Effects of cholecystokinin-8s in the nucleus tractus solitarius of vagally deafferented rats

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ABSTRACT

We have shown recently that cholecystokinin octapeptide (CCK-8s) increases glutamate release from nerve terminals onto neurons of the nucleus tractus solitarius pars centralis (cNTS). The effects of CCK on gastrointestinal-related functions have, however, been attributed almost exclusively to its paracrine action on vagal afferent fibers. Since it has been reported that systemic or perivagal capsaicin pretreatment abolishes the effects of CCK, the aim of the present work was to investigate the response of cNTS neurons to CCK-8s in vagally deafferented rats. In surgically deafferented rats, intraperitoneal administration of 1 or 3 µg/kg CCK-8s increased c-Fos expression in cNTS neurons (139 and 251 % of control, respectively), suggesting that CCK-8s’ effects are partially independent of vagal afferent fibers. Using whole cell patch clamp techniques in thin brainstem slices, CCK-8s increased the frequency of spontaneous and miniature excitatory postsynaptic currents in 43% of the cNTS neurons via a presynaptic mechanism. In slices from deafferented rats, the percentage of cNTS neurons receiving glutamatergic inputs responding to CCK-8s decreased by approximately 50%, further suggesting that central terminals of vagal afferent fibers are not the sole site for the action of CCK-8s in the brainstem. Taken together, our data suggest that the sites of action of CCK-8s include the brainstem, and in cNTS the actions of CCK-8s are not restricted to vagal central terminals but that non-vagal synapses are also involved.
INTRODUCTION

Cholecystokinin, which is released from intestinal cells following ingestion of nutrients (34; 37; 47), has profound effects on gastrointestinal functions. In fact, it is well known that cholecystokinin, and in particular the cleaved octapeptide cholecystokinin-8s (CCK-8s), induces vagally-mediated gastric relaxation, increases pancreatic exocrine secretion and induces short-term satiety (36; 38; 41; 42; 48; 62).

Many studies have proposed that the vagally-mediated effects of CCK-8s are almost exclusively due to a paracrine action of CCK-8s on peripheral, capsaicin-sensitive vagal afferent fibers (8; 14; 41). This postulate stems mainly from the observations that C-fiber ablation induced by perivagal capsaicin treatment greatly attenuates, or abolishes altogether, the effects of systemic administration of CCK-8s on gastric motility, gastric acid secretion and pancreatic exocrine secretion (32; 33; 41; 58). One must keep in mind, however, that the localized perineural application involves the use of very high concentrations of capsaicin, usually 33mM (i.e. 1% solution)(9; 23; 25; 41; 56; 63; 67). Such a high concentration of capsaicin raises the possibility that its observed effects are due to the toxic effects of massive calcium influx on fibers or neurons other than C-fibers due to the activation of VR-1 receptors. Although it is well known that the effects of capsaicin on ionic conductances are transient (reviewed in (7; 26)), it is also accepted that administration of elevated doses of capsaicin induces degenerative changes in CNS neurons and fibers, including vagal efferent fibers, that do not express VR1 receptors and even involves neuronal areas that do not receive sensory inputs from the periphery (24; 26; 49).

Furthermore, the very composition of the vagus, where more than 85% of the fibers are afferent, implies that even minor damages induced by perivagal application of capsaicin to the efferent fibers could not only be overlooked easily, but would also have a potentially major impact on the vagal motor output, leading to an overestimation of the contribution of vagal afferent fibers to the CCK-8s mediated gastrointestinal effects.

Finally, should the physiological effects of CCK on gastrointestinal and feeding-related circuits be due exclusively to a paracrine action of the hormone, then the response to systemic administration of CCK-8s falls short of explaining its mechanism of action. In fact, it is highly unlikely that systemically administered CCK-8s could cross the lamina propria at a concentration sufficient to induce a paracrine effect. It is very likely, therefore, that the actions of CCK are not limited to its paracrine effect on peripheral vagal afferent fibers, but other sites of action, including the second order neurons of the NTS, should be taken into account. Indeed we, and others, showed recently that CCK-8s increases glutamatergic excitatory inputs to NTS neurons of animals with intact vagal afferent fibers (2; 3). Since the activity of CCK is supposedly mediated in its entirety by vagal C-fibers (32; 33; 41; 58), chemical
(i.e. perivagal capsaicin) or surgical (rhizotomy) deafferentation should prevent the response of brainstem vagal circuits to application of CCK.

We used the NTS subnucleus centralis (cNTS) as a convenient model for our studies because: 1) the neuronal population of cNTS is quite homogeneous since it comprises almost exclusively sensory neurons from the esophagus (1; 5); and, 2) its rostral location makes it highly unlikely that it receives a significant innervation from the contralateral NTS, unlike the more caudally located subnuclei medialis and commissuralis, which receive a robust input from the contralateral NTS (30).

In the present manuscript, we use different experimental approaches with the aim of characterizing the response of cNTS neurons to exogenous applications of CCK-8s in rats that underwent either chemical or surgical vagal sensory deafferentation.

