Behavioral, autonomic, and endocrine outputs of the CNS are subject to important feedback modulation by viscerosensory signals that are conveyed initially to the hindbrain nucleus of the solitary tract (NST). In the present study, noradrenergic (NA) neurons [i.e., those that express the NA synthetic enzyme dopamine β hydroxylase (DbH)] in the caudal NST were lesioned to determine their role in mediating anorexigenic responses to gastric stimulation and in conveying gastric sensory signals to the hypothalamus and amygdala. For this purpose, saporin toxin conjugated to an antibody against DbH was microinjected bilaterally into the caudal NST in adult rats. Control rats received similar microinjections of vehicle. Several weeks later, rats were tested for the ability of systemic cholecystokinin octapeptide (CCK) (0 or 10 μg/kg) to inhibit food intake. CCK-induced anorexia was significantly attenuated in toxin-treated rats. Rats subsequently were used in a terminal cFos study to determine central neural activation patterns after systemic CCK or vehicle and to evaluate lesion extent. Toxin-induced loss of DbH-positive NST neurons was positively correlated with loss of CCK-induced anorexia. Hypothalamic cFos expression was markedly attenuated in lesioned rats after CCK treatment, whereas CCK-induced neural activation in the parabrachial nucleus and amygdala appeared normal. These findings suggest that hindbrain NA neurons are an integral component of brainstem circuits that mediate CCK-induced anorexia and also are necessary for hypothalamic but not parabrachial or amygdala responses to gastric sensory stimulation.

Key words: gastrointestinal; vagus; nucleus of the solitary tract; hypothalamus; amygdala; parabrachial nucleus; saporin toxin

Introduction

Behavioral, autonomic, and endocrine outputs of hypothalamic and limbic forebrain circuits are subject to important feedback modulation by viscerosensory signals from the gut, and bidirectional interactions between gastrointestinal function and behavioral–cognitive state are well recognized. These interactions contribute to the clinical problem of anorexia after chemotherapeutic cancer treatment and may underlie the high comorbidity of gastrointestinal disorders with anxiety, depression, and other psychiatric disorders (Ballenger et al., 2001; Mayer et al., 1998). Small doses of CCK in rats and in humans mimic the physiological satiety effects of endogenous CCK released from the gut postprandially (Schwartz and Moran, 1998). Conversely, large doses of CCK (i.e., 10–100 μg/kg) produce anorexia that generally is accompanied by nausea, pituitary release of stress hormones, and, in humans, anxiety (Verbalis et al., 1986a,b; Kamiliris et al., 1992; McCutcheon et al., 1992; Greenough et al., 1998). Combined radiofrequency lesions of the area postrema (AP) and portions of the medial NST in rats virtually eliminate the anorexigenic effect of low doses of exogenous CCK (i.e., 1–2 μg/kg) (Edwards et al., 1986). However, these lesions spared a portion of the caudal medial and commissural NST, and a higher dose of CCK (i.e., 8 μg/kg) was still effective in inhibiting food intake (Edwards et al., 1986).

In rats, systemic CCK activates ascending projections to the hypothalamus and amygdala that arise primarily from noradrenergic (NA) [i.e., dopamine β hydroxylase (DbH) positive] neurons of the A2 cell group, located within the caudal medial and commissural NST (Rinaman et al., 1995; Myers and Rinaman, 2000). CCK also activates a smaller contingent of noncatecholaminergic NST neurons with ascending projections; these are immunoreactive for glucagon-like peptide 1 (GLP-1) (Rinaman, 1999; Myers and Rinaman, 2000). Other ascending viscerosensory pathways activated by CCK include NST projections that are relayed through the ventrolateral medulla (VLM) and pontine parabrachial nucleus (PBN). Ascending NA pathways are implicated in viscerosensory activation of the hypothalamus.
Materials and Methods

Animals. All procedures conformed to National Institutes of Health guidelines and were approved by the University of Pittsburgh Animal Care and Use Committee. Data from 20 adult male Sprague Dawley rats (Harlan Sprague Dawley, Indianapolis, IN) are included in this report. Rats were singly housed with ad libitum access to water and pelleted rat chow (Purina, St. Louis, MO) except during food intake experiments. Colony rooms were maintained at 22–23°C and kept on a 12 hr light/dark cycle, with lights on from 7:00 A.M. to 7:00 P.M. Rats weighed 180–225 g at the time of toxin injection surgery and 200–325 g at the time of food intake experiments and the terminal cFos study.

