Role of Vasoactive Intestinal Polypeptide in the Internal Anal Sphincter Relaxation of the Opossum

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Abstract

The nature of the inhibitory neurotransmitter responsible for internal anal sphincter (IAS) relaxation in response to rectoanal reflex is not known. The objective of the present investigation was to examine the role of VIP in IAS relaxation in response to the rectoanal reflex in intact opossums with the use of VIP antagonists, [4Cl-d-Phe\(^6\),Leu\(^7\)] VIP and (N-Ac-Tyr\(^{\beta}\),D-Phe\(^{\gamma}\))-GRF (1-29)-NH\(_2\). Intraluminal pressures from the sphincter were monitored using low-compliance, continuously perfused catheters. VIP and the antagonists were administered close-intraarterially. The responses to VIP, rectoanal reflex, sacral nerve stimulation, and local intramural stimulation were examined before and after the VIP antagonists. The present studies in intact animals show: (a) VIP causes a dose-dependent fall in the IAS pressures by a direct action at the IAS smooth muscle; (b) VIP antagonists selectively and significantly antagonized the inhibitory action of VIP; and (c) VIP antagonists caused significant antagonism of the IAS relaxation caused by rectoanal reflex and the other neural stimuli. The antagonism of the IAS relaxation by the VIP antagonists, depending upon the volume of rectal distension used, ranged from 46% to 62% (P < 0.05). From these results, we conclude that VIP acts as an inhibitory neurotransmitter for IAS relaxation during the rectoanal reflex.

Introduction

It has been previously shown that internal anal sphincter (IAS) relaxation is mediated through the release of a noncholinergic, nonadrenergic inhibitory neurotransmitter (1-5). Such noncholinergic, nonadrenergic nerves play a significant role in the relaxation of a variety of smooth muscle tissues (6-7). However, the exact nature of the inhibitory neurotransmitter responsible for the relaxation of these smooth muscle tissues and IAS relaxation is not known. It has been suggested that vasoactive intestinal polypeptide (VIP) acts as a neurotransmitter in certain systems (8-13). Previous in vivo and in vitro studies in the lower esophageal sphincter (LES) (11, 14) and in vitro studies in the IAS (3) using VIP antisera have suggested that VIP may play a role in IAS relaxation. However, studies using newly discovered VIP antagonists (15, 16) to examine the participation of VIP as an inhibitory neurotransmitter have not been performed in vivo or in vitro for any smooth muscle including the IAS.

The aim of the present investigation was to examine the possibility of VIP as a noncholinergic, nonadrenergic inhibitory neurotransmitter responsible for the relaxation of the internal anal sphincter in response to the rectoanal reflex (mimicked by rectal balloon distension).

Methods

Animal preparation. Studies were performed on 23 opossums (Didelphis virginiana) of either sex weighing between 1.6 and 2.8 kg with a mean of 2.1 kg. The opossum was chosen as the animal model for a number of reasons. The high pressures in the anal canal in this animal species are primarily due to IAS smooth muscle (17). The detailed studies of myenteric (18) and submucosal (19) plexuses of the gut have been carried out in the opossum. The animals were fasted overnight but allowed free access to water. On the day of the experiment each animal was initially anesthetized with pentobarbital sodium (40 mg/kg intraperitoneally) and then restrained supine on an animal board equipped with a water circulating heat pad (model K-20, Gorman-Rupp Industries Div., Belleville, OH) to maintain the body temperature at 36°C. Either femoral vein was cannulated to provide central venous access for intravenous (i.v.) administration of different agents. The left femoral artery was cannulated and the catheter advanced ~ 10 cm (jut cephalad to the aortic bifurcation) for monitoring the blood pressure and localized intraarterial (i.a.) administration of various agents. The position of the i.a. cannula was confirmed at the end of the experiment. Subsequent anesthesia was provided with alpha-chloralose (70-200 mg i.v.) as needed.

The animals were intubated with a cuffed endotracheal tube (3.0 mm i.d.; 4.3 mm o.d.), and placed on an artificial ventilator (model 661, Harvard Apparatus Co., Millis, MA) at 20 strokes/min. The tidal volume was determined from the Harvard Apparatus ventilation nomogram based on body weight.

