Stimulating Effects of KW-5092, a Novel Gastroprokinetic Agent, on the Gastric Emptying, Small Intestinal Propulsion and Colonic Propulsion in Rats

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Received July 8, 1994 Accepted October 21, 1994

ABSTRACT—KW-5092 (11-[[2-[[5-(piperidinomethyl)-2-furanyl]methyl]amino]ethyl]-2-imidazolidinylidene}propanedinitrile fumarate) is a novel gastroprokinetic agent with acetylcholinesterase (AChE) inhibitory activity and acetylcholine (ACh) release facilitatory activity. The present study examined the effects of KW-5092 on gastrointestinal (GI) propulsion in rats. KW-5092 at 1 to 30 mg/kg, p.o. dose-dependently enhanced the gastric emptying, small intestinal propulsion and the proximal and distal colonic propulsion. Metoclopramide, a dopamine D2-receptor antagonist with ACh release facilitatory activity, dose-dependently enhanced the gastric emptying at 0.03 to 1 mg/kg, p.o., whereas this drug did not affect the small intestinal propulsion, or the proximal and distal colonic propulsion. Neostigmine, an AChE inhibitor, dose-dependently enhanced the small intestinal propulsion and the proximal and the distal colonic propulsion at 0.3 to 10 mg/kg, p.o., whereas it delayed the gastric emptying at 10 mg/kg, p.o. The present results demonstrate that KW-5092 enhances the GI propulsion from the stomach to the colon and that metoclopramide or neostigmine enhances only the upper or the lower GI propulsion, respectively. Thus, KW-5092 may be a gastroprokinetic drug of a novel type for the treatment of GI motility dysfunctions in a wide range from the stomach to the colon.

Keywords: KW-5092, Metoclopramide, Neostigmine, Gastrointestinal propulsion

Metoclopramide has been demonstrated to possess gastroprokinetic activity and improve the gastrointestinal (GI) motor dysfunctions entailed in gastric stasis, gastroparesis or gastrosophageal reflux disease (1). Its action is ascribed to an antidopaminergic property (2, 3) and/or an increase in acetylcholine (ACh) release from postganglionic nerve endings (4, 5). On the other hand, neostigmine is a potent acetylcholinesterase (AChE) inhibitor (6) and increases colonic motility in human (7).

KW-5092 (11-[[2-[[5-(piperidinomethyl)-2-furanyl]methyl]amino]ethyl]-2-imidazolidinylidene}propanedinitrile fumarate) is a newly synthesized gastroprokinetic agent. In anesthetized rabbits (8) and in conscious and anesthetized dogs (9), KW-5092 enhanced GI motility, which was examined by a force transducer and was calculated by integrating the area between the baseline and the contractile wave, in a wide range from the gastric antrum to the colon. In the in vitro study, KW-5092 was a potent inhibitor of AChE derived from rat brain, the inhibitory activity being equipotent to that of neostigmine (8). Moreover, KW-5092 concentration-dependently enhanced the ACh release and the contraction of the isolated longitudinal muscle-myenteric plexus preparation from guinea pig ileum (10). Thus, the stimulation by KW-5092 of GI motor activity in vivo as well as in vitro has been ascribed to the AChE inhibition and the ACh release facilitation.

If the stimulation of GI motor activity does not lead to peristaltic contractions, GI propulsion may not be enhanced. Since KW-5092 possesses a novel and unique profile as a gastroprokinetic agent, it seemed of interest to examine the effect of this drug on the GI propulsion. Prior to the present paper, however, the effects of KW-5092 on the GI propulsion have not been reported. In the present study, we investigated the effects of KW-5092 on the gastric emptying, small intestinal propulsion and the colonic propulsion in rats and compared them with those of metoclopramide and neostigmine.
MATERIALS AND METHODS

Animals
Male Sprague-Dawley rats weighing 150 to 250 g were purchased from Japan SLC, Inc. (Hamamatsu). The animals were maintained on ordinary laboratory chow and tap water ad libitum under a constant 12-hr light-dark cycle.

Drugs
KW-5092 ([1-[2-[[5-(piperidinomethyl)-2-furanyl]-methyl]amino]ethyl]-2-imidazolidinylidene propanedinitrile fumarate) and metoclopramide hydrochloride were synthesized in our laboratories. Neostigmine methylsulfate was purchased from Sigma Chemical Co. (St. Louis, MO, USA). Phenol red, charcoal and trichloroacetic acid were purchased from Wako Pure Chemical Industries (Osaka). Arabic gum was purchased from Nacalai Tesque, Inc. (Kyoto). Test drugs were dissolved in saline and were orally administered to rats at a volume of 5 ml/kg.

Gastric emptying
The gastric emptying was elicited with a modification of the reported procedure (11). The animals were deprived of food 24 hr prior to the experiment but allowed free access to water until 3 hr before the experiment. A solution of 0.05% (w/v) phenol red in aqueous sodium carboxymethyl cellulose (1.5% w/v) was used as a test meal. The test drug was administered p.o. 1 hr before the test meal was given. Fifteen minutes after the test meal was given, the animals were sacrificed by cervical dislocation. The stomach was then exposed by laparotomy and the percentage traverse of charcoal meal in the stomach was determined.