Experimental groups. Rats received bilateral injections of anti-DbH–saporin toxin (toxin; n = 10; Advanced Targeting Systems, San Diego, CA) or 0.15 M NaCl (vehicle; n = 10) into the caudal NST. Each rat was used in a food intake experiment and in a terminal cFos study.

NST microinjection. Toxin or vehicle microinjections were made into the NST at six sites (Fig. 1). Injection sites encompassed the NST region that contains the highest incidence of cFos expression by DbH–positive neurons after CCK treatment, corresponding to the location of the A2 cell group within the caudal medial NST (Sawchenko and Swanson, 1982; Rinaman et al., 1993; Rinaman et al., 1995). Rats were anesthetized with halothane (2–5% in 100% oxygen) and secured in a stereotaxic frame using blunt ear bars, with the head ventroflexed. The skin over the dorsal neck surface was shaved and incised, and the neck muscles were retracted and bluntly dissected to expose the meninges overlying the dorsal surface of the caudal medulla. With the aid of a surgical microscope, the meninges were cut with a sterile needle to reveal the AP. Under visual guidance, a glass micropipette tip (outer diameter of 50–75 μm) filled with toxin or vehicle and affixed to a 0.5 μl Hamilton syringe was positioned on the midline at the caudal limit of the AP and was then moved 0.25 mm lateral and 0.5 mm below the medullary surface for the first, most caudal injection site. The second injection site was located 0.25 mm below the lateral border of the AP at its midrostrocaudal extent, and the third site was located 0.25 mm below the lateral border of the AP at its most rostral extent (Fig. 1). The three NST injection sites were then duplicated contralaterally. At each injection site, 50 nl of sterile 0.15 M NaCl vehicle containing 0 or 5 ng of toxin was delivered by manual pressure injection over ~30 sec. Toxin was freshly prepared from a frozen stock solution within 2 hr of injection. Results from a published report (Madden et al., 1999) and additional pilot studies confirmed that this concentration and volume of toxin destroyed the majority of DbH–positive neurons within the caudal NST (see Results). Lower toxin doses were less effective, whereas higher doses produced marked tissue necrosis. After each microinjection, the pipette tip was left in place for 1–2 min and then withdrawn. The skin incision was closed with stainless steel clips after the final injection. Rats were returned to their home cages after recovery from anesthesia.

The fact that saporin toxin microinjected near NA cell bodies effectively lesions those cells provides empirical evidence for vesicular release and concomitant extracellular exposure of DbH within the toxin injection site (Madden et al., 1999). The dorsal vagal complex (DVC) receives relatively sparse input from other medullary and pontine NA cell groups but is richly innervated by DbH–positive terminals of presumably local origin. Somatic and/or dendritic vesicular release from NA neurons also may occur, as has been reported for local monoamine release within the locus ceruleus and other central sites (Pudovkina et al., 2001).

Posturgical maintenance. Rats with NST vehicle injections attained their presurgical body weights (BW) within 2–3 d after surgery and continued to gain BW thereafter (Fig. 2). Conversely, although the 10 rats with bilateral toxin injections also attained their presurgical BWs within 2–3 d, several began to lose BW beginning 6–7 d after surgery. This effect is consistent with the delayed time course of toxin-induced neuronal death (Madden et al., 1999). To encourage food intake in these rats while maintaining consistent dietary treatment within both groups, all vehicle- and toxin-injected rats were given daily overnight access to 25 ml of
palatable liquid diet (vanilla-flavored Ensure) in addition to ad libitum chow, regardless of their postsurgical BW profile. Liquid diet access was initiated 7 d after NST injection surgery and discontinued after 2 weeks, by which time the majority were consuming their full allotment of liquid diet plus additional chow each day, and all had surpassed their prestudy BWs. Rats continued to gain at least 1 gm of BW per day after termination of liquid diet access. Three toxin-injected rats did not consume adequate amounts of chow or liquid diet to maintain BW. These rats were killed without data collection and are not included in the reported n values.

