Effect of modulation of serotonergic, cholinergic, and nitrergic pathways on murine fundic size and compliance measured by ultrasonomicrometry

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The three main motor functions of the human stomach are to accommodate food ingested at mealtimes, triturate food, and mix the content with digestive enzymes and acid. Gastric accommodation includes receptive relaxation and adaptive relaxation. Receptive relaxation allows the stomach to initially accommodate the volume of a meal without a rise in intragastric pressure (IGP) (9, 21), whereas adaptive relaxation is a slower response that may be modulated by the physical and chemical properties of the meal ingested (13, 25) and the neurohormonal responses to the meal.

Impaired fundic accommodation can occur as a result of vagal injury or a vagotomy and is also present in about a third of patients with functional dyspepsia (3, 34, 36, 37). Fundic accommodation requires an intact extrinsic innervation, predominantly but not exclusively, through vagovagal reflexes and a complex interaction among enteric nerves, gastric mucosa, muscularis propria, smooth muscle cells, and interstitial cells of Cajal (4, 35, 39). In several species, including humans, the neurotransmitters involved include nitric oxide (NO) and serotonin (5-HT) (18, 19, 32, 40). An intact nitrergic pathway is also required for fundic relaxation in the isolated murine fundus (11, 14, 28, 31, 33). However, the mouse fundic data were obtained in muscle strips. The 5-HT receptor agonists buspirone and sumatriptan have been used in clinical studies (7, 16) to treat functional dyspepsia by targeting fundic relaxation. Buspirone is a 5-HT1A agonist and sumatriptan is a 5-HT1D agonist, although it is also clear that both act on more than one 5-HT receptor subtype (24). The relative contributions of central and peripheral mechanisms of action of both drugs have not been well defined. Research on the neuronal and nonneuronal pathways that lead to fundic accommodation is hampered by the lack of an accurate, reliable, and reproducible method to study fundic accommodation in small animals. In vivo experiments are complicated by the lack of an accurate method to assess gastric accommodation in larger animals, including humans, but their utility is severely limited in smaller animals such as mice. A recent study (30) developed a miniaturized method to assess gastric tone and compliance using a barostat. The ready availability of knockin and knockout mice makes the mouse model an attractive model to dissect out the pathways that contribute to fundic accommodation. Testing and validation of a method to accurately measure in vivo changes in murine fundic size and determination of the effect of cholinergic, nitrergic, and serotonergic pathways on fundic size in the intact innervated murine stomach are therefore of considerable interest.

In the present study, we utilized an ultrasound technique, known as ultrasonomicrometry, to determine changes in distance between piezoelectric crystals attached to the serosal surface of the murine fundus. The methodology was used to determine the effect of modulation of the major pathways that control fundic size in larger intact animals.

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Ultrasonomicrometry has previously been successfully used to determine changes in volume in the murine heart (22, 23) and the rat gastric body (1, 2). As the piezoelectric crystals are small, several can be placed on the serosa of the murine stomach, allowing the measurement of ultrasound waves as they pass through the thin wall of the murine fundus without needing to traverse the air-filled lumen. This avoids the current major limitation of conventional transabdominal ultrasonography. Our results show that ultrasonomicrometry can be successfully used to measure fundic size in vivo and that the measured changes accurately reflect changes in volume. These data suggest that the technique may be of use in small animal in vivo research on fundic and gastric volumes.

METHODS

Animals

All experiments were approved by the Mayo Institutional Animal Care and Use Committee. Adult (6–8 wk) male 129 SvEv mice (Taconic; Gemantown, NY; 20–25 g body wt) were used in all experiments. Their diet was changed to Ensure Plus (Abbott Laboratories; Columbus, OH) with free access to water 48 h before each experiment to make sure the stomach was empty of solid food at the time of the experiment. Mice were anesthetized with ketamine (100 mg/kg im) and xylazine (10 mg/kg im). Every 40 min, half of the initial dosage was readministered. Animals were placed on a custom-built heating pad set at 38°C to control body temperature, and an initial dosage was readministered. Animals were placed on a custom-built heating pad set at 38°C to control body temperature, and an abdominal midline incision was made. A silicon catheter (outer diameter: 0.9 mm, inner diameter: 0.6 mm) was inserted through a small incision (~1–2 mm) made in the proximal jejunum about 3–5 cm distal to the pylorus, and the residual stomach content was flushed out with saline. After the stomach was emptied, the tip of the catheter was placed in an area between the fundus and corpus. The catheter was ligated in place with a silk suture that also closed the jejunal incision. The other end of the catheter was connected to a saline catheter. Pressure signals were recorded into the mouse stomach along with the catheter. The pressure transducer was calibrated before each experiment using a water column (Fig. 1).

