EFFECT OF VASOACTIVE DRUGS ON GASTRIC BLOOD FLOW MEASURED BY A CROSS THERMOCOUPLE METHOD IN RATS

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Abstract—To make a continuous recording of gastric blood flow (GBF) in rats, application of a cross thermocouple method was investigated together with related pharmacological studies. When secretagogues (tetragastrin, histamine and methacholine) were given intravenously in a dose sufficient to stimulate acid secretion, the increases in GBF observed were much the same as those seen when the aminopyrine clearance technique was used. When epinephrine and norepinephrine were administered via a close intrarterial route in the stomach, there was an initial decrease followed by an increase in the GBF. This biphasic response was antagonized by phentolamine, but not by propranolol, thereby suggesting that those agonists predominantly stimulate α-adrenoceptors. On the contrary, isoprenaline produced an increase in GBF which was attenuated by propranolol. Acetylcholine produced an increase in GBF, which was blocked by atropine. Histamine increased the GBF, and such was inhibited by diphenhydramine, but not by cimetidine, suggesting a stimulation of H-1 receptors by histamine. Serotonin, in a lower dose, and tetragastrin also elicited an increase in GBF. However, with a higher dose of serotonin, there was an increase followed by an apparent decrease in GBF. From these results, it was concluded that the cross thermocouple method is practical for a continuous recording the rat GBF in response to vasoactive drugs.

Gastric blood flow is closely associated with gastric physiological functions or pathophysiological changes of gastric mucosa subjected to ulcerogenic conditions (1-4). With respect to methods for continuously measuring gastric blood flow, however, few methods provide such estimations of gastric blood flow in small animals such as rats (5, 6) which are widely used for pharmacological studies. A cross thermocouple method is a simple, practical one that enables a continuous monitoring of tissue blood flow on the basis of heat clearance (7), and has so far been applied to many tissues (8). However, if thin or small tissues have to be used for study, there is a considerable difficulty in implanting a thermocouple element in the exact position and the responses are thus less pronounced.

The present work was an attempt to demonstrate that the cross thermocouple method is applicable to the rat gastric mucosa and can be compared to findings with the aminopyrine clearance technique. Using the cross thermocouple method, the effect of various vasoactive drugs on gastric tissue blood flow (GTBF) was studied regarding features of microcirculation in the
MATERIALS AND METHODS

Male Wistar rats weighing 300 to 330 g were deprived of food 16 hr before each experiment but water was allowed ad libitum. After anesthesia with urethane (2 g/kg, i.p.), the trachea was cannulated and a midline laparotomy done to expose the stomach. Polyethylene tubings were inserted to three positions; the right carotid artery for blood sampling and/or measurement of blood pressure, the left femoral vein for an i.v. administration of drugs, and the splenic artery, retrogradely, for a close intra-arterial (i.a.) injection of drugs to the stomach (Fig. 1).

Cross thermocouple method: A wire-typed cross thermocouple element (W-41, Unique Medical) was transversally implanted, using a sewing needle, into the gastric wall of the glandular portion with a serosal access (Fig. 1). In this way, the thermocouple elements were microscopically demonstrated to be located mostly within the mucosal layer.

Electric potential, which was theoretically considered to correlate with tissue blood flow on the basis of heat clearance (7), was monitored on a recorder (056, Hitachi) via an amplifier (UM 2000, Unique Medical). Changes of GTBF by gastric secretagogues and by antagonists to vasoactive agents were expressed as a potential change (mV) from the potential levels in the basal state and as a percentage (%) of the potential levels of those control responses, respectively.

Aminopyrine clearance method: Measurement of gastric mucosal blood flow (GMBF) using aminopyrine clearance was made by the method of Jacobson et al. (9) on the basis of the principle described by Shore et al. (10). After a loading dose (20 mg/kg) of aminopyrine had been infused through the femoral vein over a period of 10 min, a maintenance dose (5 mg/kg/hr) was continuously infused using a perfusion pump (KN-201 D, Natsume). The stomach was washed several times with saline (0.9% NaCl solution) through an opening made in the forestomach. A rubber catheter for perfusing acidic saline (pH 3.0) was transorally introduced into the stomach, and the tip was placed below the cardia. Another polyvinyl catheter for sampling gastric perfusates was also inserted into the pyloric portion through the opening in the forestomach. After pyloric ligation, perfusion of the acidic saline was started at a rate of 0.5 ml/min using a perfusion pump (TMP-15 E, Toyo-Kagaku-Sangyo). Gastric perfusates were collected every 30 min, their acid concentrations were titrated with 0.02 N NaOH using a pH stat (RAT-11S, Hiranuma-Sangyo), and the concentrated perfusates, as well as blood samples, were used for the determination of aminopyrine concentrations. Blood samples were collected at the end of each experiment, as our preliminary studies demonstrated that aminopyrine blood concentrations of each rat were fairly constant.
during a period of 5 hr from the 1st to the 6th hour after aminopyrine infusion.

