Behavioral/Systems/Cognitive

Discrete Innervation of Murine Taste Buds by Peripheral Taste Neurons

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The peripheral taste system likely maintains a specific relationship between ganglion cells that signal a particular taste quality and taste bud cells responsive to that quality. We have explored a measure of the receptoneural relationship in the mouse. By injecting single fungiform taste buds with lipophilic retrograde neuroanatomical markers, the number of labeled geniculate ganglion cells innervating single buds on the tongue were identified. We found that three to five ganglion cells innervate a single bud. Injecting neighboring buds with different color markers showed that the buds are primarily innervated by separate populations of geniculate cells (i.e., multiply labeled ganglion cells are rare). In other words, each taste bud is innervated by a population of neurons that only connects with that bud. Palate bud injections revealed a similar, relatively exclusive receptoneural relationship. Injecting buds in different regions of the tongue did not reveal a topographic representation of buds in the geniculate ganglion, despite a stereotyped patterned arrangement of fungiform buds as rows and columns on the tongue. However, ganglion cells innervating the tongue and palate were differentially concentrated in lateral and rostral regions of the ganglion, respectively. The principal finding that small groups of ganglion cells send sensory fibers that converge selectively on a single bud is a new-found measure of specific matching between the two principal cellular elements of the mouse peripheral taste system. Repetition of the experiments in the hamster showed a more divergent innervation of buds in this species. The results indicate that whatever taste quality is signaled by a murine geniculate ganglion neuron, that signal reflects the activity of cells in a single taste bud.

Key words: taste bud; geniculate; ganglion; innervation; mouse; gustatory

Introduction

The peripheral taste system likely maintains a specific relationship between taste bud cells selectively responsive to one taste quality and the ganglion cells signaling that particular quality. This explains the response specificity of some individual taste nerve fibers (Boudreau et al., 1971; Frank, 1973; Pfaffman et al., 1979; Frank et al., 1988; Sollars and Hill, 2005), particularly because sweet, amino acid, and bitter receptors are expressed in different neuron fibers (Boudreau et al., 1971; Frank, 1973; Pfaffman et al., 1979; Frank et al., 1988; Sollars and Hill, 2005), particularly because sweet, amino acid, and bitter receptors are expressed in distinct populations of taste cells (Chandrashekar et al., 2000; Nelson et al., 2001, 2002). Although anatomical evidence for such an exclusive relationship is lacking at the level of single receptor and ganglion cells, the relationship between single buds and their innervating ganglion cells is tractable neuroanatomically. We explored this receptoneural relationship in the fungiform and palatal taste bud system of the mouse.

Fungiform and palatal taste buds are innervated by the geniculate ganglion cells of the facial nerve. The structural basis for taste quality coding may involve a topographic representation of taste qualities in the geniculate ganglion or the brain. In other senses, a map of the peripheral receptor sheet is represented centrally in an orderly anatomical manner; the peripheral map sometimes exhibits a patterned array of receptors (Tilney et al., 1992). However, in olfaction, the peripheral receptors are represented centrally, not on the basis of topography, but by the spatial convergence of synapses of neurons expressing specific receptors onto central neurons (Mombaerts et al., 1996; Gao et al., 2000; Vosshall et al., 2000). In taste, attempts to identify a spatial representation of taste receptors or taste qualities have revealed only an indistinct functional topography in the brain (Hamilton and Norgren, 1984; McPheeters et al., 1990). Nevertheless, taste ganglion cells must distribute peripheral fibers to particular receptor cell types and disseminate impulses centrally in a structurally organized manner. Previous studies of the peripheral fiber distribution have characterized receptive fields of geniculate ganglion cells or stained fibers in the tongue (Miller, 1971, 1974; Whitehead et al., 1999). These studies of how ganglion cells innervate taste buds have some limitations because of the techniques used; none has analyzed the mouse.