Experimental Protocols

After stable traces from the piezoelectric crystals were established, drugs were injected via the tail vein or intramuscularly. When more than one drug was used, a minimum of 20 min separated administration of each drug.

For the ex vivo experiments, the mouse stomach (n = 5) was removed and placed in normal Krebs solution (at 38°C) bubbled with 3% CO2 and 97% O2. A pair of crystals was glued to the fundus (4–7 mm apart). The distal esophagus was ligated with a silk suture, and the proximal duodenum was connected to a 1-ml syringe. Boluses of saline (100 μl) were injected into the stomach in a stepwise fashion.
The changes in distance between the crystals was recorded digitally and plotted against the injected volume. In the in vivo experiments, the mice (n = 5) were anesthetized, the stomach exposed, and the crystals were placed on the fundus in situ. A similar protocol to the one described above was then used.

To measure compliance, IGP was adjusted at the beginning of an experiment to 60 mmH2O (≈4.41 mmHg) by infusing prewarmed saline solution into the stomach; 60 mmH2O was chosen as the initial IGP because, in agreement with the literature (30), at this pressure both contractions and relaxations could be most easily observed. After a 10- to 15-min equilibration period, the IGP was increased in a stepwise fashion by 10 mmH2O (0.73 mmHg) to a maximum of 100 mmH2O (Fig. 2). At least 6 min separated each increase in IGP. The 6-min time period was chosen based on preliminary experiments that showed that steady state was reached in this time period. Changes in distances between crystals (dI) and IGP were continuously recorded, and the intercrystal distance (ICD) and the change in IGP (10 mmH2O) were used to calculate compliance (dI/dP) and plotted (Fig. 2B,a).

The gastric fundus is not only compliant but also actively relaxes to accommodate food. Disorders in fundic function may therefore not only include abnormal compliance but also impaired, delayed, or slowed receptive relaxation. To quantify the rate of relaxation, we also measured the rate of the change in distance between the crystals, i.e., the fundic distension rate (FDR), in response to a given initial pressure. The FDR was calculated from the change in length over time (dI/dt). Values of dI/dt were determined by calculating the fit of the slope of the trace (25–75%) in response to any given pressure change (fig. 2B,b).

Drugs

Atropine sulphate, bethanechol chloride, and Nω-nitro-L-arginine (L-NNA) were purchased from Sigma (St. Louis, MO). Nitroglycerin was purchased from American Regent Laboratory. Buspirone hydrochloride was purchased from Tocris (Ellisville, MO), and sumatriptan succinate was purchased from GlaxoWellcome (Research Triangle Park, NC).

All drugs were given intravenously via the tail vein except for atropine, which was given intramuscularly. The doses used were 0.2–0.4 mg/kg atropine, 0.15 mg/kg bethanechol, 0.1–0.3 mg/kg sumatriptan, 0.1–0.3 mg/kg buspirone, 0.03 mg/kg nitroglycerin, and 0.5–1.5 mg/kg t-NNA. Doses were selected based on our previous work and on published data (5, 6, 10, 15, 17, 20, 26, 42).

Statistical Analysis

All results are reported as means ± SE. The number of individual experiments is indicated by the n value. Statistical significance was determined using paired Student’s t-tests for changes in ICD in response to a drug. An unpaired t-test was used for the compliance and FDR data. A P value of <0.05 was considered significant.

RESULTS

Validation Studies

A series of validation studies was carried out to determine whether the recorded values for the change in distance between the crystals correlated to the actual changes in the distance between the crystals. In the first set of experiments, the upper part of a finger of a rubber glove was cut off and attached to the tip of a 12-ml syringe. Two crystals were glued on the glove tip 10 mm apart using VetBond. The setup was placed under a dissecting microscope equipped with a micrometer. The syringe was used to inject water into the glove tip, and the distance between the two crystals at different injected volumes was directly measured and also measured using the ultrasonomicrometry system (n = 3). As can be seen in Fig. 3, there was a 1:1 correlation between the two measurements, suggesting that the ultrasonomicrometry measurements accurately reflected changes in distance between the two crystals.

