Ongoing ingestive behavior is rapidly suppressed by a preabsorptive, intestinal “bitter taste” cue

Lindsey A. Schier*, Terry L. Davidson, Terry L. Powley
Department of Psychological Sciences and Ingestive Behavior Research Center
Purdue University
West Lafayette, Indiana

Running head: intestinal bitter taste aversion

*To whom correspondence should be addressed:
Lindsey A. Schier
Department of Psychological Sciences
703 Third Street
Purdue University
West Lafayette, IN 47907
Phone: (765) 494-6268
Email: LSchier@purdue.edu
ABSTRACT

The discovery that cells in the GI tract express the same molecular receptors and intracellular signaling components known to be involved in taste has generated great interest in potential functions of such post-oral “taste” receptors in the control of food intake. To determine if taste cues in the GI tract are detected and can directly influence behavior, the present study used a microbehavioral analysis of intake, in which rats drank from lickometers that were programmed to simultaneously deliver a brief yoked infusion of a taste stimulus to the intestines. Specifically, in daily 30 min sessions, thirsty rats with indwelling intraduodenal (ID) catheters were trained to drink hypotonic (0.12M) sodium chloride (NaCl) and simultaneously self-infuse a 0.12 M NaCl solution. Once trained, in a subsequent series of intestinal taste probe trials, rats reduced licking during a 6 min infusion period, when a bitter stimulus Denatonium Benzoate (DB; 10mM) was added to the NaCl vehicle for infusion, apparently conditioning a mild taste aversion. Presentation of the DB in isomolar lithium chloride (LiCl) for intestinal infusions accelerated the development of the response across trials and strengthened the temporal resolution of the early licking suppression in response to the arrival of the DB in the intestine. In an experiment to evaluate whether CCK is involved as a paracrine signal in transducing the intestinal taste of DB, the CCK-1R antagonist Devazepide partially blocked the response to intestinal DB. In contrast to their ability to detect and avoid the bitter taste in the intestine, rats did not modify their licking to Saccharin ID probe infusions. The intestinal taste aversion paradigm developed here provides a sensitive and effective protocol for evaluating which tastants—and concentrations of tastants—in the lumen of the gut can control ingestion.

Keywords: chemoreceptor, gastrointestinal, denatonium benzoate, food learning, cholecystokinin
INTRODUCTION

Chemoreceptors are indispensable to the control of food intake. As taste receptors on the tongue and in the oral cavity, they supply signals critical for accepting and rejecting foods for consumption, promoting intake of palatable foods, and preparing the digestive system for the arrival of nutrients. Complementary to this early oral sensory processing, chemoreceptors in the stomach and intestines transduce signals from the lumen to adjust motility, secrete digestive enzymes, and initiate satiation [cf., 24]. Accordingly, the GI tract has been conventionally considered to have a more or less continuous sensory surface for tracking meals to engage and revise apposite and coordinated responses particular to the properties of the food and the body’s physiological state. Despite this arrangement, relatively little work has been done to compare the encoding processes of oral taste receptors with those of the chemoreceptors situated in the post-oral GI tract.

Recently, however, the discovery of taste-like receptors in the GI tract, like members of the T1R (sweet) and T2R (bitter) receptor families, as well as their taste-specific intracellular signaling components (e.g., α-gustducin and TRPM-5) [14, 30, 20, 50, 4, 5, 46, 22, 23, 26] has revealed some parallels among oral and post-oral chemoreceptors and has generated interest in the functional relevance of “taste” receptors in the gut. Such interest has only been reinforced by the mounting evidence that post-oral satiety signals are disrupted in obesity [e.g., 9, 10, 13, 12]. While the anatomical and molecular correspondences have accordingly inspired a rethinking of how chemicals are detected and encoded by post-oral chemoreceptors to affect physiological, and possibly behavioral, controls of intake, these speculations have outpaced experimental analysis.

At least one key complicating factor has commonly impeded investigations of the functional aspects of gut taste. When nutrients are ingested or infused, the specific contributions of the preabsorptive chemoreceptors in the gut are generally confounded with the later postabsorptive consequences, with respect to both within meal feedback and the longer term development of food preferences. Complicating this problem, in many cases, it has also been customary or even necessary to deliver a large stimulus infusion over a prolonged period of time (e.g., 30 mins to several hours) in order to demonstrate overt effects on behavior, but given that almost as soon as the stimulus arrives in the gut, it is transformed and transported, at varying rates dependent in part on the properties of the compound, the stimulus is likely permitted to engage both pre- and post-absorptive pathways during that period. Indeed, feedback from both these sources are accessed and incorporated into ingestive behavior. For example, rats readily establish a conditioned flavor preference (CFP) for an oral flavor that has been repeatedly paired with GI nutrient
infusions [40, 53, 41, 1]. This CFP develops even when vagal and/or splanchnic afferents are surgically or chemically disrupted, suggesting nutrients sufficiently engage post-absorptive signals to affect behavior [40, 53]. At the same time, vagal deafferentation abolishes rapid discrimination of GI stimulus properties for CFP [53] and attenuates the short term satiating effects of some, but not all, nutrients in the GI tract, suggesting auxiliary information is relayed by preabsorptive receptors [49, 40].

In addition to illustrating that both pre- and post-absorptive signaling pathways are involved in the postingestive controls of intake, the CFP example highlights yet another consideration with implications for behavioral analyses of GI infusions. Test subjects appear to be quite flexible with respect to selecting different sources of feedback in order to solve the task. That is, when limited by surgical or chemical intervention to use post-absorptive information in order to associate a flavor with its postingestive outcomes, rats seem to accordingly attend to those features of the stimulus to make appropriate changes in behavior. Consistent with this, intact test subjects may be forced to preferentially use one set of cues (e.g., preabsorptive signals from the intestinal receptors) by simply arranging the demands of the behavioral task to render those (preabsorptive) cues most informative to response decisions [see 33 for a review].

Thus, in light of such experimental issues, the purpose of the present set of experiments was to develop a microbehavioral paradigm to more directly study the stimulus encoding and functional consequences of GI chemoreception. In a modified conditioned taste aversion preparation, rats were forced to use early “taste” cues arising from the intestines to control ongoing ingestive behavior. More specifically, to attempt to assess preabsorptive sensory events, the protocol was designed to (1) probe intestinal chemoreceptive responses to non-caloric, non-readily absorbable tastants presented in simple hypotonic salt solutions, (2) measure short latency responses to tastants as they reach the intestines, thus presumably monitoring events that occur prior to extensive processing of the stimulus within the gut, (3) limit the duration of the taste infusion period within a longer session, in order to track lick response recovery after the stimulus terminates and, thus, in conjunction with a discrete onset, potentially indicating a “stimulus bound” response, (4) render oral cues non-informative, which when combined with the early brief infusion period, forces rats to use those early taste cues arising from the intestine to solve the task, and (5) use an associative approach, i.e. a modified conditioned taste aversion, to amplify responses to weak or innocuous intestinal stimuli, without the need to increase infusion volumes or prolong infusion periods. This
procedure was developed with a representative bitter taste agonist, Denatonium Benzoate (DB), and was then expanded to examine a representative sweet taste agonist, Saccharin (Sacc).