Drugs: Drugs used were acetylcholine chloride (Daiichi), DL-norepinephrine hydrochloride, L-epinephrine hydrochloride (Sankei), L-isoprenaline hydrochloride (Tokyo Kasei), methacholine chloride, histamine dihydrochloride, atropine sulfate, serotonin (Wako), tetragastrin (Teikokuzohki), phen tolamine mesylate (CIBA), propranolol hydrochloride (Sumitomo), diphenhydramine hydrochloride (Kowa), and cimetidine (SKF). These drugs were dissolved or suspended in saline. Vasoactive agonists were given via an i.a. route. Their antagonists, when necessary, were given i.v. 5-10 min before the administration of agonists. Drug doses were expressed in terms of the salt.

Statistical significance was evaluated using Student’s t-test.

RESULTS

1. Evaluation of the cross thermocouple method for determination of gastric blood flow and a comparison of the findings with use of the aminopyrine clearance technique:

Two methods for measurement of blood flow, i.e. cross thermocouple and aminopyrine clearance, were compared by simultaneously determining responses of gastric blood flow to three different secretagogues. According to the cross thermocouple method, an increase in GTBF occurred at the first 5-min period and subsided at the second 5-min period after administration of each secretagogue (Fig. 2). When measured by the aminopyrine clearance technique, an increase in GMBF appeared at the first 30-min period and disappeared at the next period (Fig. 3). As to responses to individual secretagogues, histamine (300 µg/kg, i.v.) and tetragastrin (7.5 µg/kg, i.v.), but not methacholine (8 µg/kg, i.v.), produced a significant increase in gastric blood flow by means of either method. On the other hand, all secretagogues, in tested doses, significantly stimulated gastric acid secretion (Fig. 3).

**Fig. 2.** Responses of GTBF to secretagogues by the cross thermocouple method. Sal: saline. Hist: histamine (300 µg/kg). T.G.: tetragastrin (7.5 µg/kg). MeCh; methacholine (8 µg/kg). All drugs were given i.v. The results of GTBF were estimated as the difference in the averaged potential level at each 5-min interval from that observed before treatment, and expressed as the mean±S.E. of 9 experiments. *P<0.05, **P<0.01, when compared with the corresponding value of saline treatment.

**Fig. 3.** Responses of GMBF and acid secretion to secretagogues by the aminopyline clearance method. Pre: pretreatment. Other abbreviations or treatments are shown in Fig. 2. Results of GMBF and acid output were expressed as the mean±S.E. of 9 experiments at each 30-min interval. The acid output was estimated as the difference in the amount of hydrogen ion from that observed during pretreatment. *P<0.05, **P<0.01, ***P<0.001, when compared with the value of pretreatment.
2. Responses of GTBF to vasoactive drugs with or without their antagonists, using the cross thermocouple method:

Typical patterns of GTBF in response to each vasoactive drug are shown in Figs. 4 and 5. The antagonisms of responses of

Fig. 4. Typical patterns of responses of GTBF to vasoactive drugs with or without their antagonists. Nor; norepinephrine. Epi; epinephrine. Iso; isoprenaline. Phent; phentolamine. Prop; propranolol. Agonists were given i.a., while antagonists were given i.v.

Fig. 5. Typical patterns of responses of GTBF to vasoactive drugs with or without their antagonists. ACh; acetylcholine. Hist; histamine. 5-HT; serotonin. T.G.; tetragastrin. Atr; atropine. Cimet; cimetidine. Diphen; diphenhydramine. Agonists were given i.a., while antagonists were given i.v.
Table 1. Inhibition of the increasing or decreasing response of GTBF to vasoactive drugs by pretreatment with their antagonists