The animals were then turned to a prone position. A middorsal incision was made over the pelvisacral region, and the lateral paraspinal muscles were dissected and retracted to expose the sacroiliac joint and sacrum. As described previously (2), the overarching portion of the pars lateralis and the sacroiliac joint were resected to expose the sacral nerve roots as they emerged from the sacral foramina. Bipolar, silver-silver chloride hook stimulating electrodes were placed in contact with the fourth sacral nerve root to provide sacral nerve stimulation. Bipolar stainless steel needle electrodes (2 mm apart) were in-
serted into the muscularis of the distal anal canal, ~1 cm from the anal verge, to provide local intramural stimulation. A balloon constructed as part of the catheter assembly was placed in the rectum as described below. Inflation of the balloon was intended to produce rectal distension. Pancuronium bromide (Organon Inc., W. Orange, NJ) 1 mg/kg i.v. was administered to eliminate the electromyographic activity of the external anal sphincter (17).

**Intraluminal pressure measurements.** A specially designed, six-channel manometry catheter assembly continuously perfused with bubble-free water through a low compliance, pneumohydraulic valve system was used to record internal anal sphincter pressures. The catheter assembly consisted of six polyvinyl tubes (i.d. 0.7 mm, o.d. 1.3 mm; Insultab, Woburn, MA) arranged around a central polyvinyl tube (i.d. 1.6 mm, o.d. 2.4 mm) to which an inflatable balloon constructed from a finger cot was attached. The assembly provided a linear array of six side-hole openings situated at 0.6-cm intervals with an inflatable balloon attached 2 cm beyond the distal port. The total outside diameter of the assembly was 4.7 mm. The catheter was positioned so that the pressures from the distal rectum and the entire length of the anal canal could be recorded simultaneously. A pull-through was performed until the highest pressure zone of the IAS was identified. The rectal balloon could be inflated with varying volumes of air without disturbing the position of the catheter assembly. For the sake of uniformity, resting sphincter pressures from the top of the rhythmic fluctuations were recorded in the zone that gave the highest pressures. Pressures were recorded on a Beckman Dynograph recorder (model R711, Beckman Instruments Inc., Schiller Park, IL) using Statatham transducers. The pressure dynamics of the perfused catheters were such that abrupt occlusion of the catheter tip produced a pressure rise of 200 mmHg in 0.1 s. The baselines were set at atmospheric pressure. Details of the catheter assembly and the procedure for recording pressure have been described before (17).

**Stimulus and drug administration.** Baseline responses to isoproterenol hydrochloride (Winthrop Laboratories, New York; mol wt 211.24) 4.8-9.5 × 10^-9 mol/kg (1-2 μg/kg) administered i.a. were obtained. Control responses of the fall in internal anal sphincter pressures (IASP) by sacral nerve stimulation from 0.5 to 20 Hz (5 mA; 0.5-ms pulse duration; 2-s train) and local intramuscular stimulation with 20 and 40 Hz (5 mA; 0.5-ms pulse duration; 2-s train) were obtained using a stimulator (model S11A, Grass Instrument Co., Quincy, MA). Inhibitory responses to rectal balloon distension on IAS were obtained by inflating the rectal balloon with volumes of 2-15 ml of air for 8 s.

Dose–response curves were obtained after the i.a. administration of vasoactive intestinal polypeptide (VIP) (VIP porcine sequence; Sigma Chemical Co., St. Louis, MO; mol wt 3,326) in doses ranging from 1.1 × 10^-12 to 7.2 × 10^-9 mol/kg.

The influence of i.a. administration of the neurotoxin tetrodotoxin (TTX) (Calbiochem-Behring Corp., San Diego, CA; mol wt 319.28) on the VIP dose–response curve was then examined. The initial dose was 1.6 × 10^-8 mol/kg (5 μg/kg), and it was then repeated as necessary until the rectoanl reflex response to balloon distension was abolished. In all the experiments where TTX was used, the animals were provided with cardiovascular support with a continuous i.v. infusion of lactated Ringer’s at 9 ml/h. The above-mentioned doses of VIP were administered again once the rectal distension-induced IASP relaxation was abolished.