**METHODS**

Research reported in the present manuscript conforms fully to National Institute of Health guidelines and was approved by the Pennington Biomedical Research Center-LSU System Institutional Animal Care and Use Committee.

*Immunohistochemistry*

Experiments were conducted on 24 Sprague-Dawley rats of either sex (200-250g). One group of rats (N=12) underwent a complete resection of the subdiaphragmatic anterior vagal trunk and surgical removal of the sensory rootlets of the right vagus (supranodose afferent rhizotomy; see below for surgical techniques). The other group of rats (N=12) was used as control, i.e. no surgical procedures were performed. Three days later, all rats were injected with fluorogold (Fluorochrome, Englewood, CO; 20µg/1ml saline/rat, ip) to label preganglionic neurons innervating the subdiaphragmatic viscera allowing delineation of the boundaries of the DMV and, in deafferented rats, to ascertain that the surgical procedures did not damage DMV neurons in those animals that underwent supranodose deafferentation (40; 66). Experiments were conducted 3-5 days after fluorogold injection.

On the day of the experiment, the rats were injected either with saline, 1 or 3µg/kg CCK-8s (0.5ml i.p.). Ninety minutes later, the rats were anesthetized deeply (abolition of foot pinch withdrawal reflex) with isoflurane, and perfused transcardially with chilled saline followed by Zamboni’s fixative (see below). After extraction, the brainstems were stored overnight in Zamboni’s fixative at 4°C. Following wash-out of the Zamboni’s solution, the brainstems were cut in 40µm-thick coronal sections and every third slice mounted onto gelatin-coated coverslips. The slices were incubated at 37°C for 2hrs with the primary antibody (rabbit-α-cFos; 1:5000 in phosphate buffered saline (PBS) containing
0.15% triton-100 (TX) and 0.1% bovine serum albumin (BSA)]. The slices were rinsed with PBS-TX-BSA and incubated again at 37°C for 30 minutes with secondary antibody (goat-α-rabbit Texas red, 1:100 diluted in PBS-TX-BSA). The specimens were again rinsed with PBS-TX-BSA solution containing 1% goat antiserum, before being allowed to air dry, and mounted with Fluoromount-G (Southern Biotechnology Associates, Birmingham, AL). Specimens were coded and c-Fos immunoreactivity (-IR) was counted manually using a calibrated grid by an investigator unaware of the treatment. In each rat we counted, on average, 6.7±1.2 slices from the caudal DVC, 6.5±1.1 slices from the intermediate DVC and 4.4±0.9 slices from the rostral DVC, while 9.3±1.7 slices were used to count the c-Fos expression in cNTS. Photomicrographs were taken using a Nikon E-400 microscope (x100 final magnification) equipped with TRITC (to visualize cFos-IR) and UV (to visualize fluorogold-IR) fluorescent filters and a SPOT Insight® camera and software (Diagnostic Instruments, Sterling Heights, MI, USA) connected to a PC. Overlapping panels of the whole DVC area were digitally enhanced and montaged using ImageJ (developed at the U.S. National Institutes of Health and available from the Internet at http://rsb.info.nih.gov/ij) and Adobe Photoshop® software (Adobe Systems, Inc.).

**Electrophysiology**

The methods of slicing the brainstem and the identification of cNTS neurons have been described previously (3; 4; 59). Briefly, 25-35 day old Sprague-Dawley rats of either sex were anesthetized with isoflurane (abolition of the foot pinch withdrawal reflex) before being killed by severing the blood vessels in the chest. The brainstem was removed and glued to the platform of a vibratome, and three coronal slices (300µm-thick) were cut starting from the posterior area postrema moving rostrally. The slices were stored at least one hour in carboxygenated (95% O₂/5% CO₂) Krebs’ solution (see below) at 30°C before use. A single slice was then transferred to a perfusion chamber (volume 500µl; Michigan Precision Instruments, Parma, MI), kept in place with a nylon mesh and maintained at 35±1°C by perfusion with warmed Krebs’ solution at a rate of 2.5-3.0ml.min⁻¹.

Whole cell recordings were conducted on putative cNTS neurons identified as per their location in close proximity (within 100µm) to the tractus solitarius at a level from the posterior tip of the area postrema to approximately 0.5mm rostral to the anterior portion of the area postrema. Recordings were made with patch pipettes (6-8MΩ resistance) filled with a potassium gluconate solution (see below) by using an Axoclamp 2B amplifier (Axon Instruments, Union City, CA). Data were sampled at 10 kHz and filtered at 2 kHz, digitized via a Digidata 1322 interface (Axon Instr.), acquired with a PC utilizing pClamp8 software (Axon Instr) and analyzed with Mini Analysis software (Jaejin Software, Leonia, NJ).
Rise and decay kinetics of the EPSC were calculated for each event using a subroutine of the Mini Analysis software; the data were then averaged and their mean was compared in the presence and absence of CCK-8s. Recordings were accepted only if the series resistance was < 20\ohm. In the experiments in which the events’ kinetics were analyzed, the series resistance was checked every 10min and the data were discarded if there was a variation of series resistance of more than 25%.

In some experiments (N=19), a bipolar tungsten electrode was placed in the tractus solitarius. Pulses of DC current (up to 2ms long) were delivered while recording from cNTS (HP=-50mV) of rats which underwent vagal deafferentation (see below).