GENERAL METHODS

Subjects. Male Sprague-Dawley rats (Harlan, Indianapolis, IN) were individually housed in hanging wire mesh cages in a climate controlled colony room on a 12:12 hour light: dark schedule (lights on at 06:00). Experiments were conducted in the light phase. Rats were fed ad libitum powdered chow (Purina #5001) and tap water during a one week acclimation period and in a post-surgery recovery period, before being switched to a restricted food and water access schedule for the remainder of the experiment (see Deprivation Schedule). All animal care, surgical, and experimental procedures were approved by the Purdue University Animal Care and Use Committee.

Surgery. Following acclimation to the colony room (1 week), rats were fasted overnight for surgery and then anesthetized with Nembutal (sodium pentobarbital, 60 mg/kg, ip) and laparotomized for implantation of an intraduodenal (ID) catheter. First, a puncture wound was made in the greater curvature of the forestomach, and a Silastic catheter (ID=0.64 mm, OD=1.19 mm, Dow Corning, Midland, MI) was advanced through the stomach and pyloric sphincter. The end of the catheter was tethered to the intestinal wall approximately 4 cms distal to the pylorus with a piece of Marlex mesh and a single stay suture. Then, the puncture wound in the forestomach was closed around the catheter with a purse string suture and a serosal tunnel. The free end of the catheter was tunneled subcutaneously to an interscapular exit site, where it was exteriorized for connection to a Luer Lok adapter, which was mounted in a harness worn by the rat (Quick Connect Harness, Strategic Applications, Inc). Rats were treated with postoperative analgesic and antibiotic.

Apparatus. Daily sessions were conducted in 9 identical operant chambers that were each equipped with a controlled access lickometer and infusion line (Habitest, Coulbourn Instruments, White Hall, PA). The lickometer comprised a sipper spout that was located in a recessed magazine in the center of the endwall, approximately 2 cms above the grid floor and a photobeam that was mounted at the base of the sipper spout to record licks. Each lick was counted and programmed (Graphic State, v3.03) to operate a pump to deliver 5?l of 0.12 M sodium chloride (NaCl) solution to the sipper spout via a polyethylene connection tube. This 5 ?l/lick delivery rate approximated a 1 ml/min consumption rate. Rats had access to the active sipper spout for the entire 30 min session. Throughout all
experimental sessions (as described below), 0.12 M NaCl was the only solution delivered to the sipper spout for oral consumption.

In the final phase of pretraining and then for the remainder of the experiment (see below), a brief ID infusion was added. During the first 6 mins of the drinking session, each lick also operated a second pump and syringe connected to the ID infusion line. The line consisted of polyethylene tubing to a single channel swivel (Instech Solomon, Plymouth Meeting, PA), which was joined to a polyethylene tube encased in a spring tether; this arrangement permitted the rat to move freely about the chamber. Prior to the start of the session, the free end of the spring tether was fastened to the Luer Lok on the rat’s harness to establish connection with the indwelling ID catheter. Direct delivery of the ID infusate (5µls/lick) was yoked to active licking at the sipper spout. Importantly, these yoked infusions were only available for the first 6 mins of the session; once this period elapsed, the ID infusion pump was rendered inactive for the remainder of the 30 mins. The ID infusate consisted of either a 0.12 M NaCl or isomolar lithium chloride (LiCl) vehicle alone, or with a taste stimulus added, depending on the session type (see below). Prior to each session, the infusion lines were thoroughly flushed with 0.12 M NaCl to rinse out any remnants of the previous session’s infusate from the line. The dead space in the line was filled with 0.12 M NaCl (~0.40 mls), and then the line was attached to the syringe containing the ID stimulus; this prevented the rats from being exposed to the ID stimulus as they were connected to the infusion line. Additionally, at the conclusion of the daily session, the ID catheter was flushed with 2 mls of isotonic saline and capped.

**Stimuli and Drugs.** Stimulus properties are shown in Table 1. On all training and test sessions, the rats licked for NaCl, while being infused with NaCl or LiCl. LiCl was mixed in order to achieve the approximate final per session dosage of 8 mg, while matching the molarity of the NaCl. For example, since rats self-infused 5 mls of ID NaCl in pretraining, then 0.12 M LiCl (e.g., 160 mg LiCl in 31.5 ml deionized water) was diluted with 0.12 NaCl (e.g., 480 mg NaCl in 68.5 ml deionized water) for a final LiCl concentration of 1.6 mg/ml. LiCl was delivered at this low dose (8 mg) to prevent rats from shutting down intake altogether following the first exposure to LiCl and to keep rats drinking at a relatively high and stable rate day to day. Denatonium Benzoate (10 mM, DB) or Sodium Saccharin (9.75 mM, Sacc) were added to either the ID NaCl or LiCl, depending on group assignment (see Experiments). All reagents were purchased from Sigma Aldrich (St Louis, MO). All solutions were prepared fresh each day with deionized water and delivered at room temperature. Devazepide (DEV, Tocris Bioscience), a CCK-1R antagonist, was prepared by first dissolving 10 mg in 0.1 ml of DMSO, then adding 0.1 ml of Tween-80,
followed by 0.8 mls of isotonic saline. This 10 mg/ml solution was then diluted to 75 μg/ml by adding isotonic saline, aliquoted in 1 ml vials, and frozen at -20°C. Just prior to administration, DEV was thawed and sonicated. Equivalent volumes of the DMSO, Tween-80, and saline were mixed and stored for use as the vehicle control (VEH).

Table 1 goes here

_Deprivation schedule._ Upon recovery from surgery (~ 2 weeks), rats were gradually accustomed to a water deprivation routine in the home cage; ultimately, on this schedule, rats were given one 30 min drinking session per day (at the same time each day), followed approximately 30 mins later by access to powdered chow and a 10 ml supplement of deionized water for 5 hours. In addition to motivating animals to drink the majority of their daily fluid in a 30 min period, this schedule minimized the amount of food and water in the stomach and intestines during the drinking session (by ~ 18 hour deprivation), while still permitting the rats to gain ~ 2 gms of bodyweight per day. After the initial acclimation to the schedule and for the remainder of the experiment, rats were transferred to a lickometer chamber for the daily 30 min drinking session.

_Pretraining._ Rats were pretrained to drink at the sipper spout for 0.12 M NaCl for 30 mins for 6 days (no ID infusions were made during these initial sessions). This was followed by 8 days in which rats were allowed to drink from the spout for NaCl (for 30 mins) and an ID infusion of 0.12 M NaCl was yoked to licking in the first 6 mins only of each session to establish extensive oral and intestinal experience with the safe salt solution.

_Probe Training._ At the conclusion of pretraining (within Experiments 2 and 3), two day average (last two days of pretraining) bodyweight, lick rate, total licks in the first 6 mins, and total licks over the entire 30 mins were calculated for each rat; then, rats were assigned to groups matched on the mean ±SD on each of these four factors for probe training. Each probe trial consisted of two sessions, which were run on consecutive days (presented in a counterbalanced order across training) and differed only on the type of ID infusate that was delivered; otherwise, these sessions were run in an identical manner as the latter phase of pretraining. In general, for each trial, rats received an ID 0.12 M NaCl infusion on one probe session and an ID 0.12 M LiCl infusion on the alternate probe
session (but note one exception in Experiment 3 below). For some experiments, taste stimuli were added to one or both of these infusates (see Experiments for descriptions of specific pairings). One to three baseline re-stabilization sessions, in which rats licked for NaCl and were infused with NaCl vehicle (no probe tastants added), were interposed between each two session probe trial. Re-stabilization sessions were identical to those described for the latter part of pretraining.

