serve to inhibit the pro-apoptotic factor bax and to attenuate the transcription of genes encoding for pro-apoptotic proteins, such as bax. Taken together, regulating expression of genes encoding Bcl-2 during CARE could be relevant to explain the beneficial effects of BHSST in treatment of CARE. Nevertheless, the mechanism of action of BHSST remains to be unambiguously defined and further research is required.

In conclusion, data from the present study indicates that the down-regulation of TNF-α or the suppression of NADPH oxidase activity affects the esophageal apoptotic response in CARE. Meanwhile, BHSST treatment was shown to exert beneficial anti-apoptotic effects in CARE, as shown in Figure 6. Accordingly, BHSST may be a promising herbal formula for the prevention or treatment of GERD. However, the relationship between the prevalence of esophageal mucosal ulcerations and other risk or etiological factors, such as gastrointestinal motility, gastric acid and pepsin secretion, or esophageal sensitivity is unknown, and further study is necessary in order to adequately understand this relationship.

**COMMENTS**

***Background***

Gastroesophageal reflux disease (GERD), including reflux esophagitis, is mainly caused by excessive or continuous exposure of the esophageal mucosa to the gastric contents. The existing therapeutic strategy for GERD is primarily acid suppression, and commonly involves the use of pharmacological antacids, H2-receptor antagonists, and proton pump inhibitors (PPIs). Despite their well-known efficacies, recurrence among the treated patients is considerably high. Recent studies have reported that oxidative stress plays an important role in the pathogenesis of reflux esophagitis. Hence, the authors sought to determine whether banhasasim-tang (BHSST) treatment exerts a protective effect under oxidative stress status of chronic acid reflux esophagitis (CARE) using a rat model.

***Research frontiers***

GERD, which causes symptoms of heartburn and acid regurgitation, is a common disease affecting modern life, despite the great achievements that have increased our understanding of the pathophysiology and treatment of the disease. Incidence of GERD has grown worldwide in recent decades, and the quality of life for a patient with GERD. The findings from this research may help health care professionals in dealing with GERD. Consequently, BHSST, one of more safe and effective herbal formulations, may prove useful as a therapeutic to increase the quality of life of GERD patients.

***Innovations and breakthroughs***

This novel study to demonstrated that BHSST plays a protective role against esophageal mucosal ulcer, possibly through modulation of apoptotic proteins *via* suppression of oxidative stress.

***Applications***

This study may provide a future strategy for therapeutic intervention for esophageal mucosal ulcer that is induced in CARE model-rats.

***Terminology***

The esophageal mucosal ulcer is involved in CARE. CARE leads to oxidative stress. But suppression of oxidative stress can affect apoptosis in the esophagus. As the result, BHSST administration exerts a protective effect through the inhibition of reactive oxygen species (ROS).

***Peer-review***

This is a well written and planned study demonstrating the protective effects of BHSST on a rat CARE model system. The protective effects of BHSST seem to arise from regulation of ROS-dependent apoptosis. The results of this study are very interesting and good.

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**P-Reviewer:** Dong H, Guo XZ, Teramoto-Matsubara OT **S-Editor:** Ma YJ **L-Editor:** **E-Editor:**