A second validation study was carried out to determine the correlation between the changes in distance between two crystals placed on the curved surface of the mouse fundus and the volume of the stomach. This set of experiments was carried out both ex vivo on the excised stomach and in vivo as outlined in Methods. As can be seen in Fig. 3B, increases in gastric volume...
within the physiological range of the murine stomach also resulted in proportional changes in linear distance between two crystals \( (n = 5 \text{ for each experiment}) \).

**Pharmacological modulation.** In a third set of experiments, to determine the biological responsiveness of the technique, bethanechol was used to contract the fundus, and nitroglycerin, a NO donor, was used to relax the fundus (Fig. 4). These experiments were carried out in vivo. Bethanechol (0.15 mg/kg iv) caused a rapid decrease in the distance between the crystals, indicating fundic contraction (9.7 ± 2.9%, \( n = 3, P < 0.05 \); Fig. 4). Nitroglycerin (0.03 mg/kg iv) caused an increase in the distance between the crystals, indicating relaxation (4.6 ± 1.8%, \( n = 5, P < 0.05 \); Fig. 4). Atropine (0.2 mg/kg im) also caused an increase in the distance between the crystals (5.5 ± 0.9%, \( n = 8, P < 0.05 \); Fig. 4), suggesting that there was endogenous cholinergic input to baseline fundic tone.

It is possible that a change in fundic volume may reflect a passive response to a contraction or relaxation in other regions of the stomach. A strong antral contraction may displace enough gastric content proximally to distend the fundus. To investigate this possibility, an additional two crystals were placed on the antral surface. Changes in fundic ICD were not accompanied by inverse changes in the antrum, suggesting that the results represented a direct effect of the drugs on the fundus (data not shown).

**Reproducibility.** We next determined the reproducibility of our measurements. In this set of experiments, IGP was increased in a stepwise fashion from 60 to 100 mmH\(_2\)O as previously described, and distance between the fundic crystals was measured (Fig. 5). The IGP was then returned to baseline, and the experiment was repeated. As can be seen in Fig. 5, there was close to 1:1 correlation both between the rate of change and absolute change in distances between the two sets of data.

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**Fig. 3.** In vitro, ex vivo, and in vivo validation of the accuracy of ultrasonomicrometry. A: correlation between distance between two crystals attached to a glove tip, measured using a micrometer, to changes in distance measured using the ultrasonomicrometry system \( (n = 3) \). A 1:1 ratio was obtained, suggesting that ultrasonomicrometry accurately measured distance. B,a: correlation between distance between two crystals placed across the dome of the fundus and volume changes in an isolated stomach \( (n = 5) \). B,b: similar experiment but in vivo with the crystals placed on the stomach after laparotomy \( (n = 4) \). Known volumes of saline were added to the isolated mouse stomach to induce volume changes. There was good correlation between the two measurements in both experiments.

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**Fig. 4.** Representative traces of changes in ICD during the administration of bethanechol (**A**), atropine (**B**), and nitroglycerin (**C**). The initial IGP was set to 60 mmH\(_2\)O. An upward deflection reflects an increase in ICD (increased fundic size), and a downward deflection reflects a decrease in ICD. Bethanechol decreased the ICD, suggesting that it decreased fundic size, whereas nitroglycerin administration increased the ICD, suggesting that it increased fundic size. Atropine also increased the ICD, suggesting a baseline cholinergic regulation of fundic tone.
In Vivo Measurement of Changes in Baseline Fundic Volume

As described in METHODS, after the midline incision and placement of the crystals, the anesthetized mice were left to recover for about 30 min when the traces from all the channels reached a stable level. In about half of all animals assessed, regular oscillations in ICD, reflecting a change in fundic size, accompanied by changes in IGP (Fig. 6A), were recorded. These oscillations initially suggested spontaneous fundic contractions. Changes in ICD were about 25–33% of those recorded from the antrum (data not shown). However, peaks in IGP preceded each increase in the distance between the crystals (Fig. 6B). The peaks in apparent fundic contraction (smallest distance between the crystals) coincided with the lowest IGP recordings, suggesting that the observed “contractions” superimposed on the slower changes in fundic size reflected contractile changes in the distal stomach and not spontaneous fundic contractions.