The effects of two VIP antagonists, VIP analogue, [4Cl-D-Phe^4,Leu^7] VIP (a generous gift from Dr. J. Rivier of the Peptide Biology Laboratory of the Salk Institute, San Diego, CA; mol wt 3,342.09) (15) and GFR analogue, (N-Ac-Tyr^3,D-Phe^2)-GRF (1-29)-NH_2 (Peninsula Laboratories, Inc., Belmont, CA; mol wt 3,476.3) (16) on IAS relaxation were investigated. Both antagonists were administered in incremental doses until the fall in IASP caused by near D_{90} of VIP was significantly antagonized. At that point a constant i.a. infusion of the antagonist was started. After VIP antagonism was achieved, IASP responses to isoproterenol, VIP, sacral nerve stimulation, rectal balloon distension, and local intramuscular stimulation were repeated.

In one series of experiments, after obtaining significant VIP antagonism with (N-Ac-Tyr^3,D-Phe^2)-GRF (1-29)-NH_2, and after administration of the above-mentioned stimuli, the dose of the antagonist was increased fourfold, and the stimuli were repeated. The purpose of this experiment was to determine if the GFR analogue acted as a competitive antagonist. Such experiments were not performed with [4Cl-D-Phe^4,Leu^7] VIP because of the limited amount of the antagonist available.

In another series of experiments baseline responses to VIP, 1.5 × 10^-10 mol/kg, isoproterenol hydrochloride, 2.9 × 10^-10 mol/kg, adenosine (Calbiochem-Behring Corp.; mol wt 267.2), 7.9 × 10^-7 mol/kg, and peptide histidine isoleucine (PHI) (Peninsula Laboratories, Inc.; mol wt 2,995.85), 6.0 × 10^-10 mol/kg administered i.a. were obtained, before and after the administration of the VIP antagonists.

**In vitro experiments.** In another series of experiments some in vitro studies were performed to examine the possibility of VIP as an inhibitory neurotransmitter in the neurally mediated relaxation of isolated smooth muscle strips of the IAS of the opossum. For these experiments, standard methodology for recording resting tension and its changes in response to electrical field stimulation (EFS) and VIP was employed (3). The changes in the IAS tension were expressed on a percentile basis. In this series of experiments, the control responses to EFS (50 V, 2 ms for 4-s train) at different frequencies (0.5-10 Hz) and different doses of VIP (1.0 × 10^-7 to 1.0 × 10^-6 M) on the resting tension of the IAS strips were examined. The responses to these stimuli were then repeated in the presence of [4Cl-D-Phe^4,Leu^7] VIP (30 μM) and (N-Ac-Tyr^3,D-Phe^2)-GRF (1-29)-NH_2 (10 μM). The doses of both antagonists were chosen on the bases of previous experiments by other investigators studies in vitro (15,16).

The i.a. catheters, syringes used for injection, and vials containing different peptide solutions and muscle baths were all rinsed with 2.5% bovine serum albumin before use to avoid binding of peptides to polyvinyl tubing, plastic, and glassware. All agents employed in the study were dissolved in physiologic saline. The i.a. or i.v. administration of different volumes of physiological saline produced no significant change either on the resting tone of the IAS or in response to any stimuli.

**Data analysis.** All results are expressed as mean ±SE of different observations. Changes in IASP and tensions are expressed on percentile as absolute bases. The statistical analysis was performed using one-tail t tests or multivariate analysis where appropriate (20).

**Results**

**Effect of VIP on the resting internal anal sphincter.** Intrarectal administration of VIP produced a dose-dependent fall in IASP (Fig. 1). The threshold dose (1.1 × 10^-12 mol/kg) of VIP produced 19.2±3.7% fall (from 61.2±2.9 to 48.5±2.0 mmHg) in IASP. The maximal effective dose (MED) was 1.8 × 10^-9 mol/kg, which produced a fall in IASP of 90.6±0.7% (from

![Figure 1](image_url)
the TTX antagonism of VIP response was 8.0±0.8 and 111.5±12.0 mmHg, respectively. These values were not significantly different (P > 0.05; n = 6 in three animals). The entire dose–response curve with VIP showing a fall in IASP obtained during control experiments was compared with the one obtained after TTX administration by polynomial comparison. No significant difference between the two curves was found (P > 0.05; Fig. 3).