**Specialty type:** Gastroenterology and hepatology

**Country of origin:** South Korea

**Peer-review report classification**

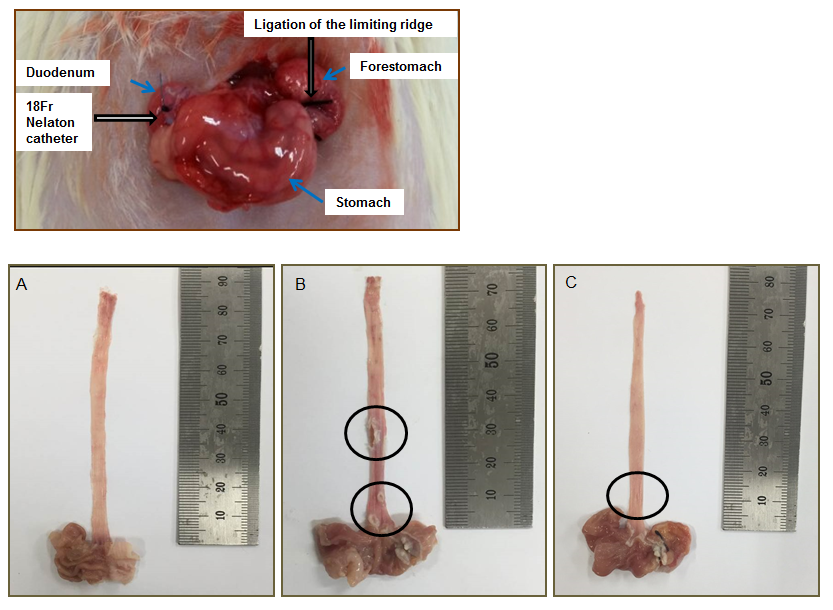
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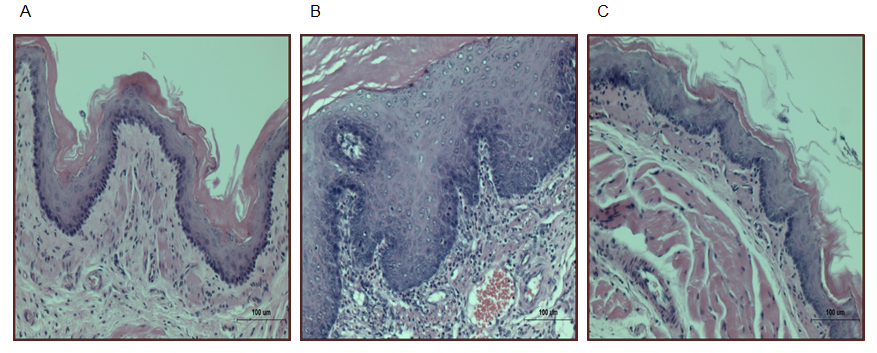
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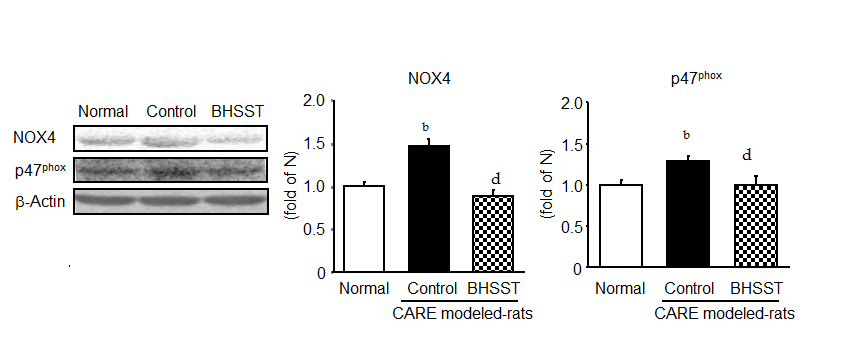
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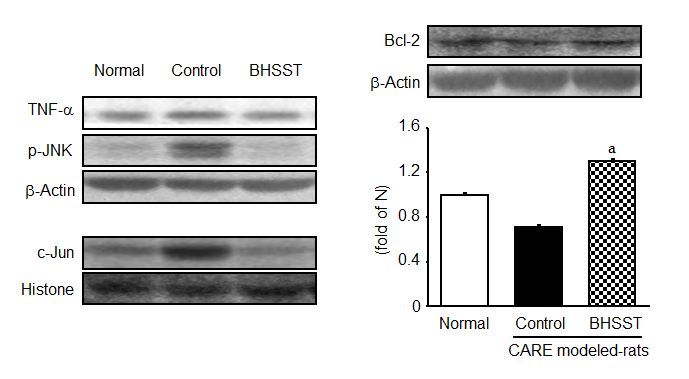
**Figure 1 Surgical induction of chronic acid reflux esophagitis and representative image of the degree of esophageal mucosal ulcer in each group of rats.** A: Normal group; B: Untreated chronic acid reflux esophagitis modeled group; C: BHSST-treated group. BHSST: Banhasasim-tang.