<table>
<thead>
<tr>
<th>Antagonists</th>
<th>Agonists</th>
<th>µg/rat</th>
<th>No. of rats</th>
<th>Decrease (% Control)</th>
<th>Increase (% Control)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1) Phentolamine</td>
<td>Norepinephrine</td>
<td>0.125</td>
<td>4</td>
<td>51.7±12.5*</td>
<td>14.0±9.9**</td>
</tr>
<tr>
<td>(1 mg/kg)</td>
<td>Epinephrine</td>
<td>0.25</td>
<td>4</td>
<td>51.4±7.9**</td>
<td>28.0±10.1**</td>
</tr>
<tr>
<td></td>
<td>Epinephrine</td>
<td>0.50</td>
<td>4</td>
<td>36.7±1.9***</td>
<td>14.9±5.4***</td>
</tr>
<tr>
<td></td>
<td>Epinephrine</td>
<td>0.125</td>
<td>3</td>
<td>41.3±12.9*</td>
<td>47.2±9.9*</td>
</tr>
<tr>
<td></td>
<td>Epinephrine</td>
<td>0.25</td>
<td>10</td>
<td>38.9±8.4***</td>
<td>31.5±8.8***</td>
</tr>
<tr>
<td>2) Propranolol</td>
<td>Norepinephrine</td>
<td>0.125</td>
<td>3</td>
<td>102.0±16.1</td>
<td>122.6±21.6</td>
</tr>
<tr>
<td>(1 mg/kg)</td>
<td>Epinephrine</td>
<td>0.25</td>
<td>3</td>
<td>121.1±23.6</td>
<td>129.1±21.2</td>
</tr>
<tr>
<td></td>
<td>Epinephrine</td>
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<td>3</td>
<td>96.8±23.1</td>
<td>142.4±54.1</td>
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<tr>
<td></td>
<td>Epinephrine</td>
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<td>5</td>
<td>100.0±12.4</td>
<td>85.3±19.9</td>
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<tr>
<td></td>
<td>Isoprenaline</td>
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<td>40.9±12.4*</td>
<td>45.9±3.5***</td>
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<td></td>
<td>Isoprenaline</td>
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<td>11</td>
<td>6.2±3.8***</td>
<td>8.5±3.7***</td>
</tr>
<tr>
<td>3) Atropine</td>
<td>Acetylcholine</td>
<td>0.25</td>
<td>4</td>
<td>46.8±17.9***</td>
<td>53.8±13.5***</td>
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<tr>
<td>(0.1 mg/kg)</td>
<td>Acetylcholine</td>
<td>0.50</td>
<td>5</td>
<td>46.8±17.9***</td>
<td>57.5±14.1***</td>
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<tr>
<td>4) Diphenhydramine</td>
<td>Histamine</td>
<td>1.0</td>
<td>11</td>
<td>103.5±8.0</td>
<td>109.4±12.3</td>
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<tr>
<td>(10 mg/kg)</td>
<td>Histamine</td>
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<td>90.0±6.5</td>
<td>90.0±6.5</td>
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<tr>
<td></td>
<td>Histamine</td>
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<td>78.5±5.8</td>
<td>78.5±5.8</td>
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<tr>
<td>5) Cimetidine</td>
<td>Histamine</td>
<td>1.0</td>
<td>9</td>
<td>30.5±5.8***</td>
<td>30.5±5.8***</td>
</tr>
<tr>
<td>(5 mg/kg)</td>
<td>Histamine</td>
<td>5.0</td>
<td>9</td>
<td>38.5±6.6***</td>
<td>38.5±6.6***</td>
</tr>
<tr>
<td></td>
<td>Histamine</td>
<td>10.0</td>
<td>9</td>
<td>51.0±1.8***</td>
<td>51.0±1.8***</td>
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</tbody>
</table>

a) Responses of GTBF to an agonist were obtained before and after its antagonist(s). The extent of the latter responses was expressed as a percentage (the mean ± S.E.) of the former potential level (control responses, i.e. 100%). The results were analyzed statistically using a paired t-test. *p<0.05, **p<0.01, ***p<0.001, when compared with controls.

GTBF between vasoactive drugs and their antagonists are presented in Table 1.

Adrenergic agonists: Norepinephrine and epinephrine, which were tested in 0.125 and 0.50 µg/rat, produced a biphasic response of GTBF; a decrease was initiated about 1 min after injection, followed by an increase about 2 min later. When the same dose was injected, epinephrine produced a more prominent action than did norepinephrine. Pretreatment with phentolamine, 1 mg/kg, significantly antagonized this biphasic response to norepinephrine and epinephrine, while pretreatment with propranolol, 1 mg/kg, produced no appreciable change. Isoprenaline, 0.125 and 0.25 µg/rat, produced an increase in GTBF, and the peak of the increase was about 1 min after injection. This response was significantly blocked by propranolol, 1 mg/kg.

Acetylcholine: Administration of acetylcholine, 0.25 and 0.50 µg/rat, caused an increase in GTBF. The peak of the response occurred within 1 min after injection. Pretreat-
ment with atropine, 0.1 mg/kg, greatly inhibited this increase.

Histamine: Histamine, 1–10 µg/rat, elicited an increase in GTBF. The peak of the increase was 1–2 min after injection. This was significantly antagonized by pretreatment with diphenhydramine, 10 mg/kg, but not with cimetidine, 5 or 20 mg/kg. A combination of these antagonists resulted in a somewhat greater inhibition as compared with the administration of diphenhydramine alone.

Serotonin: Patterns of responses to serotonin varied with the dosage. An increase in GTBF was elicited at 0.25 µg/rat of the drug, but was preceded by an apparent decrease when injected at 1.0 µg/rat. With a higher dose, the peak of the decrease was about 1 min after injection and that of the increase about 2 min after injection.

