Effect of antipyrine on the gastric emptying of liquid in rats

Abstract

Antipyrine (At) and dipyrone (Dp) delay gastric emptying (GE) in rats. The objective of the present study was to assess the effects of intravenous (iv) and intracerebroventricular (icv) administration of At and Dp on the GE of liquid by rats. GE was assessed in male Wistar rats (5-10 in each group) 10 min after the icv or iv drug injection by measuring percent gastric retention (%GR) of a saline test meal labeled with phenol red 10 min after administration by gavage. The At iv group was significantly higher (64.4 ± 2.6%) compared to control (33.4 ± 1.5%) but did not differ from the Dp group (54.3 ± 3.8%). After icv administration of At, %GR (34.2 ± 2%) did not differ from control (32.6 ± 1.9%), but was significantly higher after Dp (54.5 ± 2.3%). Subdiaphragmatic vagotomy significantly reduced %GR in the At group (30.2 ± 0.7%) compared to the sham group, but was significantly higher than in the controls (23.0 ± 0.5%). In the animals treated with At iv, baclofen significantly reduced %GR (28.3 ± 2.4%) compared to vehicle-treated animals (55.2 ± 3.2%). The same occurred in the animals treated iv with vehicle and icv with baclofen. Although vagotomy and baclofen reduced %GR per se, the reduction was twice more marked in the animals treated with At. The results suggest that At administered iv, but not icv, delays GE of liquid in rats with the participation, at least in part, of the vagus nerve and that this phenomenon is blocked by the activation of GABA_B receptors in the central nervous system.

Introduction

Gastric emptying (GE) is the process of the transfer of the gastric content to the small intestine as the result of the motor activity of the stomach, pylorus and duodenum under control of inhibitory and stimulatory mechanisms (1).

Non-steroidal anti-inflammatory drugs are among the most frequently prescribed drugs in the world and have been extensively studied because of their gastrointestinal toxicity. Among the effects of these medications, changes in GE have been reported to be induced by aspirin, acetaminophen and indomethacin (2-6). Dipyrone (Dp), a phenylpyrazolone derivative mainly used for its antipyretic and analgesic properties, causes delayed GE in rats when administered at high doses (7). This effect was confirmed
using a dose of 80 mg/kg (240 µmol/kg) administered intravenously (iv) and saline solution as a test meal (8). A recent study in which Dp was administered iv and intracerebroventricularly (icv) to rats has suggested that this effect may be due to the action of the drug on the central nervous system (CNS), with the participation of the paraventricular nucleus and of the vagus nerve (9). Additional observations have demonstrated that this phenomenon is blocked by icv administration of baclofen (bac), a GABAB receptor agonist (10).

Takanaka et al. (11), in a study on the effect of antipyrine (At, phenazone), another phenylpyrazolone derivative, on the GE of rats, observed delayed GE after its administration. However, the site of action and the mechanisms involved in this phenomenon are unknown.

The objective of the present study was to assess the effect of At on the GE of rats, using saline solution as a test meal. The following procedures were carried out: 1) comparative studies with Dp using equivalent doses administered iv (240 µmol/kg) and icv (4 µmol/animal); 2) study of the dose-response effect and time-response effect of At administered iv; 3) study of the effect of subdiaphragmatic vagotomy (VgX) on the iv administration of the drug; 4) study of the effect of icv administration of bac on the iv administration of the drug.

Material and Methods

The study was conducted on 8-week-old male Wistar rats weighing 250-300 g, adapted to laboratory conditions for 4 weeks and receiving water and food ad libitum. The experimental protocols used in the present study followed the recommendations of the Brazilian College of Animal Experimentation (COBEA).

The drugs used, dipyrone, antipyrine, and baclofen (all from Sigma, St. Louis, MO, USA) were diluted in sterile physiological saline as a vehicle at the time of the experiments.

The GE test was carried out in animals fasted for 24 h but with free access to water, which was removed 30 min before the experiments.

For the comparative study of At and Dp injected iv the animals were divided into three groups: At and Dp groups, respectively receiving iv 45.17 and 80 mg/kg of the drug corresponding to the dose of 240 µmol/kg, and control group (C) receiving vehicle (1 mL/kg body weight). Drugs were injected iv through a tail vein and GE was assessed 10 min after administration.