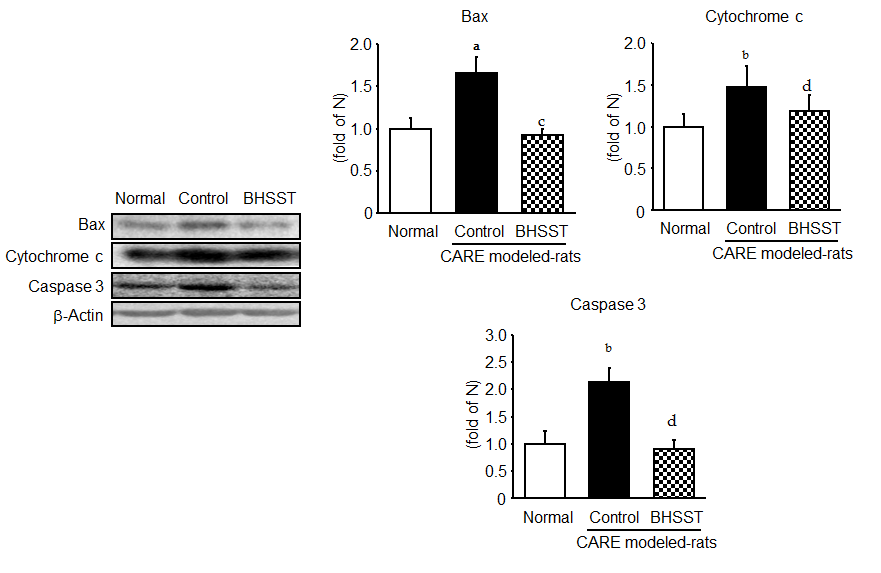
**Figure 2 Chronic acid reflux esophagitis and effects of banhasasim-tang treatment.** A: Normal, normal rats; B: Control, untreated CARE modeled-rats; C: BHSST, BHSST-treated (1 g/kg body weight per day) CARE modeled-rats. Histology of the non-CARE esophagus revealed thin epithelium with few inflammatory cells. Whereas, CARE esophagus showed mucosal thickening with basal cell hyperplasia and marked inflammatory cell infiltration. These histological changes were improved by BHSST treatment. H&E staining; original magnification ×200. CARE: chronic acid reflux esophagitis; BHSST: Banhasasim-tang.



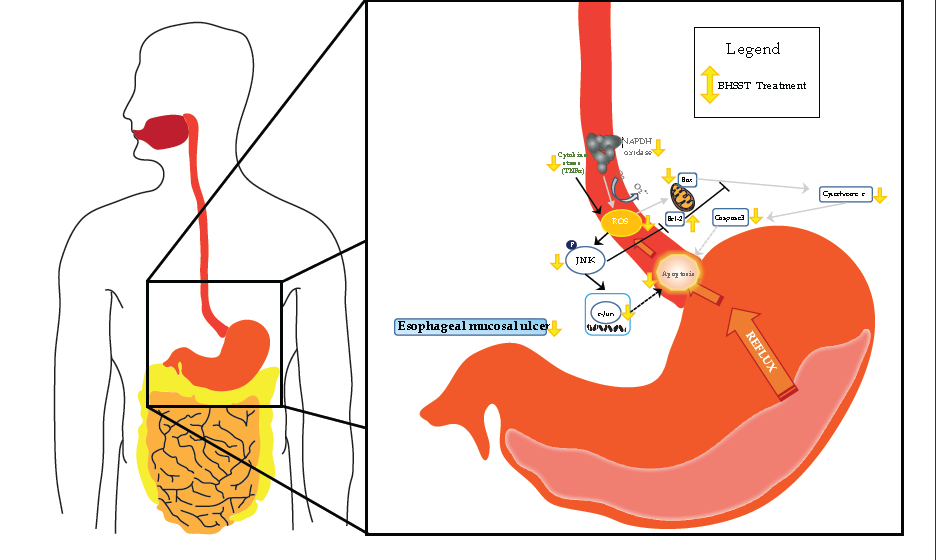
**Figure 3 Expression of NOX4 and p47phox in chronic acid reflux esophagitis rats and effects of banhasasim-tang treatment treatment.** Normal, normal rats; control, untreated CARE modeled-rats; BHSST, BHSST, BHSST-treated (1 g/kg body weight per day) CARE modeled-rats. Data are presented as mean ± SE. (*n* = 6). b*P* < 0.01 *vs* normal rats; d*P* < 0.01 *vs* untreated CARE modeled-rats. CARE: chronic acid reflux esophagitis; BHSST: Banhasasim-tang.