Drugs were applied to the bath via a series of manually operated valves at concentrations shown previously to be effective (3). Neurons were allowed to recover fully between additions of agonists (minimum wash-out period of 10min). Cells were classified as CCK-8s responsive if perfusion with CCK-8s (100nM) increased the frequency of spontaneous excitatory postsynaptic currents (sEPSC) by a minimum of 50% above baseline.

Vagal deafferentation

Both chemical and surgical methods were used to remove vagal sensory inputs to the brainstem (vagal nerve deafferentation). Chemical vagal deafferentation was conducted following methods described previously (23, 41). Briefly, 20 day old rats of either sex were anesthetized with a mixture of ketamine/xyalzine/acepromazine (80/1.6/5mg/ml/kg i.p.). Under aseptic conditions, a ventral midline incision on the neck exposed approximately 4mm of the right cervical vagal trunk, which was isolated over a strip of Parafilm®. A cotton pellet soaked in a 1% capsaicin (CAP) solution (8:1:1 saline: DMSO: ethanol) was applied to the nerve for 30 min; the area was then dried, the Parafilm® removed and the incision closed with 5/0 suture. The electrophysiological experiments were performed 7 to 10 days after the perivagal capsaicin treatment.

Surgical vagal deafferentation was achieved by sectioning the vagal afferent nerve rootlets (supranodose afferent rhizotomy) using a technique similar to that described previously (61). Rats were anesthetized as above and placed in a stereotaxic frame. Following a dorso-lateral incision at the level of the occipital bone, muscle tissue was blunt dissected to expose the occipital bone and the first cervical vertebra; the three supranodose vagal dorsal afferent rootlets are located ~1mm medial to the occipital condyle, the occipital bone needs to be trimmed with a #6 dental drill to expose the vagal rootlets (32, 61). The dorsal rootlets are located beneath the caudal portion of the occipital bone. Once visualized, the supranodose dorsal rootlets on one of the vagal trunks were sectioned under microscopic guidance using a 27 gauge surgical needle. The complete resection of the rootlets was assessed and confirmed routinely by the person assisting the surgery. In one group of rats, 4 days after
vagal deafferentation, the nodose ganglia were exposed by an incision in the neck using an aseptic technique (n=3). The neck muscles were blunt dissected to expose the internal carotid artery and the attached cervical vagus nerve, which was dissected carefully from the carotid artery in the rostral direction. Once exposed, a small (less than 0.3mm) incision was made in the sheath surrounding the nodose ganglion to allow insertion of a micropipette containing a solution (5%) of rhodamine dextrane (lysine fixable, MW 3000). The tip of the pipette was guided through the incision in the sheath of the nodose ganglion, and pressure pulses applied to inject the dye (total injected volume was ~0.1µl). The pipette was withdrawn, the area blotted dry and the cervical incision closed with 4-0 suture. Three-5 days later, the rats were anesthetized deeply (isoflurane 5%) and perfused transcardially with chilled saline followed by Zamboni’s fixative. The brainstem was extracted and postfixed in Zamboni’s fixative overnight at 4°C. Following wash-out of the Zamboni’s solution, coronal sections (40µm thick) containing the DVC were cut using a freezing microtome, and every third slice placed onto gelatin-coated coverslips and mounted with Fluoromount® (Southern Biotechnology Associates, Birmingham, AL) to reduce fading. Confocal microscopic images were collected by using a Zeiss 510 confocal scanning laser microscope equipped with a Kr/Ar-ion laser and filters for the selective visualization of Texas red. Two Z-sections separated by 5µm were taken (final magnification, x100) and their projections merged. Overlapping panels of the entire DVC area were digitally enhanced and montaged using ImageJ and Adobe Photoshop® software (figure 1).

When conducting immunocytochemistry experiments, contralateral subdiaphragmatic vagotomy was performed by removing 3-5mm of the anterior branch of the vagus at a level rostral to the hepatic branch bifurcation or by removing 3-5 mm of the posterior vagal branch at mid-esophageal level. The incisions were closed using 5/0 suture and the rats allowed to recover for 5-10 days prior to experimentation.

**Solution composition**

*Krebs’ for electrophysiology* (in mM): 126 NaCl, 25 NaHCO₃, 2.5 KCl, 1.2 MgCl₂, 2.4 CaCl₂, 1.2 NaH₂PO₄, and 11 dextrose, maintained at pH 7.4 by bubbling with 95% O₂-5% CO₂.

*Intracellular solution* (in mM): K gluconate 128, KCl 10, CaCl₂ 0.3, MgCl₂ 1, Hepes 10, EGTA 1, ATP 2, GTP 0.25. Adjusted to pH 7.35 with KOH.

*Zamboni’s fixative* (in mM): 1.6 % (w/v) paraformaldehyde, 19 KH₂PO₄ and 100 Na₂HPO₄.7H₂O in 240 ml saturated picric acid-1600 ml H₂O; adjusted to pH 7.4 with HCl.