For the assessment of the effect of At and Dp administered icv, a cannula was implanted into the right lateral ventricle of each animal 8 days before the experiments using previously established coordinates (9,10). For icv injection, 10 µL saline (C), Dp (4 µmol = 1333.2 µg), or At (4 µmol = 752.8 µg) was used. The dose (in µmol) and the technique employed for icv administration had been established in a previous study (9). GE was assessed 10 min after the end of the icv injection.

For the study of the dose-response effect, iv doses of 60 (11.25 mg/kg), 120 (22.5 mg/kg), and 240 µmol/kg (45.17 mg/kg) At were used. Control animals (C) received vehicle by the same route. GE was assessed 10 min after administration. For the study of the time-response effect, At or vehicle was administered iv at the dose of 240 µmol/kg (45.17 mg/kg), and GE was assessed 10, 30, 60, and 120 min after administration.

To assess the participation of the vagus nerve in the effect of At on GE, the rats were submitted to subdiaphragmatic section of the ventral and dorsal branches of the vagus (VgX group) 2 weeks before the experiment. The sham group consisted of rats submitted to surgery without section of the vagal branches. Each group was divided into subgroups of animals receiving vehicle (C) or At at the dose of 240 µmol/kg (45.17 mg/kg).
kg) iv. GE was assessed 10 min after administration.

To assess the action of bac on the effect of At, a solution of 2 µg bac (bac2) or vehicle (bac0) was applied icv (10 µL) and the animals then received a solution of At, 240 µmol/kg, or vehicle (C) iv. The baclofen dose and the technique employed for icv administration had been established in a previous study (10). GE was assessed 10 min after the end of the iv injection.

In all experiments, GE was evaluated indirectly by measuring percent gastric retention (%GR) of a saline test meal labeled with phenol red (60 µg/mL) in a volume of 2 mL/100 g body weight administered by gavage using a standardized technique (12). The absorbance of the dye was determined with a spectrophotometer at 560 nm. At the end of the experiment all animals were sacrificed and the brain of those submitted to icv injection was injected with 10 mL 1% Evans blue through the cannula. The brain was removed and fixed in 10% formalin for 24 h. After fixation, coronal sections were obtained and the proper site of injection was confirmed when the dye was observed in the fourth ventricle.

Statistical analysis

Data are reported as means ± SEM. Statistical analysis was carried out using ANOVA followed by the Tukey test when necessary. When it was not possible to use parametric tests, the Kruskal-Wallis test was used, followed by the multiple comparisons Dunn test when necessary. The level of significance was set at $\alpha = 5\%$ for all tests.

Results

The data regarding the effect of Dp and At administered iv are presented in Figure 1A. The At and Dp groups presented a significantly more elevated %GR compared to control. The At group showed higher %GR values (mean ± SEM = 64.4 ± 2.6%) which, however, did not differ significantly from Dp (54.3 ± 3.8%). The data regarding the effect of Dp and At administered icv are presented in Figure 1B. There was a significant increase in %GR in the Dp group (54.5 ± 2.3%) compared to the C (32.6 ± 1.9%) and At (34.2 ± 2%) groups, whereas the At group did not differ significantly from control.

The dose-response study (Figure 2A) showed a dose-dependent increase in %GR in the animals treated with At ($r = 0.86$) which, however, was significant only at the dose of 240 µmol/kg (64.5 ± 4.7%) compared to control (37.4 ± 1.2%). Regarding the time-response study, the At group showed a maximum effect of the drug on %GR at 10 min (67.6 ± 3%) compared to control (37.4 ± 1.3%). A progressive reduction occurred, with the maintenance of a significant difference from control at 30 and 60 min which, however, no longer existed at 120 min (Figure 2B).