Small intestinal propulsion
The small intestinal propulsion was determined according to the reported procedure (12). The animals were deprived of food 24 hr prior to the experiment but allowed free access to water until 3 hr before the experiment. A suspension of 10% (w/v) charcoal in aqueous Arabic gum (5% w/v) was used as a test meal. The test drug was administered p.o. 1 hr before the test meal was given. Ten minutes after the test meal was given, the animals were sacrificed by cervical dislocation. The small intestine was then exposed by laparotomy and the percentage traverse of charcoal meal in the small intestine was determined.

Proximal colonic propulsion
The proximal colonic propulsion was determined with a slight modification of the reported method (13). Each animal was anesthetized with pentobarbital sodium (50 mg/kg, i.p.), and the cecum was exposed by laparotomy. A vinyl tube of 1 mm in diameter was inserted into the cecum at the beginning of the colon. The other end of the tube was then taken out of the back. The animals were kept in individual cages for 4 to 5 days and deprived of food 24 hr prior to the experiment.

In the experiment examining the proximal colonic propulsion, a suspension of 5% (w/v) charcoal in aqueous Arabic gum (10% w/v) was used as a test meal. Each animal was lightly anesthetized with ether and administered 0.5 ml of the test meal into the colonic tubing. The test drug was administered p.o. 1 hr before the test meal was given. One hour after the test meal were given, the animals were sacrificed by cervical dislocation. The colon was then exposed by laparotomy, and the percentage traverse of charcoal meal in the colon was determined.

Distal colonic propulsion
The distal colonic propulsion was determined with the procedure reported previously (14). Each animal was lightly anesthetized with ether, and a teflon ball of 3 mm in diameter was inserted into the colon 3 cm proximal to the anus. The test drug was administered p.o. 1 hr before the teflon ball was inserted. The time required to evacuate the teflon ball was determined as an index of the distal colonic propulsion.

Statistical analyses
The result is expressed as the mean±S.E.M. Differences between the mean values in each drug treatment group and control group were analyzed by the Steel multiple comparison test following the Kruskal-Wallis test. A P value of less than 0.05 was considered statistically significant.
RESULTS

**Effects of drugs on the gastric emptying**

KW-5092 at 1 to 30 mg/kg, p.o. dose-dependently enhanced the gastric emptying and significantly enhanced it at 10 and 30 mg/kg, p.o. (Fig. 1A). Metoclopramide at 0.03 to 1 mg/kg, p.o. also dose-dependently enhanced the gastric emptying and significantly enhanced it at 0.3 and 1 mg/kg, p.o. (Fig. 1B). On the other hand, neostigmine significantly delayed the gastric emptying at 10 mg/kg, p.o. (Fig. 1C).

**Effects of drugs on the small intestinal propulsion**

KW-5092 at 1 to 30 mg/kg, p.o. dose-dependently enhanced the small intestinal propulsion; and at 10 and 30 mg/kg, p.o., the effect was statistically significant (Fig. 2A). Neostigmine at 0.3 to 10 mg/kg, p.o. also dose-dependently enhanced the small intestinal propulsion and significantly enhanced it at 3 and 10 mg/kg, p.o. (Fig. 2C). In contrast, metoclopramide at up to 100 mg/kg, p.o. did not affect the propulsion (Fig. 2B).

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**Fig. 1.** Effects of KW-5092, metoclopramide and neostigmine on the gastric emptying in rats. Each bar represents the mean ± S.E.M. of 10 rats. *: P<0.05, **: P<0.01, compared with the value in the control (0 mg/kg, p.o.) group (Steel multiple comparison test).

**Fig. 2.** Effects of KW-5092, metoclopramide and neostigmine on the small intestinal propulsion in rats. Each bar represents the mean ± S.E.M. of 8 rats. **: P<0.01, compared with the value in the control (0 mg/kg, p.o.) group (Steel multiple comparison test).
Effects of drugs on the proximal colonic propulsion

KW-5092 at 1 to 30 mg/kg, p.o. dose-dependently enhanced the small proximal colonic propulsion and significantly enhanced it at 3 to 30 mg/kg, p.o. (Fig. 3A). Neostigmine at 0.3 to 10 mg/kg, p.o. also dose-dependently enhanced the proximal colonic propulsion, and a significant effect was observed at 10 mg/kg, p.o. (Fig. 3C). On the other hand, metoclopramide at 3 to 100 mg/kg, p.o. did not affect the proximal colonic propulsion (Fig. 3B).