*Phosphate Buffer Solution (PBS)* (in mM): 115 NaCl, 75 Na₂HPO₄.7H₂O, 7.5 KH₂PO₄ and 0.15% Triton-X.
Drugs and chemical

Capsaicin and tetrodotoxin were purchased from TOCRIS Cookson Inc. (Ellisville, MO, USA) and Alomone labs (Jerusalem, Israel), respectively. Fluorogold was purchased from Fluorochrome, LLC (Denver, Colorado, USA); c-Fos antibodies were purchased from Calbiochem (San Diego, CA); Alexa 568-conjugated goat-α-rabbit IgG was purchased from Molecular Probes (Eugene, OR). All other salts were purchased from Sigma (St. Louis, MO, USA). Stock solutions were prepared freshly and diluted to the final concentration in Krebs’ solution just before use.

Statistical Analysis

Results are expressed as means ± S.E.M. with significance defined as \( P < 0.05 \). Results were compared before and after drug administration, with each neuron serving as its own control (Student's paired \( t \) test). Intergroup comparisons were conducted using the Student's grouped \( t \) test or the \( \chi^2 \) test.

RESULTS

CCK-8s increases c-Fos expression in cNTS of deafferented rats.

c-Fos expression was measured in the cNTS following i.p. administration of saline or CCK-8s. Rats were divided in two groups of 12 animals; one group was used as control while the second group comprised rats that underwent surgical vagal deafferentation (i.e. afferent rhizotomy and contralateral subdiaphragmatic vagotomy). Both groups were subdivided in three subgroups consisting of four rats each that received the following treatments: 1) saline; 2) CCK-8s 1µg/kg; and 3) CCK-8s 3µg/kg.

Deafferentation did not affect the baseline c-Fos expression in NTS. In control animals that received saline injections, the whole NTS expressed 126±50 c-Fos positive neurons while the cNTS displayed 16±7 c-Fos positive neurons; in deafferented rats, following saline injection there were 108±42 and 16±5 c-Fos positive cell in the NTS and cNTS, respectively (\( P > 0.05 \) vs control). After injection of 1 µg/Kg CCK-8s in control rats, c-Fos expression increased to 229.5±53.1% of control in cNTS (193.2±44.5% for the whole NTS; \( P < 0.05 \) vs control). In deafferented animals, administration of 1µg/kg CCK-8s increased c-Fos expression to 138.9±30.2% of control in cNTS and 143.1 ± 28.3 % for the whole NTS (\( P < 0.05 \) vs control; figure 2). Injection of 3 µg/Kg CCK-8s in control rats increased c-Fos expression to 359±56.7% of control in cNTS (275±42.0% for the whole NTS; \( P < 0.05 \) vs control). In deafferented animals, administration of 3µg/kg CCK-8s increased c-Fos expression to 251±30.4% of control in cNTS and 281±48.6% for the whole NTS (\( P < 0.05 \) vs control; figure 2). These data suggest that part of the c-fos expression in cNTS neurons induced by peripheral administration of a
physiological (1µg/kg) or a pharmacological (3µg/kg) dose of CCK-8s occurs independently of vagal afferent fiber activation.

Differences were never observed in the CCK-8s induced cFos expression on the right (rhizotomy) vs the left (subdiaphragmatic vagotomy) side (table 1). These data further suggest that the residual c-Fos expression observed in deafferented rats is determined by CCK-8s' activation through routes other than the sectioned vagal afferent fibers or the projections from the contralateral NTS.

**CCK-8 increases spontaneous and miniature postsynaptic current (PSC) frequency in a subpopulation of cNTS cells.**

In neurons voltage clamped at -50 mV, 2-5min perfusion with 100nM CCK-8s increased both the frequency and amplitude of the inward spontaneous postsynaptic currents (sPSC) from 2.08±0.3 to 11.3±1.4 events.s⁻¹ (p<0.05) and from 26.8±0.8pA to 35.4±2pA (p<0.05), respectively, in 45 out of 105 cNTS neurons (i.e. 43%; figure 3). The frequency and amplitude of the sPSC recovered to baseline upon washout. In the remaining 60 neurons, perfusion with CCK-8s changed neither the frequency nor the amplitude of the sPSC (1.6±0.3 to 1.7±0.3 events.s⁻¹ and 24.6±0.7 to 23.7±0.7pA in control and CCK-8s, respectively).

When voltage clamped at -50mV, perfusion of either picrotoxin (40µM) or bicuculline (30µM) with TTX (0.5µM) did not eliminate the PSCs, which, instead, were abolished completely by perfusion with the non-NMDA antagonist CNQX (10µM; N=8; figure 3), suggesting that the PSC are due to glutamate receptor activation and can be classified as excitatory postsynaptic currents (EPSCs).

In 8 cells responsive to CCK-8s, in the presence of either picrotoxin (40 µM) or bicuculline (30 µM) with TTX (0.5 µM), CCK-8s increased the frequency of miniature EPSC (mEPSC) from 1.93±0.74 to 5.8±1.6 events.s⁻¹ (p<0.05) but not their amplitude (27.5±2 and 28.5±2.1pA in control and following CCK-8s, respectively; p>0.05; figure 2). Furthermore, perfusion with CCK-8s did not change either the 10-90% rise time (1.15±0.09 and 1.02±0.08ms in control and following CCK-8s, respectively) or the 90-37% decay time (3.2±0.37 and 3±0.3ms in control and following CCK-8s, respectively) of the mEPSC (figure 4).