**Statistical analyses.** All analyses were conducted with Statistica (Statsoft, Inc, v10, Tulsa, OK) and data were plotted with Graphpad (Prism 5, La Jolla, CA). For all comparisons, an initial trial by probe infusate by min (Experiments 1-2) or trial by group by min (Experiment 3) repeated measures ANOVA was conducted. Then, to break down significant main effects and interactions, separate repeated measures ANOVAs or post-hoc Newman Keuls tests were run. Because intake was reliably the highest at min 2 on all infusate session types and for all groups, the difference in lick rate between the two probe sessions across the next 6 mins (mins 3-8) was calculated at each trial for each group in Experiments 1-2 in order to examine the development of the early response across training. Additionally, following analyses that compared lick patterns between the two probe infusate session types, separate repeated measures ANOVAs were conducted on each probe infusion session across trials in order to determine changes in response to the particular probe over repeated exposure. An alpha level of 0.05 was used for all analyses.
EXPERIMENTS

Experiment 1: Rats rapidly suppress ongoing intake in response to bitter DB in the intestine

The GI mucosa is the final barrier between the internal and external milieus, where preventing toxins from entering circulation is as important as digesting and absorbing nutrients. Although the latter functions have been the primary focus of GI chemoreceptor research to date, the presence of bitter (T2R) receptors in the post-oral GI tract is consistent with a role for chemoreceptors in defensive GI functions as well [45]. In the gustatory taste system, bitter taste receptors are wired to rejection responses to limit the ingestion of noxious foods [25], but, in the event these foods are ingested, GI T2Rs may provide a second line of defense. Indeed, application of the bitter compound Denatonium Benzoate (DB) directly to GI cells stimulates the release of CCK and slows gastric emptying [8, 18, 17]. In addition to initiating these local reflexes, GI DB conditions avoidance of an oral flavor with which it has been repeatedly paired [17, 19], suggesting (1) that the stimulus has aversive properties in the GI tract and (2) that these signals are integrated centrally, such that the rat avoids the associated flavor upon subsequent opportunities to consume that same flavor. Still, it is unclear whether the bitter stimulus engages a preabsorptive chemoreceptor pathway to affect motility and intake. In fact, in an experiment with some similarities to the present experiments, Glendinning et al [17] suggested that the underlying signal was of a humoral basis, because rats did not suppress intake until approximately 6 mins after the start of the yoked gastric DB infusions and maintained this suppression for an extended period of time. However, because these gastric DB infusions in the experiment of Glendinning and colleagues were continuously yoked to intake across the 30 min sessions, the persistent suppression could reflect repeated preabsorptive stimulation of the bitter receptors, in addition to or even separate from a humoral response. It remains unclear then whether DB is aversive due to its bitter taste properties in the GI tract or some other delayed aversive side effect produced by the compound (e.g., malaise).

Thus, in the procedure developed here, thirsty rats licked at a spout for a safe 0.12 M NaCl solution for 30 min sessions each day, while receiving a brief (6 min) yoked ID infusion of the same salt solution. On probe sessions, the ID infusate contained either 0.12 M NaCl or was replaced with an isomolar LiCl solution. Since at isomolar concentrations, NaCl and LiCl have common early sensory properties (e.g., salt taste), but very different unconditioned effects (i.e., LiCl is a malaise inducing toxic agent) [38], these two salts were ideal candidates for establishing a task in which rats could not use oral cues at the sipper spout (because that solution was always salty plain NaCl) and presumably could not likewise use the early sensory properties of the two infusion vehicles in order
to determine when they would receive an aversive stimulus (i.e., LiCl), unless a specific aversive taste cue (i.e., DB) was added into one of ID infusion vehicles (e.g., NaCl). ID infusions were yoked to drinking for the first 6 mins only of the 30 min drinking session, because (1) voluntary intake is greatest and steadiest in the early part of the daily drinking session—this ensured that the rats self-administered a significant infusion volume at a relatively stable rate, (2) by keeping the infusion to the first 6 mins, rats were forced to use immediate “taste” cues to detect the ID stimulus, rather than delayed effects of the stimulus, and (3) the voluntary intake conditions permitted the rats to control the amount of infusion they received, so that if rats learned to predict the later aversive consequences from the early sensory cues (over the course of 5 exposures), then they could escape or minimize those consequences by discontinuing intake during the infusion period.

In Experiment 1, DB was added as the intestinal taste cue to the ID NaCl infusate on one set of probe sessions, while the ID LiCl infusate was left unadulterated on the interposed, alternative set of probe sessions. While both ID probe stimuli used here (DB and LiCl) are known to be aversive, they are also believed to engage different postingestive pathways. In particular, because LiCl acts primarily by a postabsorptive route to its targets in the CNS (e.g., area postrema) to produce aversive consequences [32,33] and was delivered so as to be difficult to discern on the basis of early sensory properties from safe NaCl [38], it was expected that rats would inhibit intake in the final minutes only of the LiCl probe sessions, reflecting the onset of the malaise. On the other hand, assuming DB activates bitter taste receptors in the GI tract, it was predicted rats would rapidly curb intake in response to the arrival of the bitter stimulus in the intestine. Rats received 5 sessions with each of these aversive probes to examine the development of the response over several trials.

Results and Discussion

Mean licks/min on DB in NaCl and LiCl probe sessions, collapsed across 5 trials are presented in Figure 1. Overall, ANOVA yielded a significant probe infusate by min interaction, $F(29, 174)=6.58, p=0.00001$. Post hoc comparisons indicated that this was due to a lick rate reduction during mins 4-10 of the DB in NaCl session, relative to the alternative LiCl session (largest $p=0.03$ at min 8). This lick rate reduction was transient enough so as not to significantly affect total intake over the entire 30 min session [DB in NaCl: 121.00±30.13 licks/min, LiCl: 141.52±23.80 licks/min], $F(1, 6)=2.51, p=0.16$. The early transient lick rate suppression in response to the arrival of DB developed with repeated probe trials, $F(116, 696)=1.43, p=0.004$. Rats did not discriminate among the two probe infusate types on trials 1-3 [smallest $p=0.13$ on trial 3). On trial 4, rats began to significantly curb licking in
response to the arrival of DB in the intestine (mins 7 and 9, \( p_s=0.02 \) and 0.03, respectively), \( F(29, 174)=3.37, p=0.00001 \). The response was more rapid and robust (mins 4-6, largest \( p=0.03 \) at min 4) on the fifth and final trial of probe training, \( F(29, 174)=5.22, p=0.00001 \). A separate analysis conducted on the DB in NaCl probe session across the five trials confirmed this emergence of the response to DB in the later trials, \( F(116, 696)=2.00, p=0.00001 \). The same analysis was likewise conducted on the plain unadulterated LiCl probe sessions across five trials and further indicated that the response to LiCl infusions did not change from the first exposure to the last, \( F(116, 696)=0.91, p=0.74 \).