**Figure 4 Detection of TNF-α, p-JNK, Bcl-2 and c-Jun in chronic acid reflux esophagitis rats and effects of banhasasim-tang treatment.** Normal, normal rats; Control, untreated CARE modeled-rats; BHSST, BHSST-treated (1 g/kg body weight per day) CARE modeled-rats. Data are presented as mean ± SE. (*n* = 6). a*P* < 0.05 *vs* untreated CARE modeled-rats. CARE: chronic acid reflux esophagitis; BHSST: Banhasasim-tang.



**Figure 5 Expression of bax, cytochrome c and caspase 3 in chronic acid reflux esophagitis rats.** Normal, normal rats; control, untreated CARE modeled-rats; BHSST, BHSST, BHSST-treated (1 g/kg body weight per day) CARE modeled-rats. Data are presented as mean ± SE. (*n* = 6). a*P* < 0.05, b*P* < 0.01 *vs* normal rats; c*P* < 0.05, d*P* < 0.01 *vs* untreated CARE modeled-rats. CARE: chronic acid reflux esophagitis; BHSST: Banhasasim-tang.



**Figure 6 Possible mechanism of the banhasasim-tang protective mechanism in the esophagus of chronic acid reflux-induced esophageal ulcer in rats.** BHSST: Banhasasim-tang; ROS: reactive oxygen species.

**Table 1 Composition of the Banhasasim-tang formulation used in this study**

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
|  | **Herb** |  | **Amount in (g)** | | |
| Pinelliae rhizoma | | | | | 1.67 |
| Scutellariae radix | | | | | 1.00 |
| Zingiberis rhizoma siccus | | | | | 0.83 |
| Ginseng radix | | | | | 1.00 |
| Glycyrrhizae radix | | | | 1.00 | |
| Jujubae fructus | | | | 1.00 | |
| Coptidis rhizoma | | | | 0.33 | |

Banhasasim-tang Ex. granule was obtained from Hankook Shinyak Corp (Nonsan-si, Chungcheongnam-do, South Korea) in a pack of 3.5 g.

**Table 2 Body weight change and food intake**

|  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| **Group**  **Drug treatment** | **Body weight change (g) (after surgery)** | | | | | | | | | | | | | **Food intake**  **(g/d)** | | |
| **0 d** | | | **4 d** | | **7 d** | | **14 d** | | | **21 d** | | |
| **-** | | | **-** | | **(1 d)** | | **(8 d)** | | | **(15 d)** | | |
| Normal rats | |  | 169 ± 2.4 | | 170 ± 2.4 | | 219 ± 2.6 | | 278 ± 3.9 | | | 329 ± 5.6 | | | 27.32 ± 0.47 | |
| CARE modeled- rats | | | | | | | | | | | | | | | | |
| Untreated | |  | 166 ± 2.2 | 137 ± 2.0b | | 142 ± 2.7b | | | | 150 ± 7.5b | | | 178 ± 12.8b | | | 12.27 ± 0.63b |
| BHSST-treated | |  | 167 ± 1.8 | 135 ± 1.8 | | 146 ± 3.8 | | 169 ± 10.1 | | | | | 194 ± 19.1 | | | 14.99 ± 0.58d |

Data are presented as mean ± SE (*n* = 7). b*P* < 0.001 *vs* normal rats; d*P* < 0.01 *vs* untreated CARE modeled- rats.

**Table 3 Hematological analyses**

|  |  |  |
| --- | --- | --- |
| **Group** | **ROS**  **Fluorescence/min·mL** | **TBARS**  **nmol/mL** |
| Normal rats | 182 ± 14.5 | 3.56 ± 0.13 |
| CARE modeled- rats |  |  |
| Untreated | 289 ± 31.9b | 7.60 ± 0.77b |
| BHSST-treated | 134 ± 59.2c | 5.56 ± 0.36 |

Data are presented mean ± SE (*n* = 7). b*P* < 0.01 *vs* normal rats; c*P* < 0.05 *vs* untreated CARE modeled- rats. CARE: Chronic acid reflux esophagitis; ROS: Reactive oxygen species; TBARS: 2-Thiobarbituric acid reactive substance.