In Vivo Measurement of Compliance

Fundic compliance was first determined in controls. Fundic compliance was $9.1 \pm 0.69 \mu \text{mm H}_2\text{O}$ ($n = 26$ preparations). As drugs were administered intravenously in $15–30$ $\mu \text{l}$ of saline, we tested the effect of $25$ $\mu \text{l}$ NaCl 0.9% (iv) on fundic compliance. No effect was noted (Table 1). We tested the effects of atropine, bethanecol, and L-NNA on fundic compliance to determine the effect of cholinergic and nitrergic input on compliance. Atropine (0.4 mg/kg im) immediately increased the size of the fundus but did not alter compliance (Table 1). Bethanecol (0.15 mg/kg iv) and L-NNA (1.5 mg/kg iv) significantly reduced fundic compliance (Table 1).

In Vivo Measurement of FDR

The rate of fundic distension was calculated from the slope of each trace at each IGP. Bethanechol and atropine were used to assess the influence of cholinergic pathways, and L-NNA was used to assess the influence of nitrergic pathways on the fundus. The results before and after administration of each drug are summarized in Table 2. As previously shown, atropine immediately increased ICD but did not alter the rate of fundic distension. Bethanecol and L-NNA reduced the rate of fundic distension.

Nitrergic Relaxation of the Fundus

To determine the role of the nitrergic pathway in fundic relaxation in the intact mouse, we examined the effect of L-NNA on the fundus using somomicrometry in vivo. A decrease in ICD, suggesting contraction of the fundus, was only seen in mice in which there was no initial adjustment of the IGP to $60$ $\text{mm H}_2\text{O}$ (Fig. 7A). In these experiments, no saline

Fig. 5. Reproducibility of compliance measurements obtained by stepwise increases in IGP. A: changes in ICD induced by stepwise changes in IGP. After 30 min, the protocol was repeated, and the slope of both experiments was plotted (B). The slopes were nearly identical, suggesting excellent reproducibility. Numbers are in $\text{mm H}_2\text{O}$.

Fig. 6. Oscillations in fundic ICD associated with changes in IGP. A: spontaneous rhythmic changes in ICD associated with changes in IGP. B: an expanded trace to highlight the temporal relationship between the apparent fundic contractions and IGP. Peak IGP (dashed line) occurred before peak changes in ICD and was lowest when ICD was lowest, suggesting that the observed rhythmic activity observed in the fundus, superimposed on slower changes, reflected a passive response to contractions in the distal stomach.
was infused into the stomach, and IGP was left at baseline. Surprisingly, when the initial IGP was set as 60 mmH₂O, administration of l-NNA (1.5 mg/kg iv) increased ICD, suggesting relaxation of the fundus (2.8 ± 0.4%, n = 3, P < 0.05). When distal gastric size was monitored with crystals placed on the antrum, an antral contraction was always seen with l-NNA, and, when IGP was measured, IGP increased on administration of l-NNA (Fig. 7B). It is therefore likely that the paradoxical increase in fundic size on delivery of l-NNA was due to a passive distension induced by movement of fluid from the distal stomach to the proximal stomach and that the direct effect of l-NNA on the fundus was a decrease in fundic size.

As reported above, the ICD in the fundus was initially shown in Fig. 9. Although sumatriptan decreased ICD, suggesting it contracted the fundus, both d<sub>t</sub> and d<sub>dp</sub> and d<sub>lt</sub> did not subsequently change compliance (d<sub>t</sub>/dp: 9.9 ± 2.8 vs. 8.4 ± 8.4 μm/mmH₂O, n = 4 P > 0.05), nor was there any significant change in the rate of fundic distension to a given pressure (d<sub>t</sub>/dp and d<sub>lt</sub>/dr; see Tables 1 and 2 for details).