These results demonstrated that rats were able to detect the immediate stimulus properties of bitter DB applied directly to the intestine to transiently curb ongoing intake. Detection of DB was quite rapid (within 2-3 mins of the start of session). Interestingly, however, this response did not appear until the fourth exposure to the DB infusion. Prior to the emergence of this early lick rate suppression in the fourth trial, intake remained steady across the first half of the session, before gradually decaying and then completely shutting down in the final mins of the session. This pattern of results could reflect a couple of different underlying mechanisms. For one, mere repeated exposure to the bitter tasting stimulus in the intestine may have resulted in sensitization of the response. Alternatively, it may be that DB produced a delayed onset aversive stimulus (e.g., malaise) in the intestine, independent of its taste properties, and with repeated exposure, rats associated the early sensory properties of the compound with those consequent events, forming an intestinal taste aversion. In other words, the aversive consequence of the DB stimulus may have reinforced the detection of the early intestinal taste cues of DB, which in later trials permitted rats to curb intake more immediately in response to the arrival of the stimulus, minimizing further insult. Rats were not able to use the early sensory properties of LiCl, which were arranged to be indiscriminable from those of NaCl in the intestine, in this same fashion to anticipate impending malaise, consistent with the previous findings of Rusiniak et al [38].
Experiment 2: Taste aversion in the intestine: ID DB in LiCl increases the strength and temporal resolution of the response.

One possible explanation of the development of the response to ID DB across trials observed in Experiment 1 is that rats formed a taste aversion to the bitter stimulus in the intestine, following extensive experience with the aversive compound. Since the unconditioned malaise associated with LiCl has been more fully characterized and since, in Experiment 1, the late-session depression of drinking was clearer for LiCl than for DB in NaCl, Experiment 2 was designed to more directly test the intestinal taste aversion explanation. That is, the purpose of Experiment 2 was to determine if intestinal DB paired with LiCl strengthened the response to the early taste properties of the bitter compound. One group of rats received extensive experience with licking for and being infused with NaCl, before receiving bitter ID DB in 0.12 M LiCl as the probe infusate on 5 sessions. To extend our observations on LiCl, a second group of rats received unadulterated ID LiCl on the same 5 sessions. Plain ID NaCl was the comparison probe infusate for both groups. All other licking and infusion parameters were the same as those described for Experiment 1 (see General Methods).

Results and Discussion

Figure 2A presents mean licks/min on plain NaCl and DB in LiCl probe sessions, collapsed across the 5 trials. Overall, rats consumed significantly less on the DB in LiCl infusion sessions (148.01±25.81 licks/min), as compared to the plain NaCl infusion sessions (203.27±15.51 licks/min), $F(1, 6)=11.85$, $p=0.01$. This was due to a transient, but robust, reduction in licking during mins 6-8 (largest $p=0.0002$ at min 8), $F(29, 174)=2.56$, $p=0.0001$. Although this did not further vary as a function of trial, $F(166, 696)=1.18$, $p=0.11$, because it was expected on the basis of Experiment 1 that rats would develop the response to ID DB across training, each trial was subsequently analyzed. Rats did not discriminate among the two probe infusates on trial 1, $F(29, 174)=0.76$, $p=0.80$. However, by trial 2, a significant suppression in drinking in response to the arrival of DB to the intestine emerged (min 7, $p=0.007$), $F(29, 174)=1.71$, $p=0.02$. This pattern continued to develop over trials 3 and 4 (largest $p=0.02$ for main effect of probe infusate at trial 3). By the fifth and final probe trial, rats rapidly suppressed licking by min 3, completely stopped drinking at min 6 ($p=0.04$), and resumed normal intake (to ID NaCl levels) by min 9, $F(29, 174)=1.84$, $p=0.009$. A separate analysis on the DB in LiCl probe sessions across trials further confirmed the rapid development of the early transient response to the arrival of DB in the intestine across trials, $F(116, 696)=1.31$, $p=0.02$. Although the separate analyses on the plain NaCl infusion probes across trials likewise yielded a significant
trial by min interaction, $F(116, 696)=1.35, p=0.01$, subsequent inspection did not reveal any systematic changes across training for this infusate probe.

Figure 2 goes here

Figure 2B presents the mean licks/min across plain NaCl and plain LiCl ID probe sessions for the second group for which no cues or tastants were added to the infusates, collapsed across 5 trials. Overall, rats consumed comparable amounts on the two probe sessions (NaCl: 177.12±34.07, LiCl:174.90±25.57 licks/min), $F(1, 5)=0.03, p=0.86$. This group did not discriminate among the two infusates during the early minutes of the session, with the exception of a momentary dip at min 11 ($p=0.05$), where responding on the ID NaCl session fell below that of the ID LiCl session; otherwise, these rats exhibited only a late session reduction in drinking, following the LiCl infusion (min 24, $p=0.02$), $F(29, 145)=2.59, p=0.0001$. Even though ANOVA did not return a significant main effect or interaction involving trial (smallest $p=0.53$ for main effect), separate analyses were still conducted on each probe type across training to further determine whether lick patterns in response to the two probe infusates changed with repeated exposure. Indeed, intake patterns did not change in response to either ID LiCl or ID NaCl probes with repeated training, $F(116, 580)=0.91, p=0.72$ and $F(116, 580)=0.86, p=0.85$, respectively. Since it was possible that rats in Experiment 1, who received DB in NaCl and unadulterated LiCl in separate, but consecutive sessions, only failed to detect the early cue properties of LiCl because they were attending more to the (more salient) DB stimulus sessions, we sought to take the competing stimulus out of the equation in this group by presenting these rats with only with unadulterated ID LiCl and NaCl infusions; the results obtained from this group further indicated that rats were unable to discern an early LiCl taste to control ingestion in this task.

Because it was predicted that LiCl would strengthen taste aversion learning to ID DB, we compared the difference in licking in the early mins of the session (mins 3-8) on the two probe session types across trials for three groups (see Figure 3). Whereas the response to DB emerged gradually across probe training for the rats that received DB in NaCl infusions (from Experiment 1), $F(4, 24)=2.36, p=0.08$, LiCl added to the DB infusate accelerated the emergence of the early response (by trial 2 for DB in LiCl group from Experiment 2, $p=0.005$), $F(4, 24)=5.18, p=0.004$. Rats that received plain ID LiCl, opposite plain ID NaCl (from Experiment 2), did not alter licking in the early infusion period across trials, $F(4, 20)=0.35, p=0.84$. 
Although the results from the group that received plain ID LiCl and NaCl on alternate sessions strongly suggest that rats do not readily discriminate intestinal LiCl on the basis of early taste properties of the solution, it remained possible that the enhanced detection of ID DB when it was presented in LiCl (as compared to DB in NaCl in Experiment 1) was due to some interaction among the two stimulus elements of the particular infusate, whereby, for instance, the sensory properties of LiCl acted to enhance the sensory properties of DB or the DB stimulus potentiated the detection of the early sensory properties of LiCl. Therefore, in a post probe training test, ID DB was delivered in NaCl instead of LiCl for the group of rats that were previously trained with DB in LiCl. Even when DB was transferred into an ID NaCl vehicle, overall intake remained significantly reduced (140.72±34.47 licks/min), relative to that on the plain ID NaCl infusion test (227.47±22.61 licks/min), $F(1, 6)=6.49, p=0.04$. Consistent with the lick rate pattern observed during training, rats rapidly inhibited intake in response to DB in NaCl in the intestine ($p=0.03$ at min 10) and resumed normal intake shortly thereafter, $F(29, 174)=1.56, p=0.04$. These results suggest that LiCl did not act as an intestinal DB taste enhancer, but rather merely functioned as a reinforcing stimulus to establish a robust response to the intestinal taste of DB. Furthermore, when DB was removed from LiCl in a separate post probe training test, these same rats failed to discriminate a plain LiCl infusion from a plain NaCl infusion, $F(29, 174)=1.24, p=0.20$, suggesting that DB did not enhance detection of Li taste.