Tetragastrin: Administration of tetragastrin, 1.5 µg/rat, exerted an increase in GTBF, and the peak of the increase was about 1 min after injection.

Effect of i.a. administration of vasoactive drugs on blood pressure was estimated. Some of these drugs caused a moderate change in blood pressure at higher doses, but there was no case in which evaluation of their primary action on GTBF was made difficult by a modified blood pressure.

DISCUSSION

It is generally accepted that the aminopyrine clearance method is capable of assessing total mucosal blood flow in the stomach (2). In contrast with it, the cross thermocouple method measures regional blood flow at the area where a heat-sensitive element is implanted (7). Despite the thinness of the rat gastric mucosa, however, the implanted element was found to be placed mostly within the mucosal layer. Under such experimental situations, the experimental findings with the cross thermocouple method could be compared with those of the aminopyrine clearance technique regarding direction of responses of blood flow to three different secretagogues at dosages sufficient to stimulate gastric acid secretion. Moreover, it has been suggested that GMBF may occupy half the amount of total blood flow in rats (10). Thus, we considered that the gastric blood flow estimated by this method, i.e. GTBF, may well reflect the GMBF and that the cross thermocouple method is of practical value in evaluating the rat brief phasic changes.

There is apparently no documentation regarding continuously recorded rat gastric blood flow in response to vasoactive drugs. In studies using cats or dogs, drug responses are often variable, probably because of difference in dosage, route, or methodology (11, 12). The present experiments on drug responses, therefore, were designed so as to exhibit the primary action of drugs on gastric blood flow using an i.a. route of administration. Norepinephrine and epinephrine caused a biphasic response of blood flow, namely, a decrease followed by an increase in GTBF. The decreasing response, which occurs earlier, is assumed to be elicited by α-adrenergic stimulation, because the response was blocked by phentolamine. On the other hand, mechanisms of eliciting the increase in GTBF seem to be complex. There is the possibility that β-adrenergic stimulation is responsible for the increase in GTBF, as was suggested from previous studies in dogs (12) and cats (13). However, our experiments with propranolol failed to support this possibility. Another possibility is that the increasing response originates from a decrease in GTBF itself, instead of being elicited by a direct action of nor-epinephrine or epinephrine. Such may be caused by the opening of the arterio-venous shunts which respond to the decrease (14), by autoregulatory escape mechanisms in the
vessels which exhibit the decreasing response (13, 15, 16), or by reactive hyperemia (17) possibly due to metabolic feedback which occurs during the decreasing response. Of these possible mechanisms, the presence of functional arterio-venous shunts in the rat stomach was not confirmed in the in vivo microscopical study of Guth and Smith (5). Accordingly, norepinephrine and epinephrine may elicit a decrease in GTBF through α-adrenergic stimulation, leading to a secondary increase in GTBF through autoregulatory or metabolic mechanisms. On the other hand, only isoprenaline produced an increase in GTBF which was antagonized with propranolol. Thus, there probably are β-adrenergic receptors in the rat gastric vessels although the distribution is diffuse or the sensitivity is poor.

Histamine exerts an increase in gastric blood flow in association with stimulation of gastric secretion via histamine H-2 receptors (18). However, it has been reported that the increasing response of gastric blood flow to histamine develops even in the presence of H-2 receptor antagonists (1) and is reduced by pretreatment with H-1 receptor antagonists (19). We also found that the increasing effect of histamine on GTBF was significantly inhibited with diphenhydramine but not with cimetidine although the antagonism was slightly augmented by a combination of these drugs. Thus, histamine probably increases gastric blood flow predominantly through stimulation of H-1 receptors in the rat gastric vasculature. Different responses of gastric blood flow to cholinergic drugs have been reported (20, 21). Likewise, vagal stimulation was reported to increase (22) or to decrease (23) in gastric blood flow. Our present results, including the antagonism between acetylcholine and atropine, favor the idea that acetylcholine exerts an increase in gastric blood flow through excitation of specific receptors.

Serotonin and gastrin are distributed mainly in the gastrointestinal tract. However, their primary actions on gastric blood flow of rats have never been elucidated. According to our results, serotonin causes an increase in GTBF but augments a decreasing response of GTBF with increasing dosage. Tetragastrin, a gastrin-like peptide, produced an increase in GTBF immediately after administration. Similar findings with pentagastrin have also been reported in the case of the canine stomach (19). The presence of the immediate response suggests that gastrin may increase gastric blood flow through its vasoactive actions other than mechanisms accompanying the stimulation of acid secretion.

It is essential to evaluate the primary responses of gastric blood flow to vasoactive drugs in order to understand the physiological or pharmacological actions on gastric functions. As the cross thermocouple method provides for a continuous recording of the gastric blood flow in rats, this method may be suitable for such evaluation.

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