Food intake experiment. At least 10 d before the terminal cFos experiment described below, rats with bilateral NST injections of toxin (n = 10) or vehicle (n = 10) were used to determine the effects of food intake produced by intraperitoneal administration of CCK. For this purpose, rats were removed from their original housing (i.e., a new environment with the same 12 hr light/dark cycle. Rats were housed individually in clear Plexiglas boxes (25 × 30 cm floor, 22 cm height) with stainless steel rod floors, each equipped with a computer-driven pellet delivery and monitoring system (Med Associates, E. Fairfield, VT). Drinking water was available ad libitum from sipper tubes within each box. Food access was restricted to a daily 4 hr period beginning at lights out (7:00 P.M.), when a single 45 mg chow pellet (Precision Dustless Pellets; Bio-Serv; Frenchtown, NJ) was delivered automatically to a shallow feeding trough in each cage, thereby breaking a photobeam crossing the base of the trough. A new pellet was delivered automatically each time the preceding pellet was removed. The cumulative number of pellet deliveries (i.e., photobeam breaks) was recorded automatically at 15 min intervals during the 4 hr feeding period. Cage trays beneath the open rod floors were routinely inspected at the end of the feeding period to ensure that delivered pellets were being consumed. Data were collected and stored using Med PC software (Med Associates). Rats were acclimatized to the new environment and feeding schedule for 1 week before testing, by which time stable daily 4 hr food intakes were achieved. Rats also were acclimated to gentle handling and occasional intraperitoneal saline injections during this period.

On testing day 1, all rats were injected intraperitoneally with 2.0 ml of 0.15 M NaCl vehicle. Food access was initiated within 2 min, at lights out (7:00 P.M.). Cumulative pellet delivery data were collected for 4 hr. On testing day 2, all rats were injected intraperitoneally with 2.0 ml of vehicle containing CCK (sulfated octapeptide; 10 μg/kg BW; Sigma, St. Louis, MO). Food access was initiated within 2 min after CCK injection, at lights out (7:00 P.M.). Cumulative pellet delivery data were collected for 4 hr. Thus, each rat served as its own control for the effects of intraperitoneal vehicle vs. intraperitoneal CCK treatment on subsequent 4 hr food intake. Rats were kept in the same housing environment but were returned to ad libitum chow access after testing day 2.

Food intake data analysis. BWs ranged from 200 to 293 gm on testing days; thus, intake (i.e., pellet delivery) data for each rat were converted to grams (number of pellets × 45 mg) and expressed as percentage BW. Intake data were combined a priori according to NST injection group (i.e., toxin vs vehicle) and expressed as group mean ± SE. Treatment-related differences in food intake were tested for statistical significance by using ANOVA, with NST injection plus intraperitoneal treatment (i.e., NST toxin or NST vehicle plus intraperitoneal CCK or intraperitoneal vehicle) as the independent variable. When F values indicated a significant main effect of treatment on food intake, the ANOVA was followed up with post hoc t tests using Dunn’s (Bonferroni) correction for multiple comparisons. Differences were considered significant when p < 0.05. Feeding data in rats with NST toxin injections also were analyzed by correlating CCK-induced feeding suppression with toxin-induced loss of DbH-positive neurons (see Results).

CCK-induced cFos experiment. A terminal cFos study was performed in each rat at least 4 weeks after NST injection surgery and at least 3 d after the food intake experiment. For this purpose, five rats from each NST injection group were injected intraperitoneally with 2.0 ml of 0.15 M NaCl vehicle, and five rats from each group were injected with vehicle containing CCK (10 μg/kg; Sigma). Injections were made between 9:00 A.M. and 10:00 A.M.

Rats were returned to their home cages immediately after intraperito-
cFos expression. Thus, cFos labeling in the pontine PBN was quantified by determining the number of CGRP-positive neurons in the lateral PBN that were activated to express cFos, using counting criteria similar to those described above. Tissue sections were examined bilaterally through the rostrocaudal extent of cellular cFos labeling in the lateral PBN (three to four sections spaced by 180 μm per rat). Activation in the hypothalamic paraventricular nucleus (PVN) was quantified by determining the number of magnocellular and parvocellular OT-positive neurons that contained nuclear cFos labeling. In each rat, two tissue sections through the PVN (spaced by 180 μm) containing both medial parvocellular and lateral magnocellular PVN subnuclei were selected for analysis. Amygdala activation was determined using CGRP fiber immunolabeling to distinguish the cytoarchitectural boundaries of the central nucleus of the amygdala (CeA). All cFos-positive profiles within those CeA boundaries were counted in tissue sections through the rostrocaudal extent of fibrous CGRP labeling (five to seven sections spaced by 180 μm in each rat).