**DISCUSSION**

The main aims of the present study were to determine whether ultrasonomicrometry can be used to measure changes in mouse fundic size and to determine the effect of modulation of cholinergic, nitrergic, and serotoninergic pathways on fundic size and compliance in the intact innervated murine stomach. Our results show that ultrasonomicrometry accurately measures changes in fundic size and is able to capture changes in size induced by pharmacological interventions. Our results also show that there is a baseline cholinergic tone in the murine fundus and that nitrergic and serotoninergic pathways affect fundic size and compliance.

Ultrasonomicrometry has advantages compared with current methodologies used to determine changes in fundic size. The technique can be applied to small animals as the spatial resolution is very good and the individual piezoelectric crystals are small. The spatial resolution in ultrasonomicrometry is determined by the speed of the sound energy and the time intervals at which the transit of the signal is measured. The transit speed of sound in most biological material is 1.54 mm/μs. The equipment used recorded data every 15 ns, giving a spatial resolution of about 24 μm. This resolution allows

### Table 1. Effect of drugs on fundic compliance

<table>
<thead>
<tr>
<th>Drug</th>
<th>Control</th>
<th>Experiments</th>
<th>Saline (25 μl iv)</th>
<th>Sumatriptan (0.1 mg/kg iv)</th>
<th>L-NNA (1.5 mg/kg iv)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sumatriptan treatment</td>
<td>9.2 ± 2.6 (3)</td>
<td>9.9 ± 2.8 (4)</td>
<td>9.1 ± 2.5 (4)</td>
<td>8.7 ± 1.8 (3)</td>
<td>9.3 ± 1.1 (3)*</td>
</tr>
<tr>
<td>Buspirone treatment</td>
<td>8.9 ± 1.8 (3)</td>
<td>10.4 ± 1.8 (9)*</td>
<td>8.4 ± 8.4 (4)</td>
<td>7.7 ± 1.4 (4)</td>
<td>4.8 ± 0.8 (3)*</td>
</tr>
<tr>
<td>Atrpine treatment</td>
<td>5.3 ± 4.4 (4)</td>
<td>13.0 ± 11.4 (4)</td>
<td>21.1 ± 15.7 (4)</td>
<td>29.8 ± 19.1 (4)</td>
<td>21.8 ± 13.7 (4)</td>
</tr>
<tr>
<td>Bethanechol treatment</td>
<td>8.4 ± 3.0 (3)</td>
<td>13.2 ± 2.6 (3)</td>
<td>16.2 ± 1.0 (3)</td>
<td>16.1 ± 1.8 (3)</td>
<td>7.2 ± 1.8 (3)†</td>
</tr>
<tr>
<td>l-NNA treatment</td>
<td>15.5 ± 3.1 (5)</td>
<td>24.5 ± 3.4 (5)</td>
<td>38.1 ± 4.8 (5)</td>
<td>43.8 ± 5.1 (5)</td>
<td>43.8 ± 5.1 (5)</td>
</tr>
</tbody>
</table>

Values are means ± SE (in μm/mmH₂O); numbers in parentheses indicate number of animals. l-NNA, N<sup>-</sup>nitro-l-arginine. *P < 0.05 compared with control.