We used iontophoretic injection of retrograde neuroanatomical markers into single taste papillae to label the geniculate ganglion cells innervating the buds (Krimm and Hill, 1998). Our technique used three different color fluorescent lipophilic dyes injected into neighboring buds to identify the number, location, and innervation specificity of ganglion cells innervating single taste buds. Topography and degree of branching among buds of
ganglion cells were investigated through injection strategies targeting different regions of the anterior tongue. Organized patterns are a feature of diverse epithelial appendages, such as teeth, feathers, and hair (Chuong et al., 2000). In the mouse tongue, the fungiform papillae are arranged in a specific patterned array of columns and rows (Jung et al., 1999, 2004). Whether this receptor pattern relates to a corresponding pattern of ganglionic neurons was evaluated. Finally, injecting anterior tongue and palatal buds assessed the topography of geniculate neurons for their two major receptor fields.

**Materials and Methods**

**Animals.** One hundred C57BL/6 mice and five Golden Syrian hamsters 6–16 weeks of age were used. All laboratory procedures were approved by the University of California at San Diego Laboratory Animal Care and Use Committee and followed the National Institutes of Health Guide for the Care and Use of Laboratory Animals.

**Dye preparation.** Lipophilic dyes [Fast DiI (D-3899; absorption, 568 nm; emission, 564 nm), Fast DiO (D-3898; absorption, 488 nm; emission, 499 nm), Fast DiD (D-7757; absorption, 647 nm; emission, 663 nm) (Invitrogen, Eugene, OR)] were dissolved at a concentration of 3–5 mg/ml in a 1:1 solution of DMSO and ethanol. Five to 10 μl of the dye solution were loaded into glass capillaries (25 μm tip diameter) and secured on a micromanipulator (Fine Science Tools, San Francisco, CA). Dil readily went into solution, whereas DiO and DiD solutions were sonicated, allowed to sit overnight, and centrifuged before the experiment. Because of the poor solubility of DiD and DiO and their low compatibility with each other as well as with Dil, care was taken that dye mixtures were thoroughly mixed and spun down immediately before injections.

**Papillae numbering and iontophoresis.** Animals were anesthetized with a 40 mg/kg intraperitoneal injection of pentobarbital, after calming with Isoflo (Abbott Laboratories, Chicago, IL) vapors. The dorsal half of the tongue was gently pulled out of the mouth and immobilized by pressing its ventral surface against the sticky surface of a double-sided tape attached to a tongue platform. Fungiform and palate papillae were visualized with the aid of 0.5% methylene blue solution (Fisher Scientific, Houston, TX). Using a surgical microscope (Zeiss, Oberkochen, Germany) and micromanipulator, the taste papilla was selected and firmly capped with a glass pipette (150 μm thick) before iontophoretic injection. For the palatal bud injections, the animal was placed in the supine position and both the jaws and the cheeks were stretched apart using nontraumatic hooks. The taste pores were visualized with the aid of 0.5% methylene blue solution. The dye-filled capillary was lowered over the pore and pressed snugly to cover and seal the pore and pressed snugly to cover and seal the Figure 2. Injecting the same taste bud in five different animals labels a small group of scattered cells in the geniculate ganglion (dashed outline). **A,** The injection site for each animal was CV-5 (star in **A**). **B–F,** Labeled cells in the geniculate ganglion numbered 2–7 by animal; there was no consistent location of the cells across animals, notwithstanding the same injected bud. The asterisk marks the GSP nerve, whereas the + symbol marks the central (toward brain) corner of the ganglia. Vlfn marks the facial nerve.

Scale bars, 50 μm.
buds were defined as the set of labeled cells on a line running parallel to the midline of the tongue. The column closest to the midline was named CI (C for column and roman numeral one), and CV (column 5) is the farthest column from the midline (see Fig. 1). At the anterior tip of the tongue, all columns turn downward and backward ventrally. Numbering of column papillae started with the anterior-most bud (last bud on the ventral surface) and ended with the papilla at the lower margin of the intermolar eminence. For example, bud CI-10 (column 1, bud 10) is the 10th bud of the column nearest to the midline starting from the first bud on the ventral surface of the tongue (see Fig. 1). Rows RI–RV correspond to five buds of column 5 (CV). RV-1 is the first papilla of row 5 that is coincident with the last papilla of column CV-5. Rows RIII, RIV, and RV were the most accessible rows for injections.