The results from this experiment confirm those of our previous experiment in showing that DB is rapidly detected in the intestine. In addition, we replicated our previous effect with plain ID LiCl in a separate group; intake ceased approximately 20 mins after the LiCl infusion, most likely due to the onset of the adverse unconditioned GI symptoms, but rats were otherwise not able to learn to use the early sensory features of the ID LiCl stimulus to control intake. On the other hand, ID LiCl did augment the development of a response to another tastant, DB, and strengthened the temporal contiguity between the stimulus and response. Thus, interestingly, it appears that rats learn taste aversions to an intestinal taste stimulus, such that those early stimuli in the intestine associate with subsequent events (e.g., malaise) and accordingly modify ingestive behavior upon re-exposures to those same cues in the intestine.
Experiment 3: Rats do not develop a taste aversion to Saccharin in the intestine

Despite the presence of sweet taste receptors (T1Rs) in the intestine [14] and the capacity for non-metabolizable/non-absorbable artificial sweeteners (e.g., Saccharin) to stimulate incretin (e.g., GLP-1) release and upregulate glucose transport [23, 26, 44, 30, 31], other studies have failed to demonstrate a role for GI non-nutritive sweet taste in the behavioral control of intake. For instance, one study showed that rats did not suppress intake following a preload of sweetener to the stomach [51], while a separate study showed that rats did not develop a preference to a flavor that has been repeatedly paired with GI sweetener infusions [41]. Both of these behavioral measures are, however, sensitive to sweet tasting caloric substances, like glucose [51, 49, 40, 41, 1]. Because artificial sweeteners presumably only provide a sweet taste signal (i.e., no calories) in the intestine, it is possible that the typical longer term behavioral parameters used in preload and flavor preference studies are not well suited to look at rapid intestinal taste per se, in the absence of caloric stimuli. Thus, the temporal resolution offered by the current preparation might reveal detection of an artificial sweetener, like Saccharin (Sacc), by examining its immediate effects on ongoing intake. Moreover, it is possible that because artificial sweeteners have relatively few postingestive effects, the stimulus is simply too weak or lacking in salience when presented alone; pairing the sweet stimulus with a strong postingestive event (LiCl) (that lacks any confounding caloric effects) might enhance the detection of a weak Sacc stimulus in the intestine, as was observed for DB in the previous experiment.

Thus, three groups of rats were compared here. As described previously, all groups always licked only for the safe NaCl at the sipper spout, but were infused on discrete probe sessions with different infusates, depending on group assignment. One group received ID Sacc in NaCl, while another group received ID Sacc in LiCl and yet a third group of rats received plain unadulterated LiCl. Since the previous experiments showed robust and consistent lick suppression to ID DB that developed over several trials, all three groups received ID DB (in NaCl) on their alternate probe sessions for comparison. Thus, the first group provided a baseline discrimination between the sweet Sacc and bitter DB in the intestine. Sacc was delivered in LiCl for the second group to establish an intestinal taste aversion to the sweet stimulus. But, in the event that the second group appeared to show detection only after pairings with LiCl, and since the previous experiments indicated that rats do not respond to the early stimulus features of LiCl alone, we included a third group that received unadulterated ID LiCl probes in order to verify that LiCl reinforced earlier detection of Sacc (in LiCl) relative to plain ID LiCl. In the event that rats in the second
group do not detect Sacc, then their response to ID Sacc in LiCl should look as though they received plain LiCl as well.

Results and Discussion

Figure 4 shows the licks/min response on ID Sacc or LiCl probe sessions, collapsed across 6 trials, for each group. An overall ANOVA comparing the licks per minute across the six 30 min ID Sacc probe trials yielded a significant group by minute interaction, $F(58, 493)=1.75, p=0.001$. Direct post hoc comparison among the two groups that received ID Sacc showed that those for which Sacc was mixed in LiCl suppressed drinking earlier than those that received Sacc mixed in NaCl, though this interaction was only marginally significant, $F(29, 319)=1.46, p=0.06$. Whereas the Sacc in NaCl group generally began to suppress their licking around minute 14 ($p=0.01$), the Sacc in LiCl group curbed licking several minutes earlier in the session (at minute 6, $p=0.02$). Accordingly, rats in the Sacc in NaCl group ($119.13\pm18.77$ licks/min) tended to consume more than the Sacc in LiCl group ($87.97\pm14.84$ licks/min), but this difference did not achieve significance, $F(1, 11)=1.70, p=0.22$. Interestingly, however, unlike the pattern we observed for ID DB on its own (presented in NaCl) in Experiment 1 as it developed with repeated exposures, rats did not alter their lick rate pattern in response to Sacc in NaCl across probe trials, [largest $F(5, 20)=2.11, p=0.11$ for main effect of trial]. Neither did the Sacc in LiCl group alter intake in response to the arrival of Sacc to the intestine across training, despite a significant trial by minute interaction for this group [$F(145, 1015)=1.24, p=0.04$], as subsequent analyses did not reveal a systematic change in responding across trials. Even though these comparisons among the Sacc in NaCl and Sacc in LiCl groups seem to suggest that rats for which Sacc was paired with LiCl were able to proactively suppress intake, relative to the group for which Sacc was presented in NaCl, to infer that LiCl reinforced marginally earlier Sacc detection (but notably not as early as DB detection in previous experiments), two observations suggested otherwise. First, as mentioned above, LiCl did not change the response to Sacc across training. Second, additional comparisons revealed that ID Sacc in LiCl did not evidence any advantage above plain ID LiCl. As shown in Figure 4, rats in the Sacc in LiCl group drank slightly more than the plain LiCl group ($71.77\pm8.20$ licks/min) ($p=0.17$), but the pattern of licking was identical across the 30 mins. Post hoc comparisons on a significant group by min interaction [$F(29, 377)=1.54, p=0.04$] indicated one point of divergence among the two groups (at min 10, $p=0.04$), where the group that received ID plain LiCl momentarily dipped below the group that received ID Sacc in LiCl. Furthermore, like the other two groups, those that received ID LiCl did not change their response pattern across exposures, $F(145, 870)=1.14, p=0.14$. 
All three groups received ID DB mixed in NaCl on their alternate session type. As we expected on the basis of the first experiment, all groups evidenced robust lick rate suppressions in response to the early ID DB cues, $F(58, 493)=1.51, p=0.01$, and this developed in a similar manner among groups across probe trials, $F(290, 2465)=1.06, p=0.24$. In fact, all rats consumed less on these alternate ID DB sessions compared to their ID Sacc, regardless of whether it was presented against Sacc in NaCl (83.00±15.33 licks/min) or Sacc in LiCl (57.60±12.12 licks/min), or against plain LiCl (54.62±12.96 licks/min) sessions, $F(1, 17)=14.23, p=0.002$.