Regional cell count data for each rat are expressed as the number of cells counted in each region bilaterally, divided by the number of analyzed tissue sections through the region (Table 1). Treatment group data are expressed as mean ± SE. Between-group differences in count values indicated a significant main effect of treatment on cell intake at the 30 and 60 min time points (*p < 0.05 compared with intake after control treatment).

<table>
<thead>
<tr>
<th>Rat</th>
<th>NST injection</th>
<th>Intrapentoneal treatment</th>
<th>NST #DBH</th>
<th>NST #cFos</th>
<th>VLM #DBH; cFos%</th>
<th>Lateral PBN #CGRP + cFos%</th>
<th>PVN #OT + cFos%</th>
<th>CeA #cFos%</th>
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<tr>
<td>1</td>
<td>Toxin</td>
<td>CCK</td>
<td>3.2</td>
<td>172.6</td>
<td>21.5; 4.7%</td>
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<td>CCK</td>
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<td>157.7</td>
<td>19.2; 5.2%</td>
<td>42.4</td>
<td>11.3</td>
<td>206.3</td>
</tr>
<tr>
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<td>CCK</td>
<td>34.8</td>
<td>185.3</td>
<td>28.9; 14.3%</td>
<td>26.3</td>
<td>6.0</td>
<td>192.6</td>
</tr>
<tr>
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<td>Toxin</td>
<td>CCK</td>
<td>39.6</td>
<td>194.1</td>
<td>30.0; 10.1%</td>
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<td>4.3</td>
<td>267.8</td>
</tr>
<tr>
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<td>CCK</td>
<td>41.4</td>
<td>218.4</td>
<td>24.4; 20.6%</td>
<td>30.3</td>
<td>2.6</td>
<td>126.5</td>
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<tr>
<td></td>
<td>Group mean ± SE</td>
<td></td>
<td>26.1 ± 7.9</td>
<td>185.6 ± 10.2</td>
<td>24.8 ± 3.1; 11.0 ± 3.1%</td>
<td>35.8 ± 3.2</td>
<td>5.9 ± 1.5</td>
<td>218.1 ± 30.0</td>
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<td>Toxin</td>
<td>Vehicle</td>
<td>0.3</td>
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<td>21.9; 5.4%</td>
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<td>9.5</td>
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</tr>
<tr>
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<td>Vehicle</td>
<td>1.4</td>
<td>12.3</td>
<td>15.7; 13.3%</td>
<td>4.5</td>
<td>7.3</td>
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</tr>
<tr>
<td>8</td>
<td>Toxin</td>
<td>Vehicle</td>
<td>19.2</td>
<td>15.4</td>
<td>22.2; 4.5%</td>
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<td>5.6</td>
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<td>23.0; 0%</td>
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<td>14.2</td>
<td>11.3</td>
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<tr>
<td>10</td>
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<td>30.1; 3.3%</td>
<td>2.3</td>
<td>4.3</td>
<td>4.4</td>
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<tr>
<td></td>
<td>Group mean ± SE</td>
<td></td>
<td>19.0 ± 9.5</td>
<td>20.1 ± 5.1</td>
<td>22.6 ± 2.3; 5.3 ± 2.2%</td>
<td>8.9 ± 3.0</td>
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<td>16.2 ± 4.0</td>
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<td>160.6</td>
<td>35.7/24.7%</td>
<td>41.5</td>
<td>27.5</td>
<td>221.9</td>
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<tr>
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<td>CCK</td>
<td>95.7</td>
<td>174.0</td>
<td>36.5/19.4%</td>
<td>37.5</td>
<td>29.3</td>
<td>142.5</td>
</tr>
<tr>
<td>13</td>
<td>Vehicle</td>
<td>CCK</td>
<td>108.3</td>
<td>221.3</td>
<td>30.8/20.5%</td>
<td>31.3</td>
<td>48.6</td>
<td>87.8</td>
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<tr>
<td>14</td>
<td>Vehicle</td>
<td>CCK</td>
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<td>205.1</td>
<td>43.6/28.0%</td>
<td>48.3</td>
<td>42.4</td>
<td>185.3</td>
</tr>
<tr>
<td>15</td>
<td>Vehicle</td>
<td>CCK</td>
<td>120.0</td>
<td>263.4</td>
<td>31.4/31.3%</td>
<td>59.6</td>
<td>36.5</td>
<td>204.1</td>
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<td></td>
<td>Group mean ± SE</td>
<td></td>
<td>104.3 ± 6.8</td>
<td>204.9 ± 18.2</td>
<td>35.6 ± 2.3; 24.8 ± 2.8%</td>
<td>43.6 ± 4.9</td>
<td>36.9 ± 4.0</td>
<td>168.3 ± 24.1</td>
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<td>Vehicle</td>
<td>90.0</td>
<td>21.0</td>
<td>32.6/0%</td>
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<td>17</td>
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<td>Vehicle</td>
<td>107.2</td>
<td>9.2</td>
<td>29.6/4.6%</td>
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<td>18</td>
<td>Vehicle</td>
<td>Vehicle</td>
<td>105.3</td>
<td>9.6</td>
<td>28.4/9.8%</td>
<td>14.6</td>
<td>13.5</td>
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<tr>
<td>19</td>
<td>Vehicle</td>
<td>Vehicle</td>
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<td>20</td>
<td>Vehicle</td>
<td>Vehicle</td>
<td>124.3</td>
<td>3.4</td>
<td>32.5/6.1%</td>
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<td>9.1</td>
<td>17.0</td>
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<td>Group mean ± SE</td>
<td></td>
<td>108.5 ± 5.7</td>
<td>12.4 ± 3.3</td>
<td>30.7 ± 0.8; 5.8 ± 1.7%</td>
<td>9.4 ± 1.9</td>
<td>7.9 ± 1.6</td>
<td>18.6 ± 3.8</td>
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Figure 3. Food intake in rats with previous NST vehicle injections (n = 10) after control treatment (0.15 M NaCl) and after CCK (10 μg/kg). CCK treatment significantly inhibited food intake at the 30 and 60 min time points (*p < 0.05 compared with intake after control treatment).