**Tissue histology and microscopy.** After a postinjection survival time of 48 h, the animals were anesthetized and perfused with 4% paraformaldehyde in normal saline. Tongues and ganglia were removed and left in 4% paraformaldehyde for 2 h, which was then followed by PBS or 30% sucrose (for tissue that was sectioned) infiltration overnight. Whole ganglia were stained with the Nissl green (Invitrogen), a neuronal cell marker, to confirm the identities of the dye-filled cells. All labeled cells were found to be neurons, not glia (see Fig. 1 E).

**Image analysis.** All 50 confocal images (corresponding to each dye) were passed for 4–6 min (BAB-500 iontophoresis pump; Kation Scientific, Minneapolis, MN). The negative terminal of the current source was made into multiple papillae along rows and columns. Care was taken not to spill the dye on the adjacent taste buds. A cyclic current of 4 μA with 5 s on–off pulses (square anodal pulses) were passed for 4–6 min (BAB-500 iontophoresis pump; Kation Scientific, Minneapolis, MN). The negative terminal of the current source was attached to the external ear (pinna) of the animal. Similarly, depending on the nature of the experiment, additional papillae were selected and injected with the same or different dyes. In some animals, injections were confirmed the identities of the dye-filled cells. All labeled cells were found to be neurons, not glia (see Fig. 1 E).

**Figure 3.** Injecting three adjacent buds with different color markers results in three separate groups of labeled neurons in the geniculate ganglion. A. Three adjacent buds, RIV-1, RV-1, and RV-2, were injected with DiD (blue), DiO (green), and DiI (red), respectively. B. Labeled ganglion cells from three buds in A, viewed with all three fluorescence settings, are predominantly single labeled and scattered in the ganglion. A double-labeled cell from RIV-1 and RV-1 injections is marked by the cyan circle (same cell C–E). Cells innervating each bud imaged separately with the appropriate laser settings. A asterisk marks the GSP nerve. VIln marks the seventh cranial nerve. Scale bars, 100 μm.
were superimposed and flattened (along one axis) into one final image. This merged image was pseudo colored and overlapped (merged transparent) with other color images (corresponding to two other dyes) using Adobe Photoshop (Adobe Systems Incorporated). Cells were colored red, blue, and green as a result of labeling with DiI, DiD, and DiO, respectively. Likewise, confocal optical series of images were also analyzed along the x- and y-axes. Colocalization of dyes in a single neuron was affirmed by studying the merged images at all three dimensions (see Fig. 6B).

*Papillae counts.* Mouse tongues, previously fixed with 4% paraformaldehyde, were dissected and stained with 0.5% methylene blue to enhance the visual contrast between the taste papillae and the surrounding filiform papillae. The stained tongue was then studied under the surgical microscope. Maps of the distribution of taste papillae were drawn. The upper margin of the side of the intermolar eminence was chosen to be the starting point, whereas the extreme tip of the tongue was the end point. Papillae on the underside of the tip were mapped separately by flipping the tongue upside down. Because a few taste papillae on the dorsal surface and most on the ventral surface of the tip lack their typical morphology, they were difficult to visualize and count. Therefore, tongues from transgenic mice expressing green fluorescent protein (Gy13 GFP) were used for taste bud counts. In these animals, Gy13 promoter was used to drive GFP in taste receptor cells present in most taste buds. This allowed buds to be readily identified. These counts were used to ascertain the reliability of bud numbers and patterns seen in wild-type animals. GFP-expressing papillae were counted using a fluorescence dissection microscope. The same tongues were later stained with 0.5% methylene blue and recounted. The results were consistent with counts of GFP-fluorescent buds. Similarly, buds on the surface of the soft palate and inside the nasoincisor duct of Gy13/H9253 were readily identifiable. These counts were used to ascertain the reliability of bud numbers and patterns seen in wild-type animals. GFP-expressing buds. Similarly, buds on the surface of the soft palate and inside the nasoincisor duct of Gy13/H9253 were counted. The results were consistent with counts of GFP-fluorescent buds. Similarly, buds on the surface of the soft palate and inside the nasoincisor duct of Gy13/H9253 were counted and mapped. Palate counts do not include buds on the epiglottis.