### Table 2. Effects of drugs on fundic distension rate

<table>
<thead>
<tr>
<th>IGP, mmH₂O</th>
<th>10</th>
<th>20</th>
<th>30</th>
<th>40</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sumatriptan treatment</td>
<td>12.4 ± 4.9 (9)</td>
<td>11.6 ± 3.9 (9)</td>
<td>26.8 ± 8.9 (9)</td>
<td>37.5 ± 10.5 (9)</td>
</tr>
<tr>
<td>Buspirone treatment</td>
<td>7.1 ± 1.2 (9)</td>
<td>24.6 ± 5.8 (9)</td>
<td>49.3 ± 12.8 (9)*</td>
<td>59.2 ± 12.9 (9)†</td>
</tr>
<tr>
<td>Atrpine treatment</td>
<td>9.9 ± 5.9 (4)</td>
<td>15.8 ± 7.6 (4)</td>
<td>21.3 ± 9.1 (4)</td>
<td>34.2 ± 13.7 (4)</td>
</tr>
<tr>
<td>Bethanechol treatment</td>
<td>12.6 ± 8.0 (4)</td>
<td>24.1 ± 20 (4)</td>
<td>30.8 ± 21.8 (4)</td>
<td>41.1 ± 17.3 (4)</td>
</tr>
<tr>
<td>l-NNA treatment</td>
<td>5.3 ± 4.4 (4)</td>
<td>13.0 ± 11.4 (4)</td>
<td>21.1 ± 15.7 (4)</td>
<td>29.8 ± 19.1 (4)</td>
</tr>
<tr>
<td>Atrpine (0.4 mg/kg iv)</td>
<td>4.9 ± 1.3 (4)</td>
<td>11.5 ± 0.7 (4)</td>
<td>16.7 ± 7.7 (4)</td>
<td>30.8 ± 18.2 (4)</td>
</tr>
<tr>
<td>l-NNA (0.15 mg/kg iv)</td>
<td>8.4 ± 3.0 (3)</td>
<td>13.2 ± 2.6 (3)</td>
<td>16.2 ± 1.0 (3)</td>
<td>16.1 ± 1.8 (3)</td>
</tr>
<tr>
<td>l-NNA (1.5 mg/kg iv)</td>
<td>15.5 ± 3.1 (5)</td>
<td>24.5 ± 3.4 (5)</td>
<td>38.1 ± 4.8 (5)</td>
<td>43.8 ± 5.1 (5)</td>
</tr>
</tbody>
</table>

Values are means ± SE (in μm/min); numbers in parentheses indicate numbers of animals. Intragastric pressure (IGP) is the pressure above baseline (60 mmH₂O). *P < 0.05; †P < 0.001.
measurement of small changes in the size of the fundus not usually apparent with other methods. The crystals used in the present study were 1 mm in diameter, allowing several to be placed on the surface of the murine fundus and body of the stomach. Each crystal serves as both a receiver and a transmitter, enabling distances to be calculated from any one crystal to any other crystal. The crystals measure the sound signal as it passes through the wall of the organ studied, thereby avoiding the problem faced by conventional ultrasonography, which relies on sound energy transmission through the whole organ.

Fig. 7. Effect of the nitric oxide synthase inhibitor N^\text{N}-nitro-l-arginine (l-NNA) on fundic tone and wall compliance. l-NNA (1.0 mg/kg iv) decreased the ICD of crystals placed on the fundus of an empty stomach with no perturbation of the IGP, indicating a contractile effect on the fundus (A). In contrast, when the stomach IGP was set to 60 mmHg, l-NNA increased the fundic ICD (B, solid trace). This increase in ICD was preceded by a decease in the ICD of crystals placed on the antrum (B, grey trace) and by an increase in IGP (B, shaded area). l-NNA also decreased the compliance of the gastric fundus, suggesting a stiffer fundic wall (C).

Fig. 8. Representative traces of the effect of the 5-HT_{1A} receptor agonist buspirone (A) and the 5-HT_{1B/D} agonist sumatriptan (B) on the murine fundus. The initial IGP was set to 60 mmHg. Buspirone (0.12 mg/kg iv) caused an increase in ICD, suggesting an increase in fundic size, whereas sumatriptan (0.1 mg/kg iv) decreased ICD, suggesting a decrease in fundic size.

Fig. 9. Effect of the 5-HT_{1A} receptor agonist buspirone (A) and the 5-HT_{1B/D} agonist sumatriptan (B) on fundic compliance. Buspirone (A, predrug: solid line; postdrug: grey line, 0.15 mg/kg iv) initially increased the ICD, suggesting that it increased fundic size but did not subsequently alter compliance. Sumatriptan (0.1 mg/kg iv) initially decreased ICD, suggesting that it increased fundic size but, in contrast with buspirone, subsequently increased fundic compliance (B; predrug: solid line; postdrug: grey line). CTRL, control.
Therefore, air within the stomach, particularly the fundus (which often contains air), does not interfere with the ability to record the signal or to determine fundic size. A significant advantage over current methodology is that the technique can be used in small animals with intact extrinsic innervation. This is of particular importance when studying physiological processes, such as fundic accommodation, that require intact vagovagal reflexes (4, 35, 39). These reflexes are lost in muscle strip experiments or in experiments on the isolated stomach.