**Results**

The results of this study support the view that NA neurons in the caudal NST are a necessary component of central circuits that underlie inhibition of food intake after CCK treatment. NA neurons also appear to be necessary for conveying CCK-related gastric sensory signals to the PVN but appear unnecessary for conveying such signals to the lateral PBN and CeA.

**NST injection effects on CCK-induced anorexia**

Food intake after intraperitoneal vehicle (0.15 M NaCl) and after CCK (10 μg/kg) treatment was similar in rats with previous NST injections of vehicle (Fig. 3) or toxin (Fig. 4). As expected, CCK treatment suppressed food intake in rats with NST vehicle injections (Fig. 3). Analysis of data from individual rats confirmed the generality of this effect; that is, CCK significantly inhibited food intake in every rat within the
Figure 4. Food intake in rats with previous NST toxin injections (n = 10) after control treatment (0.15 M NaCl) and after CCK (10 μg/kg). CCK did not significantly inhibit food intake at any time point.

The ability of CCK to inhibit food intake was attenuated in rats with bilateral NST toxin injections. Group food intake after CCK treatment was not significantly different from intake after intraperitoneal vehicle at any time point (Fig. 4). However, the effects of CCK treatment were variable within this group. Analysis of data from individual rats revealed that CCK inhibited food intake by <15% in six rats with bilateral NST toxin injections but inhibited food intake by 30–60% in the remaining four (see Fig. 6). The ability of CCK to suppress food intake was strongly correlated with NA lesion extent, as described below.

**Toxin-induced lesions of DbH-immunopositive neurons**

Bilateral toxin injections were associated with an average 80% reduction in DbH-positive NST neurons relative to the average number of DbH cells per section in rats with NST vehicle injections (Table 1; Fig. 5, compare A, B). Loss of DbH-positive NST cells in individual toxin-injected rats ranged from 49.5 to 99.7% relative to average counts in vehicle-injected rats. These reductions were significant in every case (p < 0.05). Thus, toxin injections effectively but variably destroyed DbH-positive NST neurons.

A toxin-related decrement of DbH-positive caudal VLM neurons also was observed in many rats (Table 1). Toxin-injected rats displayed an average of 24 DbH-positive VLM neurons per section analyzed, significantly fewer than the average 33 cells per section counted in rats with NST vehicle injections (p < 0.05). Thus, in addition to destroying DbH-positive NST neurons, toxin injections often destroyed a subset of DbH-positive VLM neurons.