Taken together, these results confirm our previous data (3) and suggest that in cNTS CCK-8s can increase the frequency of EPSCs independent of action potential generation. It must be noted, however, that CCK-8s induces a larger increase in sEPSC than in mEPSC (1140.8 ± 341.3 and 357.9 ± 176.4%, respectively, p<0.05), suggesting that part of the CCK-8s' effect is indeed due to action potential generation in the intrinsic circuitry that has been preserved in the slice.

**CCK-8 increases sEPSC and mEPSC frequency in cNTS neurons from deafferented rats.**
In cNTS neurons from rats that underwent perivagal capsaicin treatment 7-10 days prior experimentation, the frequency of sEPSC in cNTS was similar to that of untreated rats, 2.2±0.4 events.s⁻¹ (N=46) and 1.83±0.2 events.s⁻¹ (N=105) in capsaicin-treated and control, respectively. These data suggest that, in the slice preparation, the basal frequency of the sEPSC in cNTS neurons is not determined exclusively by the glutamate released from the central terminals of capsaicin-sensitive C-fibers.

Following perivagal capsaicin treatment, perfusion with 100nM CCK-8s increased the frequency of sEPSC in cNTS neurons from 2.6±1.01 events.s⁻¹ to 10.6±3.14 events.s⁻¹ and the amplitude from 24±1.78 to 41±8.6pA in 9 out of 46 cells (p<0.05). Treatment with perivagal capsaicin reduced significantly the percentage of cells responsive to CCK-8s (20% compared to the 43% of control cells; P<0.05 χ² test). These data suggest that approximately 50% of the CCK-8s-mediated excitatory response is due to actions on the terminals of capsaicin-sensitive vagal-terminals.

Electrical stimulation of the tractus solitarius at current intensity and duration one order of magnitude higher than those previously shown as effective (64) did not evoke any current in cNTS neurons taken from rats that underwent surgical deafferentation 5-10 days prior to experimentation (N=19). These data clearly show the completeness of the surgical deafferentation.

In cNTS neurons from rats that underwent surgical deafferentation 5-10 days prior to experimentation the frequency of sEPSC in cNTS neurons was higher than that of untreated rats, 4.8±1.26 events.s⁻¹ (N=11) and 1.83±0.2 events.s⁻¹ (N=105) in surgically deafferented and control, respectively. These data suggest that, in the slice preparation, the basal frequency of the sEPSC in cNTS neurons is not determined exclusively by the glutamate released from the central terminals of vagal afferent fibers, which, rather appear to exert a tonic inhibitory influence, possibly via activation of local inhibitory circuits.

Following surgical deafferentation, perfusion with 100nM CCK-8s increased the frequency of sEPSC in cNTS neurons from 4.8±1.26 events.s⁻¹ to 11.4±2.14 events.s⁻¹ in 11 out of 28 cells (p<0.05). Similarly, in 7 neurons in which perfusion with CCK-8s increased the frequency of sEPSC, perfusion with 100nM CCK-8s in the presence of 1µM TTX increased the frequency of mEPSC from 2.8±1.1 events.s⁻¹ to 5.1±1.8 events.s⁻¹ in all the cells (p<0.05; figure 4). Six neurons in which CCK-8s did not increase the frequency of sEPSCs were unresponsive even when 100nM CCK-8s was tested on mESPC (data not shown).

These data suggest that CCK-8s does not act exclusively on the terminals of vagal afferent fibers.
DISCUSSION

In the present work we have shown that: 1) surgical deafferentation reduces significantly, but does not eliminate, c-Fos expression in cNTS neurons following systemic CCK-8s; and, 2) chemical or surgical vagal deafferentation does not abolish the ability of CCK-8s to increase the frequency of excitatory synaptic transmission to cNTS neurons.

Taken together these data suggest that CCK-8s activates both capsaicin-sensitive and capsaicin-insensitive vagal sensory afferent terminals in the cNTS, but implies that CCK-8s also excites cNTS neurons via a vagally independent mechanism.

The activation of brainstem vagal circuits by systemic administration of CCK-8s has been postulated as being determined exclusively by its paracrine effects, i.e. CCK, released from intestinal cells in response to a meal, activates capsaicin-sensitive vagal afferent fibers, which then carry the information to brainstem vagal circuits that trigger the efferent vagal response (8; 14; 41). Anatomical evidence certainly supports the presence of CCK receptors on vagal afferent fibers (6) and this was proposed as the sole site of action of CCK because of the observations that, as with a complete vagotomy, C-fiber degeneration induced by perivagal capsaicin treatment greatly attenuates or even abolishes the effects of CCK-8s on gastric motility, gastric acid secretion and pancreatic exocrine secretion (21; 32; 33; 41; 58). One must be aware, however, that the high concentration of capsaicin used in the perivagal treatment (9; 23; 25; 41; 56; 63; 67) may have additional effects unrelated to its recognized actions on C-fibers or neurons receiving direct C-fibers mediated inputs (24; 26; 49). In this regard, a monosynaptic connection between vagal afferent fibers and gastrointestinal-projecting DMV neurons has been described (44), raising the possibility that the lack of CCK-8s effects on gastrointestinal functions in systemically capsaicin-treated animals may be due to non-selective toxic effects of capsaicin on vagal motoneurons. In brief, major drawbacks exist in the use of capsaicin, even when administered perivagally. In consideration of this, we resolved to conduct, whenever possible, either chemical or surgical vagal deafferentation to assess the effects of CCK-8s in cNTS neurons.