*Geniculate neuron (innervating the anterior tongue) counts.* To count taste-specific neurons in the geniculate ganglion innervating the entire anterior tongue unilaterally, the chorda tympani nerve was exposed in the middle ear by removing the tympanic membrane in anesthetized animals. The nerve was cut, and the central stump was painted with the mixture of three dyes (DiI, DiD, and DiO). After a 48 h survival time, ganglia were removed and imaged under the confocal microscope. Labeled cells, corresponding to each dye, were counted from digitally magnified images.

*Ganglion mapping.* Outer margins of the triangular ganglion (merged along the z-axis) were traced, and the midpoints (x, y, and z) (see Fig. 10C) were connected to each other. The internal triangle thus formed was termed "I" (or internal) triangle. The triangle toward the greater superficial ganglion (GSP) corner was termed "G," the triangle toward the peripheral corner was termed "P," and the triangle toward the central corner was termed "C" (toward the brain). Labeled cells in each triangle were mapped and counted separately for each color.

**Results**

**Single taste bud injections.** Fungiform papillae bearing taste buds are arranged in linear arrays on the mouse tongue (Jung et al., 1999, 2004). Extending anteriorly from the intermolar eminence, a series of five parallel columns of buds can be easily recognized on each side of the tongue (Fig. 1A, B). The anterior-most buds of these columns wrap around the tip of the tongue onto its ventral surface for a short distance. Perpendicular to the columns, the buds are arranged in loosely aligned rows. Counting from the periphery toward the midline, and from the tongue tip posteriorly, identified buds for injection (see Materials and Methods). Identified buds, sometimes the same bud in different animals, were targeted for iontophoretic injections of fluorescent tracers in the present series of experiments. Injection parameters were standardized to achieve optimum labeling of the bud without any spillage into the neighboring buds. Tongue histology confirmed that injections were confined to the taste bud (Fig. 1D).

**Injecting lateral bud CV-5, a readily accessible and consistently identifiable bud, with the retrograde tracer DiI resulted in retrograde labeling of a small number (two to eight) of geniculate ganglion cells. These cells were randomly scattered in the ganglion. Injecting the same bud in five different animals revealed no consistent ganglionic location of the few labeled cells (Fig. 2). After a single bud injection, the average number of labeled cells per geniculate ganglion was four (SD, 1.85; n = 22).

**Multiple taste bud injections.** Three adjacent buds were injected to test whether a small patch of receptors was represented by a concentration of labeled cells with a discernible topography within the ganglion. The buds were injected with different color markers (DiI, DiD, and DiO) to identify the ganglion cells innervating each of the buds. The three injected buds were chosen randomly from each animal; all were near the tongue tip. Cells labeled by anterograde transport again were scattered throughout the ganglion (Fig. 3). Varying the locations of the injections did not reveal any pattern in the representation of patches on the tongue within the ganglion. Remarkably, the average number of ganglion cells labeled after injecting three buds (17 ± 1.29 SD; range, 15–18; n = 5) was quadruple the average number labeled after injecting a single bud (4 ± 1.85). The majority of ganglion cells were single labeled; only rarely were cells double labeled (see Fig. 11A). Cells in only two instances (3.6% of total cells) were triple labeled. Across all cases, including ones with many buds injected (see below), there was a close linear relationship between the number of injected buds and number of labeled ganglion cells. The number of neurons were typically multiples of four to five of the number of buds (Fig. 4). For example, injecting 15 buds label 58 (SD, 3.59; n = 7) and 25 buds label 95 (SD, 13.86; n = 6) ganglion cells.