The results of this experiment resolved an interpretive issue in Experiment 1. Rats in the first experiment received DB (in NaCl) and plain LiCl on separate, but consecutive, probe sessions within each two-day trial. Even though these rats in Experiment 1 never received direct pairings among DB and LiCl, mere repeated exposure to LiCl induced malaise on intermittent sessions may have arguably sensitized a response to DB. In the present experiment, we included a group that did not receive LiCl; instead, for this group (Sacc in NaCl) both Sacc and DB were delivered in ID NaCl. Since we did not find evidence that rats were able to detect Sacc, then this group essentially received plain NaCl and DB in NaCl on alternate sessions. Their response to DB both within and across probe trials was identical to the group that received, as in our first experiment, plain ID LiCl and DB in NaCl. Taken together, this suggests that unpaired, intermittent exposure to LiCl did not contribute to the response to ID DB (in NaCl) in Experiment 1.

While it was clear that all groups discriminated ID DB from the alternate probe stimulus (Sacc or LiCl) in the present experiment, this appeared to be entirely due to detection of DB, rather than the detection of Sacc or a true discrimination of sweet and bitter. Importantly though, that the response to ID DB was distinguishable from that of ID Sacc indicates that rats were not sensing Sacc as a similarly bitter stimulus in the intestine. However, these results must be interpreted with caution. Given the demonstrated salient stimulus properties of DB, it is possible that rather than it being the case that rats did not sense ID Sacc, that presenting a strong aversive bitter stimulus as the alternative probe impeded learning about Sacc. Future studies will need to better match the relative intensities of the discriminative ID stimuli to avoid this complication.
That said, intestinally infused Sacc’s failure to influence intake is consistent with previous behavioral studies with GI artificial sweeteners [51, 41]. While it is unknown whether the vagus nerve is electrophysiologically responsive to Sacc, artificial sweet tastes do appear to exhibit effects on local GI physiology, upregulating glucose transport and releasing GLP-1 [23, 26, 44, 30, 31]. The ID infusions were made in the proximal duodenum, where the greater part of carbohydrate absorption occurs and is correspondingly densely populated by glucose transporters and presumably their associated sweet taste sensors [37, 52]. On the other hand, GLP-1 releasing L cells are more abundant in the distal small intestine [16], a site that is not likely exposed to our infusion stimulus during the infusion period; alternative infusion sites should be examined. Of course, one final and interesting possibility is that Sacc does modulate local digestive reflexes, but does so via a pathway that is disconnected from other, higher order, responses, like intake. Future experiments will need to re-examine ID Sacc taste with all of these issues in mind.

Experiment 4: Blockade of the CCK-1R pathway partially attenuates the response to ID DB

The results of Experiments 1 and 2 suggested that rats are able to discern the presence of DB in the intestine within approximately 2-3 mins to inhibit intake. The results also suggested that the rats detected the termination of the DB stimulus and recovered intake with a similar latency (especially when DB was in LiCl). The temporal profile of this response strongly suggests DB detection in the intestine is mediated through a chemosensory route, as opposed to a humoral one. Interestingly, however, as mentioned previously, application of DB to GI cells stimulates the release of CCK [8, 18]. While CCK is generally considered a satiety hormone that exerts its effects through both vagal and non-vagal (humoral) routes at various peripheral and central targets [34, 27, 3], it is also believed to convey stimulus-specific information in a paracrine fashion at local sensory afferent terminals [15, 36]. If CCK is involved as a critical chemosensory signaling component, then pretreatment with Devazepide (DEV), a CCK-1R antagonist, should abolish the early response to ID DB, compared to pretreatment with its vehicle (VEH). Thus, following probe training in Experiment 3, rats were tested under these conditions (VEH-DB and DEV-DB). VEH-Sacc and DEV-Sacc tests were included for comparison.

Results and Discussion

Lick patterns across all four test sessions are presented in Figure 5. ANOVA yielded a significant main effect of test, $F(3, 51)=4.05, p=0.01$. Intake on the VEH-DB test (113.46±14.44 licks/min) was significantly reduced compared to the VEH-Sacc (169.18±14.18 licks/min) and DEV-Sacc (163.87±18.79 licks/min) tests ($p$s=0.01 and 0.02, respectively). Furthermore, DEV tended to increase intake on the ID DB test (145.80±19.58 licks/min) compared to the VEH-DB test (113.46±14.44 licks/min)
licks/min), relative to the VEH-DB test ($p=0.09$). These effects of DEV on intake appear to be specific to the DB infusion stimulus, since the antagonist did not (non-specifically) increase intake on the ID Sacc test ($p=0.99$).

There was an overall test by min interaction as well, $F(87, 1479)=3.01, p=0.00001$. Comparable to the probe training pattern of data from *Experiments 1-2*, rats transiently suppressed licking on the VEH-DB test during mins 5-12 (largest $p=0.05$ at min 7), compared to the VEH-Sacc test, $F(29, 493)=5.25, p=0.00001$. DEV appeared to delay this initial lick suppression by ~ 3 mins (different from ID Sacc at mins 8-10, largest $p=0.02$ at min 10), $F(29, 493)=4.52, p=0.00001$. Furthermore, DEV appeared to alleviate the later effects of DB as intake remained slightly elevated in the concluding mins of the test, though direct comparison of the VEH- and DEV-DB tests did not yield a significant test by min interaction, $F(29, 493)=1.18, p=0.24$. Taken together, the data suggest that DEV interfered the response to DB, but did not completely abolish the effect.

*Experiment 5: Rats track and respond rapidly to ID DB presented (unexpectedly) later in the session.*

Although several features of the response to ID DB point to a chemosensory transduction mechanism, since our rats received extensive experience with infusions of DB in the first 6 mins of the session, it was also possible that the rats came to expect with some probability, a DB infusion within those early mins and were, therefore, more prepared to initiate an anticipatory humoral response, upon detection. It was of interest then to determine if these rats that had received extensive prior experience with the early infusion (mins 1-6) tracked and responded to ID DB, when it was administered (unexpectedly) later in the session, after a relatively prolonged licking bout on the safe NaCl only, to then adjust intake. Thus, here, we shifted the infusion period to mins 8-13 for two tests—one in which the ID stimulus was DB and the other in which the ID stimulus was Sacc.

**Results and Discussion**

As shown in Figure 6, rats decreased their lick rate in response to the shifted DB infusion, as compared to the shifted Sacc infusion (significant at min 16, $p=0.01$), $F(29, 348)=1.81, p=0.008$. Thus, rats readily tracked the
quality of the conditions in the GI tract even later in the session to curb ongoing intake, closely replicating the
temporal stimulus response contiguity we observed for the min 1-6 infusion period used in *Experiments 1-2*.

Figure 6 goes here
GENERAL DISCUSSION

The present experiments demonstrated that rats were able to quite rapidly sense the arrival of an aversive bitter stimulus DB in the intestine to inhibit ongoing intake, which minimized further accumulation of the noxious agent in the intestine. The development of this early response to ID DB across probe trials suggested that the longer latency unconditioned aversive effects in the initial trials reinforced the detection of the immediate taste cues during the later trials. Thus, rats developed a taste aversion to DB in the intestine. Consistent with this, use of LiCl as the vehicle for to the DB infusate accelerated the emergence of the early response across trials and produced a more temporally contiguous stimulus-response pattern. That the transient lick suppression was stimulus bound is indicative of an underlying chemosensory mechanism. It appears that CCK is not an essential signaling factor, as Devazepide did not completely abolish the response to DB in the intestine.