The deafferented rats in the immunohistochemical portion of the present study underwent unilateral supranodose afferent rhizotomy and contralateral subdiaphragmatic vagotomy. The rationale for conducting this surgical approach resides in providing a complete deafferentation of the abdominal viscera prior to systemic administration of CCK-8s. If, as inferred, the activation of vagal brainstem circuits by CCK-8s is solely the result of the paracrine effect of the peptide, then complete surgical deafferentation should prevent c-Fos increase in the DVC. A post-surgical period of 5-10 days was
sufficient to allow degeneration of vagal afferent fibers (rhizotomy side) and degeneration of both afferent fibers and motoneurons (controlateral vagotomy side) (28; 29). The fluorogold staining [which is transported centrally by vagal fibers, amongst other pathways (40; 66)] confirmed that the supranodose afferent rhizotomy did not damage the efferent vagus since DMV neurons were labeled only on the ipsilateral side. The contralateral side (both motor and sensory vagotomy) exhibited little, if any, fluorogold staining, indicating the success of the subdiaphragmatic vagotomy.

The increase in c-Fos expression in vagal brainstem circuits following intraperitoneal administration of CCK-8s is well established (18; 20; 35; 45; 46). Following surgical deafferentation, however, we show that systemic administration of CCK-8s still increased c-Fos expression in the NTS including the cNTS. These data suggest that part of the c-fos expression in cNTS neurons induced by peripheral administration of CCK-8s at physiological (1μg/kg) or pharmacological (3μg/kg) doses occurs independently of the activation of peripheral vagal afferent fibers.

Although apparently at odds with previous studies conducted on vagotomized or capsaicin-treated animals (53), one should consider the technical differences between our study and the studies of Sayegh and Ritter (53), foremost the deafferentation procedures. In fact, Sayegh and Ritter did their experiments eight weeks after conducting a complete subdiaphragmatic vagotomy or following systemic administration of capsaicin; conversely, we conducted our experimental procedures five-10 days after afferent rhizotomy and subdiaphragmatic vagotomy. The longer post-surgical recovery period allowed by these colleagues (i.e. 8 weeks) before the experimentation may well have induced a degeneration of vagal afferent fibers as well as NTS and DMV neurons, while our selective surgery and relatively short recovery period is likely to have affected only vagal afferent terminals. Sayegh and Ritter (53) reported that the c-Fos expression elicited by CCK-8s was decreased in vagotomized vs control rats [see for example columns 1 (control saline), vs column 2 (control CCK-8s), vs column 5 (vagotomized CCK-8s), in table 1 of (53)]. They did not compare, however, c-Fos expression in response to CCK-8s or saline in vagotomized rats [see for example columns 4 (vagotomized saline), vs column 5 (vagotomized CCK-8s) in table 1 of (53)]. Indeed, similar to our data, systemic CCK-8s administration induced a (possibly significant) increase in cFos expression also in the vagotomized rats reported in their study (53).

In a more recent report, van de Wall and colleagues (60) demonstrated that while C-Fos expression in response to CCK administration is markedly reduced in capsaicin-treated rats, CCK increases c-Fos expression induced by gastric distention. These data suggest that CCK activates capsaicin-sensitive fibers directly but also enhances the vagally-mediated responses of distention-activated, capsaicin-insensitive fibers that synapse in the NTS.
Our electrophysiological results confirm the excitatory effect of CCK-8s on glutamatergic terminals apposing cNTS neurons reported previously (2; 3). The increase in mEPSC frequency, but not amplitude, and the lack of effect on the EPSC kinetics indicate a presynaptic site of action and suggest that CCK-8s has a profound modulatory role of the glutamatergic inputs to cNST.

In accordance with our cFos data, we observed that perfusion with CCK-8s increased glutamatergic transmission to cNTS neurons from rats that underwent chemical (perivagal capsaicin) or surgical (rhizotomy) deafferentation. These data were somewhat surprising since, as mentioned before, the excitatory effects of CCK-8s on NTS neurons have been ascribed almost exclusively to a paracrine effect of the peptide on capsaicin-sensitive fibers. Certainly, the lower percentage of CCK-8s responsive cNTS cells observed in slices from chemically deafferented rats confirms rather than negates the paracrine effects of CCK-8s, but, at the same time, indicates the involvement of other routes of NTS neuronal activation by CCK-8s. This postulate is reinforced by the data showing that glutamatergic transmission to cNTS is still enhanced when CCK-8s is applied onto cNTS neurons from surgically deafferented rats.