Effects of drugs on the distal colonic propulsion

KW-5092 at 1 to 30 mg/kg, p.o. dose-dependently decreased the time required to evacuate the teflon ball; and at 10 and 30 mg/kg, p.o., the effect was statistically significant (Fig. 4A). Neostigmine at 0.3 to 10 mg/kg, p.o. also dose-dependently decreased the time required to evacuate the teflon ball and significantly decreased it at 10 mg/kg, p.o. (Fig. 4C). In contrast, metoclopramide at 3 to 100 mg/kg, p.o. did not affect the distal colonic propulsion (Fig. 4B).

![Fig. 3. Effects of KW-5092, metoclopramide and neostigmine on the proximal colonic propulsion in rats. Each bar represents the mean±S.E.M. of 12 rats. *: P<0.05, **: P<0.01, compared with the value in the control (0 mg/kg, p.o.) group (Steel multiple comparison test).](image1)

![Fig. 4. Effects of KW-5092, metoclopramide and neostigmine on the distal colonic propulsion in rats. Each bar represents the mean±S.E.M. of 12 rats. *: P<0.05, **: P<0.01, compared with the value in the control (0 mg/kg, p.o.) group (Steel multiple comparison test).](image2)
DISCUSSION

The present study demonstrated that KW-5092 enhanced all the types of GI propulsion examined, from the stomach to the colon, in rats. In fact, KW-5092 is known to enhance the GI motility from the stomach to the colon in dogs (9). Taken together, it is reasonable to assume that the enhanced GI motility by KW-5092 can accompany the accelerated GI propulsion. On the other hand, metoclopramide enhanced only the gastric emptying, while neostigmine enhanced the small intestinal and the colonic propulsion but delayed the gastric emptying. Indeed, the inhibitory effect of neostigmine on the gastric emptying was also observed in horses (15). Thus, neostigmine seems to inhibit the gastric emptying, whereas this drug enhances the motility of the stomach as well as those of the lower GI tract (9). The inhibitory effect of neostigmine on the gastric emptying may be due to the inability of this drug to promote the gastroduodenal coordination. In any case, the present results suggest that KW-5092, unlike metoclopramide or neostigmine, may be a novel gastroprokinetic drug effective in a wide range from the stomach to the colon.

In the present study, metoclopramide, which is known to release ACh (4, 5), enhanced only the upper GI propulsion while neostigmine, an AChE inhibitor (6), enhanced only the lower GI propulsion in rats. These results are in accordance with the previous ones that metoclopramide enhances only the gastric and the duodenal GI motor activity in conscious dogs (16) and that neostigmine increases the colonic GI motility in horses (17). These observations as well as ours suggest that the enhanced propulsion and motor activity of the upper gut are mainly associated with the ACh release facilitation and that those of the lower gut are mainly associated with the AChE inhibition. The regional difference in the sensitivity between the ACh release facilitator and the AChE inhibitor may be due to the heterogeneity of cholinergic innervation and AChE distribution. Further studies, however, are required to clarify this point. KW-5092 inhibits AChE (8) and facilitates ACh release from the isolated intestine (11) at similar concentrations. Thus, the combined effect of AChE inhibition and ACh release facilitation seems to be involved in the enhancement by KW-5092 of the propulsion as well as the motor activity in a wide range of the gut. In fact, the combination of neostigmine (10 mg/kg, p.o.) and metoclopramide (1 mg/kg, p.o.) enhanced both the gastric emptying and the distal colonic propulsion in rats (N. Kishibayashi et al., unpublished observation).

In addition to metoclopramide, other gastroprokinetic agents, such as domperidone (18) and cisapride (19), have been effectively used for treatment of GI motility dysfunctions associated with gastric stasis, gastroparesis or gastroesophageal reflux disease. Domperidone enhances gastroduodenal coordination via blockade of a specific dopamine receptor (18) and enhances only the antral and the duodenal motor activity in conscious dogs (20). On the other hand, cisapride enhances ACh release without affecting dopamine receptors (19) and enhances the GI motor activity from the gastric antrum to the colon in conscious dogs (15). The stimulating effect by cisapride of the GI motor activity has been ascribed to its ACh release facilitation (19), although the precise mechanism involved remains obscure. In rats, however, cisapride enhances the gastric emptying but does not stimulate the intestinal or the colonic propulsion (N. Kishibayashi et al., unpublished observation). In contrast, KW-5092 enhances the propulsion and the motor activity in a wide range of the GI tract not only in dogs (9) but also in rats (this study). Moreover, KW-5092, but not cisapride, ameliorated the colonic propulsion delayed by clonidine or loperamide in rats (21). KW-5092 could thus be a gastroprokinetic drug of a novel type that may be a possible treatment for a variety of GI motility dysfunctions.

In conclusion, the present study demonstrates that KW-5092 enhances the GI propulsion in a wide range from the stomach to the colon, whereas metoclopramide enhances only the upper GI propulsion, and neostigmine enhances only the lower GI propulsion. KW-5092 may be a useful drug for the treatment of the GI motility dysfunctions not only in the upper but also in the lower GI tract.

Acknowledgments

We wish to thank S. Sasho for preparation of KW-5092 and thank Drs. A. Ishii and T. Hirata for encouragement and support.

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