Qualitative analysis of TH and GLP-1 immunolabeling in three lesioned rats with the most complete loss of DbH-positive NST neurons indicated that the toxin spared dopaminergic neurons in the vagal motor nucleus and caudal NST (Fig. 5C) and GLP-1-positive peptidergic neurons in the caudal NST (Fig. 5D) and adjacent reticular formation. The GLP-1 labeling was further examined quantitatively. The average total number of GLP-1 neurons counted in the three toxin-injected rats (129 ± 5.7 neurons in five to seven tissue sections per rat) was consistent with a previous report of GLP-1-positive neurons in intact rats (Rinaman, 1999).

**Relationship between extent of NST lesion and CCK-induced anorexia**

As reported above, statistical analysis of group data indicated that CCK did not significantly inhibit average food intake in rats with toxin-induced NST lesions (Fig. 4). However, in four rats within this group, CCK suppressed food intake by ~28–60% relative to intake after intraperitoneal vehicle treatment (Fig. 6). Postmortem analysis of lesion extent indicated that ~33–50% of DbH-positive NST neurons were spared in these four cases (Table 1, rats 3–5, 10) relative to average DbH cell counts in nonlesioned rats. There was a strong positive correlation between the remaining number of NST DbH cells and the magnitude of CCK-induced anorexia at its maximum extent (i.e., at the 30 min time point) (Fig. 6).

**CCK-induced cFos expression in lesioned and nonlesioned rats**

NST cFos activation was significantly greater in lesioned and nonlesioned rats after CCK treatment (i.e., range of 158–263 cells per section; Table 1) compared with intraperitoneal vehicle (i.e., range of 3–39 cells per section; Table 1). On average, somewhat more NST cFos expression was observed after intraperitoneal vehicle treatment in rats with toxin lesions (~20 cells per section) compared with cFos in rats without lesions (~12 cells per section), but this difference was not significant. Average NST cFos expression was elevated similarly in lesioned and nonlesioned rats after CCK treatment (i.e., 186 and 205 cells per section, respectively), suggesting that NST toxin lesions eliminated a relatively small subset of the NST neurons activated by CCK (Fig. 5). In a previous report, CCK administered at a dose of 100 μg/kg activated cFos expression in ~57% of GLP-1-positive neurons in intact rats (Rinaman, 1999). The 10-fold lower dose of CCK used in the present study activated cFos expression in a somewhat smaller proportion (i.e., 34.8% ± 7.9) (Fig. 5D) of GLP-1-positive neurons in the three rats with the most extensive loss of DbH-positive neurons.

Relatively few cFos-positive neurons were present in the VLM, lateral PBN, PVN, or CeA in lesioned or nonlesioned rats after intraperitoneal vehicle (Table 1). After CCK treatment, ~25% of DbH-positive VLM neurons were activated in nonlesioned rats, whereas a significantly smaller proportion (11%) were activated in lesioned rats (p < 0.05) (Table 1). VLM activation in lesioned rats after CCK was not significantly different from activation after intraperitoneal vehicle (Table 1). NST lesions were associated with even more marked reductions in CCK-induced cFos expression in the PVN (Fig. 7), including a significant attenuation of cFos expression by OT-positive neurons (Fig. 8). CCK activated an average of 37 OT-positive cells per section analyzed in nonlesioned rats but only approximately six cells per section in lesioned rats (Table 1). These PVN OT activation values were not significantly different in lesioned rats after CCK treatment compared with OT activation after intraperitoneal vehicle (Table 1).

In contrast to the reduced VLM and PVN activation after CCK treatment in lesioned rats, CCK-induced cFos expression within the lateral PBN (Figs. 8, 9) and CeA (Fig. 10) was not attenuated in lesioned rats (Table 1). Despite a clear loss of DbH-positive fibers within the lateral PBN (Fig. 9) and CeA (Fig. 10) in lesioned rats, CCK treatment in lesioned and nonlesioned rats activated statistically similar numbers of CGRP-positive PBN neurons (~36 and ~44 cells per section, respectively) and similar numbers of cells within the CGRP-rich region of the CeA (~218 and 168 cells per section, respectively) (Table 1).
Circled numbers represent individual rats; see Table 1 for cell counting data.

point (expressed as percentage of intake in the same rats after intraperitoneal vehicle injection).