Recent evidence indicates that CCK-8s might also have an effect at sites other than vagal afferent fibers. In fact, the satiating effects of CCK-8s are attenuated by CCK receptor antagonists that cross the blood brain barrier but not by CNS impermeable antagonists (43). Similarly, subdiaphragmatic vagotomy does not attenuate the reduction of food intake induced by pharmacological doses (8nmoles/Kg) of exogenous CCK-8s, but it completely abolishes the reduction of food intake by more physiological doses of CCK-8s (1-2 nmoles/Kg) (42; 43). These reports suggest that higher levels of circulating CCK, which could possibly be attained by prolonged secretion of intestinal CCK, may act at a site outside the abdominal cavity, including the brain. Indeed, electrophysiological studies have also shown that CCK-8s has profound effects on gastric-projecting dorsal vagal motorneurons (65), on NTS POMC neurons in the caudal brainstem (2), on esophageal sensory neurons of the cNTS (3), on isolated cultured sensory neurons of the nodose ganglion (54) as well as in non-identified vagal brainstem neurons (11; 12; 39). In all instances, the threshold concentration of CCK-8s (approximately 1nM), is close to the postprandial levels of plasma CCK (in the picomolar range) (55), and gastric relaxation is observed to occur following systemic administration of CCK-8s at low nM concentrations (58). It is well accepted that, in vitro, the concentration-response curve to agonists is shifted to the right by two or more orders of magnitude (17), thus placing the observed threshold for the response to CCK-8s in the physiological range.

Particular mention must be made regarding a relatively recent manuscript (54) examining, with electrophysiological tools, the response to both capsaicin and CCK-8s in isolated, identified type A
and/or type C nodose ganglion neurons. Both substances were reported to induce a depolarization in nodose neurons. Of particular interest and relevance are the observations that 1) CCK-8s depolarizes similar proportions of A- and C-type neurons; and, 2) only C-type neurons were responsive to administration of low concentrations of capsaicin. Since capsaicin is supposedly selective for C-fibers, these observations, then, fall short of explaining the in vivo results showing that capsaicin, whether administered systemically or perivagally, antagonizes completely the effects of systemic CCK-8s. The results of Simasko and Ritter, however, would support the case of a non-selective action of capsaicin that, by inducing degeneration of fibers other than C-fibers, occludes or attenuates significantly the vagal efferent response, thus leading previous investigators to conclude that vagal afferent fibers only were responsive for the physiological actions of systemic CCK-8s. These conclusions are supported by other reported observations, including our published results, showing that CCK-8s has effects on neurons of the dorsal vagal complex (2; 3; 11; 12; 39; 65) and are strengthened by the immunohistochemical and electrophysiological data reported herein.

Our data then raise the question as to the whereabouts of this source of CCK acting in the NTS. One possibility is that CCK originates from local peptide-containing interneurons; indeed, CCK is one the most widespread neuropeptides in the brain, and there is abundant neurochemical evidence showing the presence of both CCK receptors as well as CCK-containing neurons and terminals in the caudal brainstem (15; 16; 22; 31; 57; 65), suggesting that local neurons containing CCK may also contribute to its effects in vagal brainstem circuits.

Another possibility is that CCK reaches the NTS via its plasma transport. Although it has been suggested that CCK does not cross the blood brain barrier (BBB), we have to consider that the NTS is a circumventricular organ with a leaky BBB, fenestrated capillaries and enlarged perivascular space that allows the passage of large molecules (13; 19; 50; 51), raising the possibility that circulating CCK might also reach these neuronal circuits. Additionally, the adjacent area postrema, which lies entirely outside the BBB, has a series of short, communicating vessels that potentially send postremal venous drainage to the NTS (52). These morphological characteristics raise the possibility that NTS neuronal activity can be modulated by circulating molecules, including CCK. In fact, there is functional evidence that CCK can cross the BBB to activate a solitarius-nigral pathway (27), to phosphorylate CCK-A receptors in the DVC (65) and to induce short term satiety (10).

The reader, however, has to keep in mind that we do not intend to dispute the powerful and well documented paracrine effects of CCK on peripheral vagal afferent fibers. We should like, instead, to suggest that the effects of CCK-8s are due to actions not only on those fibers, but actions at other sites, including NTS neurons, must be taken into account. The cellular versus “in vivo” mismatch reported herein could be due to the different sensitivity of the experimental approaches. In fact, the cellular
approaches we used are capable of discerning the minute changes determined by CCK-8s effects on the membrane of a single neuron. Conversely, the “in vivo” approach, by measuring the overall impact of a treatment, has a lower capability of observing minute variations. One should also keep in mind that the very composition of the vagus, where the vast majority of the fibers are afferent, implies that even minor damage to the efferent fibers induced by perivagal application of capsaicin could not only be overlooked easily, but would also have a potentially major impact on the vagal motor output, leading to an overestimation of the contribution of vagal afferent fibers to the CCK mediated gastrointestinal effects.