In particular, two features of the early lick pattern support the conclusion that DB elicited a signal that was transduced via a chemosensory route. First, following some training, the latency to curb lick rate in response to ID DB was ~2-3 mins, likely preceding the reception of a humoral signal. Because humoral signals must accumulate to a threshold level in the bloodstream and circulate to distant target organs, their effects are generally well delayed from stimulus onset (e.g., latency to affect ingestive behavior, ~5-30 mins after meal instillation) [28, 2]. Moreover, it is not likely that this rapid latency to respond to DB was due to express mobilization of a humoral signal in anticipation of the consequences of the bitter compound, since rats responded with the same latency when the DB infusion was unexpectedly shifted. Second, rats generally only abstained from licking for ~4-5 mins, following DB infusions, before resuming normal intake. This is likewise inconsistent with a humoral basis, for just as these messengers require time to accrue, humoral signals wane relatively slowly; therefore, hormone-driven responses do not tend to reverse so rapidly (e.g., for ingestive behavior, 15 mins to hours, following the meal) [28, 2].

When examined with the discrete infusion procedure used in the present series, as opposed to extended infusions in previous work, the temporal response profile to ID DB is not consistent with the previously suggested humoral basis [17]. Furthermore, our estimates of response latency are particularly conservative for two reasons: (1) infusion lines contained ~0.40 mls of 0.12 M NaCl for the start of each session; thus, the vehicle in the delivery lines and catheter was infused with the first ~80-100 licks of the session, before the DB infusate even reached the
intestine, and (2) under the deprivation conditions used here, the rats, given the opportunity to replete thirst, were perhaps more tolerant of an aversive ID stimulus.

Previously, Hao et al [18] demonstrated that an infusion of a GI bitter taste mixture into the stomach produces cFos activation in the NTS, which is blocked by vagotomy, thus suggesting that these bitter stimuli engage vagal chemoreceptors to relay a code centrally. Additionally, it was proposed that for DB in particular, CCK was a critical factor in the sensory cascade, since Hao and coworkers observed that NTS cFos was completely abolished in CCK-1R null mice and mice pretreated with Devazepide [18]. But whether this pathway is essential for the control of ingestive behavior remains to be determined. Glendinning et al [17] hypothesized that GI DB signals may be relayed by an alternative humoral pathway to reinforce flavor avoidance, though this, of course, does not necessarily preclude CCK as a key humoral factor. Since both cFos activation measures as used in the Hao et al study [18] and conditioned flavor avoidance training parameters as used in the Glendinning et al [17] and Hao et al [19] studies employed long intervals between stimulus arrival and response measurement, permitting a number of different intervening events (both paracrine and endocrine), with non-specific task demands, these procedures may exploit different types of signals.

Hence, in the present study, whether CCK mediated the inhibition of drinking by the ID DB stimulus and lick response was examined. Even though this tightly coupled temporal relationship delineated with the present paradigm seemingly favored a chemosensory/paracrine CCK effect, pretreatment with Devazepide only partially attenuated the early response to ID DB. This partial effect was unexpected given Devazepide completely abolished the cFos response to GI DB in the Hao et al [18] study, but key differences between that study and the present one, including number of exposures to DB (1 versus 6), training history (although there were no training group, in NaCl versus in LiCl, differences with respect to Devazepide blockade in the present study), infusion site (stomach versus intestine), infusion volume (1 ml versus ~4-5 mls), deprivation state (food versus food and water), and species (mouse versus rat), discourage a direct comparison. Whatever the case, the partial effect was not likely due to a subthreshold Devazepide delivery, since we used both a dose and route that have most effectively mitigated the effects of high levels of exogenous CCK and nutrient-induced endogenous CCK, previously reasoned to work through a paracrine mechanism, without producing non-specific effects on intake [6, 11]. That said, it is possible that the dose of DB used here is a more potent stimulator of CCK than has been examined before with respect to exogenous and endogenous CCK, such that a higher dose of Devazepide may be needed to completely abolish the
effect. Alternatively, DB, at the concentration used in the present experiments, may additionally stimulate other non-CCK signaling systems. As such, CCK may signal only one aspect of the DB stimulus, be it bitter taste or some other stimulus property (e.g., irritation), while other features of the stimulus are signaled by factors that are yet unknown. On the other hand, rather than (or in addition to) directly transmitting information about DB, CCK may modulate a separate signaling system responding to the intestinal bitter taste cue. Indeed, CCK has been known to synergistically interact with other peptides, like 5-HT and leptin, in the transmission of post-oral meal related stimuli [21, 35]. Future work will need to determine whether these or other signaling mechanisms are involved in ID DB sensing.

**Perspectives and Significance**

Throughout this report we have referred to DB as a bitter stimulus, simply because it is a selective agonist for a receptor belonging to the bitter T2R family, which has also been localized in the post-oral GI tract, that evokes a similar sensation to other bitter tastants (e.g., quinine) upon contact with lingual receptors [7, 50, 43]. However, it is unclear whether the DB stimulus that the rats are detecting in the intestine is encoded as bitter. For instance, rats may detect and discriminate two ID infusates on the basis of some other cue property, like pH or osmolality, though it is unclear how these particular factors contributed to the discrimination observed here, since rats clearly differentiated between ID DB (in NaCl) and Sacc (in NaCl), two solutions that had near identical pH and osmolality (see Table 1). But even beyond those types of stimulus features, DB may have been sensed or coded as some other type of stimulus (e.g., irritation, anesthetization). Despite some contribution of CCK to the signal, it is not likely that DB produced a satiety-like sensation, given the transient duration of the suppression it produced. Conveniently, though, the procedure developed here can be used to determine on what stimulus quality basis discriminations are made. Indeed, more extensive work with this basic procedure, adapted for discrimination and generalization tests, with a litany of chemicals at varying concentrations, could eventually construct GI taste categories and rule out extraneous cue features. The contributions of specific taste receptors in the GI tract (e.g., T1R and T2R) and primary sensory afferents (e.g., vagal) to these responses with the use of taste receptor knockouts and selective afferent transections, respectively, will be additionally critical to further elucidating GI taste coding.

To our knowledge, the present experiments provide the first demonstration that rats condition a taste aversion in the intestine. At one time, GI chemoreceptors were hypothesized to bridge the perplexingly long temporal delay between the oral tastant and visceral malaise that conventional oral CTAs withstood, by contributing
sensory information about the tastant in closer contiguity with illness [e.g, 39, 29]. The long delay aspect was later explained by other features of CTA, but studies by Tracy et al [47] and Tracy and Davidson [48] supported the notion that information about a post-oral nutritive stimulus (e.g., Polycose) paired with LiCl converges on central taste areas, where it is later accessed when the oral taste receptors sample the same stimulus (Polycose) to, accordingly, inform the rat to avoid its ingestion. Thus, in addition to adjusting the valence of a neutral oral flavor to reflect the respective positive (e.g., caloric) or negative (e.g., toxic) consequences, as in the case of conditioned (oral) flavor/taste-preference (CFP), avoidance or aversion, it appears that chemical features of the post-oral stimulus may be encoded in these types of gustatory associations. Importantly, the results of the present study extend this view by first revealing that a post-oral stimulus is detected and transduced at a preabsorptive receptor site within the intestine, providing a putative pathway by which its chemical features are conveyed. Further, it appears that this intestinal taste stimulus directly associates with consequent events and the signal is accordingly modified and incorporated in the control of intake, as reflected in a change in subsequent responding to same stimulus upon its arrival in the intestine. Taken together, these intestinal taste aversions highlight a more dynamic role for GI chemoreceptors than was previously appreciated [42]. That is, rather than providing comparably crude, inflexible, sensory information to feedback on ingestive behavior, it appears that taste-like signals in the post-oral GI tract are themselves actively revised with experience.