Fundic compliance (d/l/dp) measures the deformability of the fundus by measuring the volume change resulting from a pressure change. Compliance is widely used to evaluate the distensibility of hollow organs such as the gastrointestinal tract, lungs, heart, and bladder. In the present study, in addition to measuring compliance, we also reported on another coefficient, FDR (d/l/dh), which was used to provide a measure of the kinetics of distensibility. The value of the FDR is inversely proportionate to the dynamic changes in resistance to distensibility of the fundic wall. This resistance has a passive component from structures that make up the fundic wall and an active component as a result of interactions between enteric nerves, interstitial cells of Cajal, and smooth muscle cells. The FDR coefficient provides information in addition to the fundic compliance coefficient as it is an expression of compliance without requiring a static measure of maximum distension. FDR gives a measure of the rate of distension in response to a given pressure. As is seen in Table 2, FDR varied at different pressure points with a threefold increase in FDR at 40 compared with 10 mmH2O, likely reflecting active accommodation, thereby reducing resistance to distension.

As we developed the techniques required to measure changes in fundic size, we encountered limitations to the methodology. Our studies were carried out in the acute setting with anesthetized mice. In separate experiments (data not shown), we used mouse antral muscle strips to determine the effect of various anesthetics on contractile activity. We tested all anesthetic drugs currently approved by our institution for use in mice, including diazepam, ketamine, pentobarbital, thiopental, and thamylal at the recommended doses. All anesthetic drugs tested affected contractile activity, decreasing contractile amplitude. The combination of ketamine and xylazine had the least effect on spontaneous activity and was therefore used in this study.

Another limitation of the ultrasonomicrometry technique is that it does not directly measure IGP. This limitation was highlighted in our experiments with l-NNA as a NO synthase inhibitor. l-NNA would be expected to reduce NO production and therefore cause the fundus to contract. In contrast, an apparent relaxation was seen. Use of an IGP monitoring device and placement of additional crystals on the distal body of the stomach showed that l-NNA caused contraction of the gastric body, resulting in displacement of fluid from the stiffer distal stomach to the more compliant proximal stomach, thereby distending the fundus. These results suggest that it is important to monitor IGP simultaneously when using ultrasonomicrometry in all experiments. Moreover, while the technique has been successfully used to isolate longitudinal and circular muscle contraction or relaxation by placing crystals along the axis of contraction (2), this is harder to accomplish in the fundus. This is due to the spherical nature of the fundus, making the axis of contraction different in different parts of the fundus. We therefore did not attempt to separate out the contribution of each muscle layer to the changes observed.

The experiments directed toward determining the effect of modulation of cholinergic, nitrergic, and serotonergic pathways on murine fundic size and compliance in the intact innervated stomach revealed different contributions of each pathway to regulation of fundic tone. As previously shown (38), there appears to be a baseline cholinergic input maintaining fundic tone as atropine resulted in an increase in ICD, suggesting relaxation. The data obtained using l-NNA to inhibit NO production while monitoring IGP and antral size are also in agreement with those obtained in other intact animals, including humans (32, 40). Furthermore, l-NNA decreased fundic compliance and markedly altered the rate of fundic relaxation to a given pressure, suggesting that there also was a baseline nitrergic input to fundic smooth muscle and that, in the absence of NO, the fundus is stiffer and nonrelaxing. A serotonergic modulation of fundic tone has been previously reported (27, 29, 41). In contrast to data obtained from humans (8) and dogs (12), sumatriptan did not relax the murine fundus. Instead, a decrease in ICD was seen, suggesting a contraction. Sumatriptan did subsequently increase fundic compliance. These data suggest that there are species differences in the serotonergic modulation of fundic size and, although they are different from human and canine data, are similar to those seen in the cat, where sumatriptan also contracts the cat fundus (J. Tack, personal communication).

In summary, the validation experiments carried out in this study show that ultrasonomicrometry accurately measures distance in the mouse fundus, can be used in the intact animal, has an excellent resolution, and can measure the biological responses to drugs when IGP is also monitored. The data obtained with sumatriptan and buspirone also suggest that ultrasonomicrometry can also be used to explore mechanisms of action of drugs. As highlighted by the results obtained with sumatriptan experiments, important species differences may be present in the response of the fundus to a given drug.

GRANTS
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