In conclusion, our data suggest that, in addition to its paracrine effects on peripheral vagal afferent fibers, CCK also activates vagal sensory afferent terminals in the cNTS, but also implies that CCK-8s excites cNTS neurons via a vagally independent mechanism. It is then possible that the site of action through which CCK influences gastric and pancreatic functions is not limited to its paracrine effect on peripheral vagal afferent fibers but involves also other areas, including NTS neurons.

Acknowledgements

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Table 1. Summary table illustrating the effects on cFos expression of i.p. injections of saline or CCK-8s.

<table>
<thead>
<tr>
<th></th>
<th>Control</th>
<th>Deafferented</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>Saline 1µg/Kg</td>
<td>CCK-8s 1µg/Kg</td>
</tr>
<tr>
<td>NTS Rx</td>
<td>100 215.6±70.21*</td>
<td>262.9±26.40*</td>
</tr>
<tr>
<td></td>
<td>100 213.3±57.40</td>
<td>287.6±55.78</td>
</tr>
<tr>
<td>NTS Lx</td>
<td>100 205.6±37.49*</td>
<td>318.1±42.05*</td>
</tr>
<tr>
<td></td>
<td>100 244.8±59.84</td>
<td>364.1±63.48</td>
</tr>
</tbody>
</table>

Rx= right side (afferent rhizotomy)
Lx= left side (subdiaphragmatic vagotomy)
*P=NS vs Lx
FIGURE LEGENDS

Figure 1. Anatomical proof of deafferentation.

A coronal section (40µm thick) at an intermediate level of the DVC. Note that on the deafferented side of the brainstem, rhodamine dextran labeled vagal afferent nerve terminals are not present within the TS and cNTS, as compared to the dense innervation within the vagally intact brainstem side. Note also that vagal preganglionic motoneurons are labeled in both sides of the brainstem, indicating the selectivity of the surgical deafferentation procedure.

AP= area postrema; DMV=dorsal motor nucleus of the vagus; XII=nucleus of the hypoglossus; cNTS= nucleus tractus solitarius pars centralis.

Figure 2. CCK-8s increases c-fos expression in controls and deafferented rats.

A. c-fos expression following injection of 0.5ml saline in a deafferented rat. Note the prominent fluorogold labeling of the left portion of the DMV and the almost complete absence of fluorogold labeling in the right DMV. B. c-fos expression in a control rat following i.p. administration of 1µg/kg CCK-8s. Note that in this brainstem the intense fluorogold labeling of both sides of the DMV. C. c-Fos expression in the NTS of a deafferented rat treated with 1µg/kg CCK-8s i.p. D. Bar graph summarizing the c-fos expression evoked by CCK-8s i.p. in the whole NTS of control and deafferented rats.

AP= area postrema; DMV=dorsal motor nucleus of the vagus; cc=central canal; cNTS= nucleus tractus solitarius pars centralis. The micrographs were taken approximately at the same level, i.e. +0.5mm from calamus scriptorium.

Figure 3. CCK-8s increases mEPSC frequency via a presynaptic mechanism.

A. Representative traces showing that perfusion with 100nM CCK-8s increases the frequency of glutamatergic mEPSC. HP=−60mV. B. Bar graph showing the percentage of CCK-8s responsive cells in cNTS. C. Averaged traces from the same cell in A in control (36 events) and in the presence of CCK-8s (104 events). The smooth line is one-exponential fitting of the mEPSC decay phase. D. Frequency and amplitude histograms for the same cell in A. Note that perfusion with CCK-8s increases the frequency but not the amplitude of mEPSC. E. Bar graphs showing the averaged results of frequency and amplitude for the mEPSC in control (40µm picrotoxin+ 0.5µmTTX), 100nM CCK-8s and 10µM CNQX (N=8 for all). Note that CNQX completely abolished the occurrence of mEPSC. * p < 0.05 vs control. F. Bar graph summarizing the results for rise and decay time of mEPSC in control and in presence of
100nM CCK-8s. Note that perfusion with CCK-8s does not affect the rise or the decay time of mEPSCs.

*Figure 4. CCK-8s increases sEPSC frequency in rats that underwent vagal deafferentation.*

A. Representative traces showing that perfusion with 100nM CCK-8s increases the frequency of mEPSC. HP=-60mV. B. Bar graph showing the percentage of CCK-8s responsive cells in cNTS. C. Frequency and amplitude histograms for the same cell in A. Note that perfusion with CCK-8s increases the frequency and the amplitude of mEPSC.


Baptista et al. Figure 3

A

B

C

D

E

F

100 ms
50 pA

10 pA
10 ms

Cumulative Fraction

Inter-event Interval (s)

Cumulative Fraction

Amplitude (pA)

Frequency (p.p.s.)

Amplitude (pA)

10-90 % Rise Time (ms)

90-37 % Decay Time (ms)
Baptista et al. Figure 4

A

Control

CCK-8s

Washout

100 ms

50 pA

B

Responsive Cells (%)

100

50

0

C

Cumulative Fraction

Inter-event Interval (s)

1.75  3.5  5.25  7.0

0  0.5  0.5  1.0

CCK-8s  Control  Washout

Cumulative Fraction

Amplitude (pA)

30  60

0  0.5  0.5  1.0

Control  CCK-8s  Washout