The extent of CCK-induced inhibition of food intake (FI) at the 30 min time remaining in 10 lesioned rats (expressed as percentage of the average number of DbH cells in remaining in 10 lesioned rats (expressed as percentage of the average number of DbH cells in nonlesioned rats) and the extent of CCK-induced inhibition of food intake (FI) at the 30 min time point (expressed as percentage of intake in the same rats after intraperitoneal vehicle injection). Circled numbers represent individual rats; see Table 1 for cell counting data.

Figure 5. Color photomicrographs depicting CCK-induced cFos expression (blue–black nuclear label) within the dorsal vagal complex in a nonlesioned rat (intact; A) and in a rat with a toxin-induced lesion of DbH-positive neurons (B–D are from the same animal). Tissue sections are double labeled for DbH (A, B), TH (C), or GLP-1 (D; arrows point out activated neurons). Scale bar in A applies also to B and C.

Figure 6. Significant positive correlation between the number of DbH-positive NST neurons remaining in 10 lesioned rats (expressed as percentage of the average number of DbH cells in nonlesioned rats) and the extent of CCK-induced inhibition of food intake (FI) at the 30 min time point (expressed as percentage of intake in the same rats after intraperitoneal vehicle injection). Circled numbers represent individual rats; see Table 1 for cell counting data.

Discussion

Saporin toxin lesion efficacy and specificity

NST microinjections of DbH-saporin toxin conjugate were designed to destroy NA neurons in the caudal medial NST, corresponding to the location of the A2 cell group and the caudal portion of the C2 cell group. Previous parametric studies attest to the efficacy and specificity of this toxin conjugate (Wrenn et al., 1996; Madden et al., 1999; Ritter et al., 2001; Fraley et al., 2002; Fraley and Ritter, 2003). In the present study, toxin-induced loss of DbH cell and terminal immunostaining within the caudal DVC and loss of terminal labeling in the PBN, PVN, and CeA offers support for lesion efficacy. Although it is possible that the toxin also destroyed non_DbH-containing neurons, it did not destroy GLP-1-positive neurons, and it also left local TH-positive dopaminergic neurons intact. Furthermore, similar numbers of cFos-positive NST neurons were observed in lesioned and nonlesioned rats after CCK treatment, consistent with evidence that NA neurons comprise a relatively small subset of the total population of NST neurons activated by systemic CCK (Rinaman et al., 1993).

A reduced number of DbH-positive neurons within the VLM was observed in a subset of toxin-injected rats (Table 1). Because NA neurons in the VLM are not known to provide synaptic input to the caudal NST, their loss may be attributed to toxin spread from injection sites into the adjacent reticular formation. However, VLM NA cell loss did not occur in every toxin-injected rat and was not necessary for either attenuation of CCK anorexia or for suppression of CCK-induced hypothalamic cFos.

A transient baseline anorexia and BW loss emerged ~7 d after toxin injections in many rats but gradually resolved when rats had access to liquid diet supplement. Speculatively, dysfunction of NST NA neurons may disrupt digestive vago-vagal reflexes for which local NA signaling mechanisms are implicated (Rogers et al., 2003), thereby leading to gastric malaise and/or early satiety. The gradual recovery of BW gain in lesioned rats suggests that compensatory feeding control systems become operational to restore baseline food intake, despite the loss of NST NA neurons. Indeed, 4-hr food intake was similar in lesioned and control rats after intraperitoneal saline injection. Patterns of ad libitum intake were not analyzed in this study, and it is possible that differences exist between lesioned and control rats in meal number or size.

Correlation between NA cell loss and CCK-induced anorexia

A strong positive correlation was found between the number of surviving NA neurons in the caudal NST and the extent of CCK-induced anorexia. As shown in Figure 6, the greater the loss of DbH-positive neurons, the more blunted the ability of CCK to inhibit food intake. A positive correlation between NA neural activation and feeding-induced gastric distension also has been reported (Rinaman et al., 1998). Gastric distension inhibits food intake (Phillips and Powley, 1996), and CCK mimics and amplifies gastric distension vagal afferent signals (Schwartz et al., 1993; Schwartz and Moran, 1998). Thus, not only are NA neurons in the caudal medial NST activated by physiological signals that inhibit food intake, they appear to be necessary for the ability of exogenous CCK to inhibit food intake.