Influence of VIP antagonist, [4CI-D-Phe⁶,Leu¹⁷] VIP on fall in IASP caused by VIP and other agonists. In the initial experiments the influence of different doses of [4CI-D-Phe⁶,Leu¹⁷] VIP (1.7 × 10⁻⁹ to 2.3 × 10⁻⁷ mol/kg) on the fall in IASP caused by VIP was examined. It was observed that this VIP analogue was required in the dose of 2.3 × 10⁻⁷ mol/kg i.a. to significantly antagonize the inhibitory effect of VIP on IAS. The antagonism of VIP responses on IAS by the single bolus of VIP analogue lasted for ~ 30 min. In the longer protocols, when needed, the duration of action of [4CI-D-Phe⁶,Leu¹⁷] VIP could be prolonged by a constant i.a. infusion (2.8 × 10⁻⁹ mol/kg/min) of the antagonist.

The dose–response curve obtained with VIP in control experiments was shifted significantly to the right after administration of the VIP analogue (P < 0.05; n = 8 in four animals at each dose level; Fig. 4). The fall in IASP after VIP (1.5 × 10⁻⁹ mol/kg) in control experiments was 42.8±5.2% (from 45.8±3.2 to 25.27±2.6 mmHg), and after [4CI-D-Phe⁶,Leu¹⁷] VIP it was 20.3±5.3% (from 48.21±3.6 to 37.8±3.4 mmHg) (P < 0.05; n = 8 in four animals). This represents an antagonism of 52%. The fall in IASP after the threshold dose of VIP (1.1 × 10⁻¹² mol/kg) was 19.2±3.3% (from 61.0±2.9 to 48.5±2.2 mmHg) and after [4CI-D-Phe⁶,Leu¹⁷] VIP, it was almost abolished to 1.9±1.6% (from 58.7±6.4 to 57.5±6.2 mmHg) (P < 0.05; n = 8 in four animals). This represents an antagonism of 90%. The fall in IASP of 89.5±1.1% (from 55.6±2.0 to 5.7±0.5 mmHg) after the MED of VIP (1.8 × 10⁻⁹ mol/kg) was significantly antagonized to 56.1±3.8% (from 63.5±3.3 to 27.5±2.6 mmHg) (P < 0.05; n = 8 in four animals). This represents an antagonism of 37%.

In a separate series of experiments, the dose of [4CI-D-Phe⁶,Leu¹⁷] VIP that antagonized the response of VIP (1.5 × 10⁻¹⁰ mol/kg) was found not to modify the inhibitory effects of isoproterenol (2.9 × 10⁻¹⁰ mol/kg), adenosine (7.9 × 10⁻⁷ mol/kg), and PHI (6.0 × 10⁻¹⁰ mol/kg). In control experiments the fall in IASP with VIP, isoproterenol, adenosine, and PHI was 56.2±11.2, 69.5±9.4, 90.7±1.6, and 56.1±14.9%, respectively. In the presence of VIP antagonist these values were 12.2±6.9 (P < 0.05; n = 4), 70.3±16.2, 93.2±0.4, and 67.2±16.0%, respectively (P > 0.05; n = 4).

The VIP antagonist, VIP analogue at a dose of 2.3 × 10⁻⁷ mol/kg (i.a.) by itself caused a transient fall in IASP of 33.8±6.2%. This fall in IASP began within 13.4±2.2 s and lasted for 49.8±13.0 s.

Figure 3. Influence of the neurotoxin TTX on the fall in the resting IASP caused by different doses of VIP. The figure shows the comparison of dose–response curves obtained with VIP before and after TTX in the same animals. The comparison of two curves revealed that TTX exerted no significant effect on the fall in IASP caused by VIP (P > 0.05; n = 6 in three animals, two observations in each animal at each dose level).