In conclusion, several features of this series of experimentation on DB encourage further use of the present paradigm focusing on early preabsorptive signals in the intestines and, theoretically, the development and use of its appetitive counterpart (intestinal flavor preference), to survey detection and discrimination of other tastants in the GI tract. Considering the close association among chemoreceptors of the GI tract and metabolic and behavioral controls of food intake, an understanding of GI taste categories will likely lead to the development of taste ligands and foods that correct deficiencies in metabolic processes associated with obesity (e.g., glucose homeostasis) and maximize post-oral feedback on satiety to prevent overconsumption.
ACKNOWLEDGEMENTS

The authors would like to thank T. Karam and A. Birge for their assistance with animal care. This report contains part of a Ph.D. thesis by L.A. Schier. A portion of this report appeared in abstract form at the 19th Annual Meeting of the Society for the Study of Ingestive Behavior, 2011.

GRANTS

This work was supported by grants R01 DK-027627 and P01 HD05211 from the National Institutes of Health.

DISCLOSURES

The authors declare no conflicts of interest.
REFERENCES


42. **Smith GP.** The direct and indirect controls of meal size. *Neurosci Biobehav Rev* 20: 41-46, 1996


FIGURE LEGENDS

Figure 1. Mean licks per min for a NaCl solution at the sipper spout collapsed across five 30 min probe trials. A brief (6 min) intraduodenal infusion was yoked to licking at the beginning of the 30 min session (shaded in gray). 10 mM Denatonium Benzoate (DB) in NaCl was infused on one session in each trial and plain unadulterated LiCl was infused on the alternative session in each trial (n=7).

Figure 2. Mean licks per min for a NaCl solution at the sipper spout collapsed across five 30 min probe trials. A brief (6 min) intraduodenal infusion was yoked to licking at the beginning of the 30 min session (shaded in gray). One group of rats (n=7) received 10 mM Denatonium Benzoate (DB) mixed into LiCl infusions on one session per trial and received plain NaCl infusions on the alternative session per trial (A). A second group of rats (n=6) received plain LiCl infusions on one session in each trial and plain NaCl infusions on the alternative session in each trial (B).

Figure 3. Mean ±SEM difference in licks per min in mins 3-8 on the two probe sessions at each trial as a function of training group. For one group rats (from Experiment 1) the difference between licking for NaCl on ID DB in NaCl sessions from the alternate plain ID LiCl sessions is plotted (DB in NaCl). For a second group of rats (from Experiment 2) the difference between licking on DB in LiCl sessions from the alternate plain NaCl sessions is plotted (DB in LiCl). For a third group (from Experiment 2) the difference between licking on plain LiCl sessions from the alternate plain NaCl sessions is plotted (LiCl).

Figure 4. Mean licks per min for a NaCl solution at the sipper spout collapsed across six 30 min probe trials. Rats received a brief yoked intraduodenal infusion during the first 6 mins of each 30 min session (shaded in gray) of 9.75 mM Sodium Saccharin (Sacc) mixed into NaCl (n=5), Sacc mixed in LiCl (n=8), or plain LiCl (n=7).

Figure 5. Mean licks per min for a NaCl solution at the sipper spout on four 30 min test sessions. Rats (n=18) were given an ip injection of the CCK-1R antagonist Devazepide (DEV) (75 µg/ml, 4 mls/kg) or its vehicle (VEH) 30 mins prior to the start of the test session. Rats received an intraduodenal infusion (yoked to licking) of either 10 mM Denatonium Benzoate (DB) in NaCl or 9.75 mM Sodium Saccharin (Sacc) in NaCl during the 6 mins (shaded in gray) of each test session.
Figure 6. Mean licks per min for NaCl at the sipper spout on two test sessions. Rats (n=13) received intraduodenal infusions (yoked to licking) of either 10 mM Denatonium Benzoate (DB) in NaCl or 9.75 mM Sodium Sacc (Sacc) in NaCl, during mins 8-13 of the session (shifted, shaded in gray).
### Table 1. Properties of the Oral Solution and Intestinal Infusates

<table>
<thead>
<tr>
<th></th>
<th>Concentration</th>
<th>pH</th>
<th>mOsm</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>ORAL SOLUTION</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>All Experiments</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>NaCl</td>
<td>0.12 M</td>
<td>6.05</td>
<td>219</td>
</tr>
<tr>
<td><strong>INTESTINAL INFUSATES</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Experiment 1</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>LiCl</td>
<td>31.5% 0.12 M LiCl + 68.5% 0.12 M NaCl</td>
<td>5.94</td>
<td>215</td>
</tr>
<tr>
<td>DB in NaCl</td>
<td>10 mM DB in 0.12 M NaCl</td>
<td>6.72</td>
<td>246</td>
</tr>
<tr>
<td>Experiment 2</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>NaCl</td>
<td>0.12 M</td>
<td>6.05</td>
<td>219</td>
</tr>
<tr>
<td>LiCl</td>
<td>31.5% 0.12 M LiCl + 68.5% 0.12 M NaCl</td>
<td>5.94</td>
<td>215</td>
</tr>
<tr>
<td>DB in LiCl</td>
<td>10 mM DB in 31.5% 0.12 M LiCl + 68.5% 0.12 M NaCl</td>
<td>6.63</td>
<td>249</td>
</tr>
<tr>
<td>Experiment 3</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>LiCl</td>
<td>35% 0.12 M LiCl + 65% 0.12 M NaCl</td>
<td>6.02</td>
<td>215</td>
</tr>
<tr>
<td>DB in NaCl</td>
<td>10 mM DB in 0.12 M NaCl</td>
<td>6.72</td>
<td>246</td>
</tr>
<tr>
<td>Sacc in NaCl</td>
<td>9.75 mM Sodium Saccharin in 0.12 M NaCl</td>
<td>6.89</td>
<td>246</td>
</tr>
<tr>
<td>Sacc in LiCl</td>
<td>9.75 mM Sodium Saccharin in 35% 0.12 M LiCl + 65% 0.12 M NaCl</td>
<td>6.25</td>
<td>235</td>
</tr>
<tr>
<td>Experiments 4&amp; 5</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>DB in NaCl</td>
<td>10 mM DB in 0.12 M NaCl</td>
<td>6.72</td>
<td>246</td>
</tr>
<tr>
<td>Sacc in NaCl</td>
<td>9.75 mM Sodium Saccharin in 0.12 M NaCl</td>
<td>6.89</td>
<td>246</td>
</tr>
</tbody>
</table>

*Note.* The LiCl: NaCl concentration ratio in Experiment 3 was adjusted to account for the lower intake and infusion volume level in that cohort of rats. The final target dose of LiCl was still 8 mg.