![Figure 1. Gastric retention of a saline test meal 10 min after administration to rats by gavage. A. Ten minutes before the evaluation of gastric retention, the animals received intravenous (iv) vehicle (C) or 240 µmol/kg dipyrrone (Dp; 80 mg/kg), or 240 µmol/kg antipyrine (At; 45.17 mg/kg). B. Eight days before the test, a metal cannula was implanted into the right lateral ventricle of each rat. The animals received intracerebroventricular (icv) 10 µL of vehicle (C) or an equal volume of a solution containing 4 µmol (1333.2 µg) Dp or 4 µmol (752.8 µg) At. Gastric retention was assessed 10 min after the icv injections. Data are reported as means ± SEM for 10 animals per group. *P < 0.05 for the comparisons indicated by the arrows (Tukey test).](image-url)
A. The bac0At group presented a significantly higher %GR (55.2 ± 3.2%) compared to the bac0C group (34.0 ± 1.8%). On the other hand, the %GR of the bac2At group (28.3 ± 2.4%) did not differ significantly from control (bac2C, 25.2 ± 3.2%). Baclofen administered icv significantly reduced %GR in the animals that received At iv (bac2At vs bac0At) and in the control groups that received saline iv (bac2C vs bac0C); however, the reduction was approximately twice higher in the animals receiving At (a 48.7% reduction in mean %GR) compared to control (a 26.1% reduction in mean %GR).

Discussion

We observed in the present study that antipyrine administered iv delayed GE of liquids in rats in a dose-dependent manner, a phenomenon that differed from the effect observed with dipyrone. First, the effect of antipyrine was more intense during the first 10 min after iv administration, whereas the effect of dipyrone was approximately constant during the first hour (9). Second, when the drug was administered icv it did not change GE, indicating that it has no direct effect on the CNS, in contrast to what was observed with the administration of dipyrone. These facts suggest that the two drugs, although having a common chemical origin as phenylpyrazolone derivatives, cause delayed GE by distinct mechanisms.

After per os administration to humans, antipyrine is rapidly and fully absorbed and is then slowly metabolized, with a mean plasma clearance rate of 6% per hour (13). The hepatic isoenzymes of cytochrome P450 participate in these metabolic processes, giving origin to metabolites which are mainly eliminated through the kidneys (14,15). If we assume that the observed delay in GE is due to the action of antipyrine and not of its metabolites, it is not clear why the drug, which continues to circulate and whose clearance is slow, has the most pronounced effect
on GE 10 min after administration. Two possibilities may be proposed to explain this discrepancy: 1) The drug is inactivated or gradually metabolized at the site where it induces the delay, or 2) the mechanism that antagonizes the effect of delayed GE is able to partially counterbalance the effect of antipyrine.

Subdiaphragmatic vagotomy is known to modify the motor activity of the stomach, blocking the arrival of afferent stimuli to the tractus solitarius and of stimulating and inhibitory efferents originating from the dorsal nucleus of the vagus (16-18). In the present study, vagotomy reduced the effect on GE induced by iv administration of antipyrine, suggesting the participation of vagal pathways in this phenomenon. Since this procedure did not fully abolish the effect, we may conclude that the drug acts by a mechanism that does not depend, at least in part, on the participation of the vagus nerve.

In addition, the effect of iv antipyrine on GE was blocked by icv baclofen. The activation of GABA_B receptors by baclofen in the CNS results in hyperpolarization of the postsynaptic membranes or in inhibition of neurotransmitter release at presynaptic nerve endings (19,20). There is strong evidence that GABA, through GABA_B receptors, may participate in the control of gastric secretion and gastrointestinal motility at the level of the nucleus tractus solitarii (19,21-26). On this basis, we may speculate that the stimulation of presynaptic GABA_B receptors by baclofen, by blocking the excitation of the non-adrenergic cholinergic (inhibitory) pathway and/or the inhibition of the cholinergic (stimulatory) pathway of the vagus determined by antipyrine, may explain the results of the present study. Probably this is not the only hypothesis if we consider that vagotomy and baclofen administration in the CNS reduced %GR per se, although the reduction was twice more marked in the animals that received antipyrine. On the basis of the present results, no other speculations would be valid here.

The present results suggest that antipyrine administered iv, but not icv, delays GE of liquid in rats with the participation, at least in part, of the vagus nerve and that this phenomenon is blocked by the activation of GABA_B receptors in the CNS.
References

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