Figure 4. Influence of VIP antagonist [4CI-D-Phe⁶,Leu¹⁷] VIP on the fall in IASP caused by VIP. The antagonist caused a significant rightward shift in the entire dose–response curve of VIP. The calculated D₅₀ for VIP before and after the antagonist were 1.5 × 10⁻¹⁰ and 2 × 10⁻⁹ mol/kg (P < 0.05; n = 8 in four animals, two observations in each animal at each dose level).
Influence of [4CI-D-Phe\(^6\),Leu\(^7\)] VIP on the fall in IASP caused by balloon distension in the rectum. The fall in IASP caused by different volumes of rectal distension tested was significantly antagonized by the VIP antagonist ($P < 0.05$; $n = 8$ in four animals; Fig. 5). The fall in IASP in response to 10 ml rectal distension in control experiments was $71.2 \pm 8.0\%$ (from 42.6±3.3 to 13.0±3.6 mmHg) and after [4CI-D-Phe\(^6\),Leu\(^7\)] VIP, it was $36.6 \pm 12.3\%$ (from 46.6±1.6 to 30.9±6.3 mmHg) ($n = 8$ in four animals; $P < 0.05$; Fig. 5). This represents an antagonism of 49%. Fig. 6 illustrates an example of the inhibitory response of rectal balloon distension on the IAS before and after the VIP antagonist.

Influence of [4CI-D-Phe\(^6\),Leu\(^7\)] VIP on fall in IASP caused by sacral nerve stimulation. At all frequencies tested, the VIP antagonist caused significant antagonism of the fall in IASP induced by electrical stimulation of the sacral nerve. Electrical stimulation of the sacral nerve at 5 Hz caused a fall in IASP of 70.9±6.0% (from 48.4±3.2 to 14.8±3.2 mmHg) in control experiments. This response was significantly antagonized after [4CI-D-Phe\(^6\),Leu\(^7\)] VIP to 36.7±8.8% (from 40.7±2.8 to 26.8±4.7 mmHg) ($P < 0.05$; $n = 12$ in four animals, three observations in each animal at each frequency; Fig. 7), as illustrated in Fig. 6. This represents an antagonism of 48%.

Influence of VIP analogue [4CI-D-Phe\(^6\),Leu\(^7\)] VIP on fall in IASP caused by local intramural stimulation. The fall in IASP caused by local intramural stimulation at 20 and 40 Hz was significantly antagonized by the antagonist. The fall in IASP with 20 and 40 Hz of local intramural stimulation was 57.3±5.0% (from 45.3±2.6 to 19.8±3.1 mmHg) and 60.6±7.7% (from 44.5±2.9 to 19.1±4.8 mmHg), respectively, in control experiments. After [4CI-D-Phe\(^6\),Leu\(^7\)] VIP, these responses were significantly antagonized to 31.6±7.4% (from 43.0±2.2 to 29.8±3.7 mmHg) after 20 Hz, and to 36.9±9.6% (from 41.8±3.7 to 27±5.4 mmHg) after 40 Hz ($P < 0.05$; $n = 12$ in four animals, three observations in each animal at each frequency; Fig. 8), as illustrated in Fig. 6. This represents an antagonism of 45 and 39% respectively.

Influence of GRF analogue (N-Ac-Tyr\(^1\),D-Phe\(^2\))-GRF (1-29)-NH\(_2\) on fall in IASP caused by VIP and other agonists. Examination of different doses ($2.9 \times 10^{-9}$ to $2.3 \times 10^{-8}$ mol/kg) (i.a.) of GRF analogue suggested that $2.3 \times 10^{-8}$ mol/kg caused a significant and consistent antagonism of VIP-induced fall in IASP (Table I). The amount of antagonism of VIP-induced fall in IASP by (N-Ac-Tyr\(^1\),D-Phe\(^2\))-GRF (1-29)-NH\(_2\) was dependent on the dose of VIP used (Table I). GRF analogue at a dose that caused significant antagonism of the inhibitory effect of VIP on the IAS had no significant effect on the fall in IASP caused by different agonists.

In a separate series of experiments the fall in IASP in response to VIP ($1.5 \times 10^{-10}$ mol/kg), isoproterenol ($2.9 \times 10^{-10}$ mol/kg) and...
mol/kg), adenosine (7.9 × 10⁻⁷ mol/kg), and PHI (6.0 × 10⁻¹⁰ mol/kg) was examined before and after (N-Ac-Tyr¹,D-Phe²)-GRF (1-29)-NH₂. The fall in IASP with these agonists before the antagonist was 68.5±9.7, 74.8±8.8, 90.2±5.6, and 63.9±13.0%, respectively. In the presence of the antagonist these values were 34.9±4.9, 78.4±7.9, 85.5±2.3, and 62.5±7.4%, respectively. The fall in IASP with VIP in the presence of the antagonist was significantly different (P < 0.05; n = 4) while effects of other agonists were not significantly modified (P > 0.05; n = 4).