Although the hypothalamic clearly is involved in food intake and energy balance (Watts, 2000; Broberger and Hokfelt, 2001; Berthoud, 2002), it is unclear whether reduced PVN activation in lesioned rats after CCK treatment is associated with the blunted anorexigenic response in those rats. CCK does not activate hypothalamic or other forebrain neurons in neonatal rats, but it inhibits food intake as early as postnatal day 1 (Rinaman et al., 1994). CCK also inhibits sucrose intake in adult decerebrate rats that lack brainstem–hypothalamic connections (Grill and Smith, 1988), and CCK anorexia is preserved in rats after PVN microin-
Dissociation of NA lesion effects on PVN and PBN–CeA activation after CCK treatment

In addition to inhibiting food intake, systemic CCK increases plasma levels of stress hormones, including ACTH and OT in rats (Verbalis et al., 1986a; Kamilaris et al., 1992) and ACTH and AVP in ferrets, monkeys, and humans (Verbalis et al., 1987; Miasz et al., 1989; Billig et al., 2001). Exogenous CCK also activates NST NA neurons in ferrets (Billig et al., 2001) and monkeys (Schreiber et al., 1997), as in rats. Plasma hormone levels were not assayed in the present study, but the blunted PVN cFos expression observed in lesioned rats after CCK treatment predicts a similarly blunted or absent stress hormone response, as reported in neonatal rats (Rinaman et al., 1994).

In contrast to the decremented PVN activation in lesioned rats after CCK, activation in the lateral PBN and CeA apparently was unaffected. The unexpected dissociation between lesion effects on the hypothalamus and amygdala suggests that NA neurons in the caudal NST are unnecessary for conveying gastric viscerosensory signals to the PBN and CeA. Although the CeA receives direct inputs from the NST that include a prominent NA component, the PBN is the main relay for ascending viscerosensory inputs to the CeA (Saper and Loewy, 1980; Milner et al., 1984; Herbert et al., 1996; Jia et al., 1994). CGRP provides a distinct neurochemical marker for the viscerosensory PBN-to-CeA pathway (Yasui et al., 1989), and this pathway appeared to be activated normally in lesioned rats after CCK treatment. Although DbH-
positive NST neurons that project to the PBN and CeA presumably were lesioned, peptidergic NST and AP neurons that project to the PBN and CeA and do not colocalize DbH (cf. Herbert and Saper, 1990) presumably were spared. These include projections arising from GLP-1-positive neurons, which survived the lesions and were activated after CCK treatment. In this regard, a recent report indicates that GLP-1 receptor signaling in the CeA contributes to the formation of conditioned taste aversion (CTA), a sign of nausea, anxiety, and/or visceral illness in rats (Kinzig et al., 2002). It will be interesting to determine whether NA neurons in the NST are necessary or dispensable for CTA formation, because experimental dissociations between the ability of nauseogenic and anxiogenic stimuli to inhibit food intake and produce CTA have been reported previously (Curtis et al., 1994; Kinzig et al., 2002).

Gastric viscerosensory transmission to the hypothalamus and limbic forebrain

Psychiatrists and gastroenterologists have long appreciated the functional links between gut and brain. For example, the evolving disease model for irritable bowel syndrome (IBS) is a self-perpetuating cycle of positive feedback between the gut and brain, potentially initiated by pathological events in either the CNS or enteric nervous system (Mayer, 1999; Ballenger et al., 2001). Autonomic outputs of the hypothalamus and limbic forebrain are thought to contribute to IBS, which leads to altered visceral signaling in the hypothalamus and limbic forebrain (Mayer et al., 2001). Anxiety, depression, and panic disorder often are comorbid with IBS and are related to altered neural signaling in the hypothalamus and limbic forebrain (Mayer et al., 2001). It will be important to elucidate the detailed neurochemical and functional organization of neural pathways that convey gastric viscerosensory signals to these forebrain regions. The present results are consistent with the view that hindbrain NA neurons are critical for conveying gastric viscerosensory signals to the hypothalamus but are unnecessary for conveying gastric signals to the amygdala. Additional studies are needed to determine the relative importance of NA and non-NA pathways in mediating the behavioral and physiological effects of viscerosensory stimuli under normal and pathological conditions.

References


McCutcheon B, Ballard M, McCaffrey RJ (1992) Intraperitoneally injected...