The antagonism of VIP in causing a fall in IASP with a single dose of 2.3 × 10⁻⁸ mol/kg (i.a.) of (N-Ac-Tyr¹,D-Phe²)-GRF (1-29)-NH₂ lasted ~ 40 min. However, this duration could be prolonged with the continuous infusion of GRF analogue (5.7 × 10⁻¹⁰ mol/kg per min). Administration of GRF analogue (N-Ac-Tyr¹,D-Phe²)-GRF (1-29)-NH₂ (2.3 × 10⁻⁸ mol/kg) (i.a.) by itself caused a transient fall in IASP of 38.4±10.8%. This fall in IASP with GRF analogue occurred within 14.4±3.3 s and lasted for 74.5±23 s.

### Table I. Influence of GRF analogue, (N-Ac-Tyr¹,D-Phe²)-GRF (1-29)-NH₂ (2.3 × 10⁻⁸ mol/kg) on the Fall in IASP in Response to Exogenous VIP

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Values expressed as mean±SE of n = 5 in five animals.

* P < 0.05; the difference between the values obtained during control and GRF analogue are significantly different.

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Figure 8. Influence of [4Cl-D-Phe⁶,Leu¹⁷]VIP on percent fall in IASP by local intramural stimulation. The fall in IASP caused local stimulation examined at 20 and 40 Hz (5 mA; 5 ms pulse duration; 2 s train) was significantly antagonized by VIP antagonist (2.3 × 10⁻⁷ mol/kg, i.a.) (P < 0.05; n = 12 in four animals, three observations in each animal at different frequencies).

Figure 9. Influence of (N-Ac-Tyr¹,D-Phe²)-GRF (1-29)-NH₂ on percent fall in IASP caused by rectal balloon distension, sacral nerve stimulation, and local intramural stimulation. Note that the VIP antagonist caused significant antagonism of fall in IASP caused by different neural stimuli at the different intensities of stimulation tested (P < 0.05; n = 15 in five animals, three observations in each animal at each level of stimulation).
(Fig. 9) \( P < 0.05; n = 15 \) in five animals). The fall in IASP after 20 and 40 Hz was 64.5±5.9% (from 36.7±2.9 to 10.9±1.0 mmHg) and 77.4±4.4% (from 42.9±1.7 to 8.6±0.8 mmHg), respectively. After the GRF analogue, it was antagonized to 38.9±5.5% (from 36.5±2.4 to 23.1±3.3 mmHg) and 50.3±6.0% (from 38.8±1.7 to 20.4±3.2 mmHg), respectively. This represents an antagonism of 40% and 35%, respectively.

**Influence of two different doses of GRF analogue, (N-Ac-Tyr<sup>1</sup>,D-Phe<sup>2</sup>)-GRF (1-29)-NH<sub>2</sub> on fall in the IASP caused by VIP.** In order to see whether the antagonism of fall in IASP caused by different stimuli with GRF analogue was dose-dependent, we compared the influence of two doses, \( 2.3 \times 10^{-8} \) and \( 9.2 \times 10^{-8} \) mol/kg (i.a.), respectively, of GRF analogue on different stimuli. We first examined the effect of different doses of VIP in causing a fall in IASP before and after two doses of GRF analogue. GRF analogue caused dose-dependent antagonism of the fall in IASP by VIP (Fig. 10). The dose of VIP (\( 1.5 \times 10^{-10} \) mol/kg) which caused a fall in IASP of 45.3±2.3% (from 33.2±1.8 to 16±1.6 mmHg) was antagonized to 27.6±5.4% (from 35.0±1.7 to 25.5±3.1 mmHg) and 18.3±1.1% (from 29.2±5.8 to 23.7±4.4 mmHg) by \( 2.3 \times 10^{-8} \) and \( 9.2 \times 10^{-8} \) mol/kg of GRF analogue, respectively. Examination of the three dose-response curves with VIP revealed that both doses of GRF analogue caused incremental and significant rightward shifts in the control curve.

**Influence of two different doses of GRF analogue, (N-Ac-Tyr<sup>1</sup>,D-Phe<sup>2</sup>)-GRF (1-29)-NH<sub>2</sub> on fall in IASP produced by rectal distension.** The fall in IASP after rectal balloon distension was further antagonized by increasing doses of GRF analogue (Fig. 11). After 5 ml the fall of IASP in the control was 81.0±2.1% (from 32.6±1.3 to 6.1±0.5 mmHg); this was antagonized to 66.2±8.4% (from 44.1±5.9 to 14.1±3.4 mmHg) and 54.1±2.9% (from 27.5±1.7 to 12.5±1.0 mmHg) after 2.3 \( \times 10^{-8} \) and 9.2 \( \times 10^{-8} \) mol/kg of GRF analogue, respectively (\( P < 0.05 \)). As can be seen in Fig. 11 the increasing antagonism to the fall in IASP after balloon distension with higher doses of GRF analogue was more evident at the lower volumes of rectal balloon distension.

**Influence of two different doses of GRF analogue on the fall in IASP produced by sacral nerve stimulation.** Increasing doses of GRF analogue antagonized further the IASP response to sacral nerve stimulation (Fig. 11). As with balloon distension, this was more evident at the lower frequencies of electrical stimulation. At 5 Hz the IASP fall in controls was 92.3±0.3% (from 32.8±1.4 to 2.5±0.2 mmHg) and was antagonized to 65.9±6.5% (from 44.3±1.9 to 15.1±3 mmHg) and 57.7±3.9% (from 23.5±0.6 to 10.0±1.1 mmHg) after 2.3 \( \times 10^{-6} \) and 9.2 \( \times 10^{-6} \) mol/kg GRF analogue, respectively (\( P < 0.05 \)).

**Influence of two different doses of GRF analogue, (N-Ac-Tyr<sup>1</sup>,D-Phe<sup>2</sup>)-GRF (1-29)-NH<sub>2</sub> on the fall in IASP produced by local stimulation.** As can be seen in Fig. 11, increasing doses of GRF analogue further antagonized the response of IASP to local stimulation. The IASP fall after 20 Hz was 80.2±0.9 (from 42.1±1.8 to 8.3±0.6 mmHg) in controls and was antagonized to 35.8±12.0% (42.5±1.1 to 27.5±5.4 mmHg) and 26.6±2.1% (from 25.8±4.1 to 19.1±3.6 mmHg) after 2.3 \( \times 10^{-8} \) and 9.2 \( \times 10^{-8} \) mol/kg GRF analogue, respectively (\( P < 0.05 \)).

**Influence of VIP antagonists on the fall in resting tension of IAS smooth muscle strips in response to VIP and EFS.** EFS and VIP caused a fall in IAS tension in in vitro strips. The inhibitory responses to EFS were frequency dependent, and those to VIP were dose dependent (Figs. 12 and 13). Both [4Cl-D-Phe<sup>6</sup>,Leu<sup>10</sup>]VIP (30 \( \mu \)M) and (N-Ac-Tyr<sup>1</sup>,D-Phe<sup>2</sup>)-GRF (1-29)-NH<sub>2</sub> (10 \( \mu \)M) caused a significant rightward shift in the VIP dose-response and EFS frequency-response curves (Figs. 12 and 13; \( P < 0.05 \)).

**Discussion**

The present studies suggest that VIP acts as an inhibitory neurotransmitter of intramural neurons in IAS relaxation in re-
response to the rectoanal reflex in the opossum. This conclusion is supported by the following findings. VIP is a potent relaxant of IAS smooth muscle. The relaxant effect of VIP is exerted by its action directly at the smooth muscle since the inhibitory response of VIP on the IAS was TTX resistant. The direct inhibitory action of VIP on the smooth muscle strips has also been shown before (1, 3, 21). However, further support for the conclusion that VIP may act as the inhibitory neurotransmitter in the IAS comes from the present investigation which used selective VIP antagonists: [4CI-D-Phe6,Leu17] VIP (15) and (N-Ac-Tyr1,D-Phe2)-GRF (1-29)-NH2 (16). The VIP analogue has been shown to selectively inhibit VIP-stimulated amylase secretion and cAMP elevation in in vitro preparations of guinea pig exocrine pancreas (15). The GRF analogue has been shown to antagonize adenylate cyclase stimulation in response to VIP receptor activation in the rat pancreatic plasma membranes (16). In both systems these antagonists were shown to behave as selective and competitive antagonists to VIP receptors. In our experiments, VIP antagonists in doses that produced significant antagonism of fall in IASP by VIP caused significant impairment in IAS relaxation in response to rectal balloon distension (which mimics rectoanal inhibitory reflex), thus providing additional support for our hypothesis.

These studies also suggest that IAS relaxation in response to rectoanal reflex is mediated through the release of VIP from intramural myenteric inhibitory neurons. Such inhibitory neurons appear to lie in the sacral inhibitory pathway. The support for this hypothesis comes from the following findings. VIP antagonists which caused antagonism of rectal balloon distension-mediated IAS relaxation also antagonized the fall in IASP in response to preganglionic sacral nerve stimulation and local intramural stimulation. We have previously shown that the fall in IASP by local intramural stimulation is due to activation of the intramural inhibitory neurons (2). The findings are further substantiated by our in vitro experiments in the IAS strips. In such studies, the antagonism of the fall in IAS tension caused by EFS and VIP in the presence of VIP antagonists suggests the release of VIP from the intramural inhibitory neurons. Interestingly, previous studies have suggested the presence of VIP immunoreactivity in intramural myenteric neurons of the IAS region (22, 23).

The conclusion of the present investigation is in agreement with that of the LES studies (11, 14) where VIP has been suggested to act as an inhibitory neurotransmitter. Furthermore, a number of studies have shown the presence of VIP-containing nerve fibers in different regions of the gut (24, 25) and the release of VIP from the venous effluent draining the rectal region in response to pelvic nerve stimulation (9, 10, 13) and rectal distension (9) in cats and pigs.

Our studies further suggest that VIP may not be the only neurotransmitter involved in the IAS relaxation by the rectoanal reflex since the VIP antagonists in the doses that almost abolished the fall in IASP caused by lower doses of VIP failed to abolish the IAS relaxation in response to lower levels of neural stimuli. It is possible that another inhibitory neurotransmitter is released in addition to VIP and is partially responsible for IAS relaxation. In that case, it would be important to further examine the role of such a substance in IAS relaxation. Purinergic substances have been shown to act as inhibitory neurotransmitters in a variety of tissues (6). Our recent studies however rule out the participation of purinergic agents as the inhibitory neurotransmitters in IAS relaxation (Rattan, S., and R. Shah, manuscript in preparation).

The present studies also show that [4CI-D-Phe6,Leu17] VIP and (N-Ac-Tyr1,D-Phe2)-GRF (1-29)-NH2 can be used successfully as VIP antagonists in in vivo as well as in vitro studies. Whether these antagonists act as competitive antagonists in vivo experiments is not clearly known. However, a number of factors point to that effect. First, the antagonists caused a parallel shift in the VIP dose–response curve showing fall in IASP. Secondly, the maximal fall in IASP by VIP in control experiments was still obtained in the presence of VIP antagonists although the dose of VIP required to do so was increased many fold. Thirdly, when higher doses of the VIP antagonist (N-Ac-Tyr1,D-Phe2)-GRF (1-29)-NH2 were used, they produced incremental antagonism of the fall in IASP by VIP.
Furthermore, the antagonists were selective in antagonizing the inhibitory responses of VIP on the IAS without modifying the responses to isoprotenerol, adenosine, and PHI.

The findings of the present investigation suggest that VIP plays a significant role as an inhibitory neurotransmitter in rectoanal reflex-mediated IAS relaxation. However, further studies seeking electrophysiologic and biochemical correlates for the inhibitory responses of rectoanal reflex, sacral nerve stimulation, local intramural stimulation, and VIP need to be performed. Information relating to the inhibitory neurotransmitter involved in IAS relaxation would lead to a better understanding of the pathophysiology of anorectal incontinence and constipation, as they relate to the reflex control mechanisms of the anorectal region.

Acknowledgments
The authors thank Drs. R. Goyal, Chafiq Moumoumi, A. Sengupta, and R. Shah for their helpful criticism, and Dr. B. Ransil for his help with the statistical analysis and computer resources. The authors also thank Mr. J. Kaczmarek for his technical assistance, and M. L. McGee and Ms. P. Pralh for typing the manuscript.

This work was supported by U.S. Public Health Service grant DK-35385 and DK-31092 and Electrophysiology Core Grant DK-34854 from the National Institutes of Health.

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