Several types of control experiments tested for the relative effectiveness of the three fluorescent markers and for possible collateral branches of ganglionic fibers that might have escaped detection with the three bud injections. Injecting a single bud with a mixture of DiI, DiO, and DiD retrogradely labeled geniculate ganglion cells, in this case with similar effectiveness; all cells...
were triple labeled (Fig. 5A–E). In most repetitions of this mixed dye injection, some multiply labeled ganglion cells contained variable amounts of each of the three markers, probably reflecting the physical chemistry of the dye mixture and possible limited miscibility of the reagents at the injection site. An additional control, on a bigger scale, labeling the chorda tympani nerve with the three markers, retrogradely labeled the majority of geniculate ganglion cells; all were triple labeled (Fig. 5F–L). Despite the close proximity of the buds in the three neighboring bud-injection cases (and the possibility that fibers innervating one bud might also innervate its neighbor by a branch), only once were double-labeled cells seen. To rule out the possibility of a thin collateral branch, harboring very little dye, and missing detection (weak fluorescence emission) or to unmask uncommon doubly innervated buds, all buds in two concentric rings (20+/H11001 buds) were injected first with one dye (DiI), except for the central bud, which was injected with a second contrasting dye (DiO) (Fig. 6). In every case, three to five ganglion cells were labeled with the second marker, whereas a vast majority was labeled only with the first marker (see Fig. 11D). Thus, collateral branches, even if sparse, were rare. Strikingly, in the typical case, only one cell (rarely two) (Fig. 6C–E, arrows) was double labeled. All of the other cells were purely single labeled (circled green in Fig. 6B–H). This indicates that only a fraction of ganglion cells send collaterals to multiple surrounding buds, whereas most of the neurons exclusively innervate one bud.

Specific innervation of buds by ganglion cells that do not branch to innervate neighboring buds was also confirmed by injecting identified, parallel rows of buds, each with a different fluorescent marker. Again, retrogradely labeled ganglion cells, with rare exceptions, bore only a single marker (Fig. 7). Double-labeled cells that were seen bore the marker of adjacent rows, never that of separated injected rows (see Fig. 11C). There was no apparent topography, no evidence that the stripes were represented by patterns in the ganglion. A similar approach was used to

**Figure 5.** A–E. Injecting a single bud (RIII-2) with an equimolar mixture of the three dyes used in the present study (DiI, DiD, and DiO), a control for the equal effectiveness of dye uptake, labeled all of the same ganglion cells with every marker. A. Injection site. B. Merged image of all three colors showing all triple-labeled cells (white containing). The pink cast of some cells reflects the variable intensity of labeling because of physical chemistry of dye miscibility at the injection site. The blue line and green patches are artifacts. C–E. Separate images of each marker viewed with the appropriate fluorescence settings. F–L. All chorda tympani nerve fibers innervating taste buds (F) were labeled with an equimolar mixture of the three dyes. G. Most ganglion cells in the merged image of all three colors were triple labeled. H–J. Separate images of each marker viewed with the appropriate settings.
compare the distribution of labeled ganglion cells after injecting columns in different locations versus rows, with no apparent topography (Fig. 8).

**Bud and ganglion cell numbers**
Counts of fungiform taste buds determined that an average of 52 ± 3.62 buds (n = 12) are present on each side of the anterior tongue and 60 ± 9.4 buds (n = 11) are present on each side of the palate (see Fig. 10 B). Counts of geniculate ganglion cells determined that an average of 213 ± 6.24 cells (n = 3) innervate the fungiform field on each side of the anterior tongue. Thus, ganglion cells outnumber the fungiform buds they innervate 4–5:1.

**Palate bud injections**
Adjacent buds in the soft palate were injected with contrasting markers to test for innervation specificity and topography of ganglion cells for comparison with those innervating lingual buds. As with lingual buds, individual buds in the palate are innervated by small groups of five to eight ganglion cells (average, 6.41; SD, 1.16; n = 11) that exclusively innervate a bud but not its near neighbor; the neighbor is innervated by a separate group of ganglion cells (Figs. 9A–E, 11 B). One pattern in the distribution of ganglion cells innervating the palate that was apparent in every case of adjacent bud injection was their peripheral concentration within the ganglion, especially near the point where the GSP exits the ganglion. Injecting many buds in the palate and many buds on the anterior tongue with a contrasting marker established that ganglion cells innervating the tongue were prevalent in the area of the ganglion adjacent to the facial nerve, surrounded by a complimentary distribution of ganglion cells innervating the palate that were concentrated near the GSP (Fig. 9 F, G).

**Preferred distribution of ganglionic cells innervating palate and fungiform papillae**
Counts of geniculate cells in four equal and precisely defined sectors of the ganglion showed that neurons innervating fungiform buds predominate (47%; SD, 8.99; n = 5) in the area of the geniculate that lies laterally near the entry point of the chorda tympani fibers (“P,” peripheral stump of facial nerve VII). In contrast, >40% (SD, 16.98; n = 5) of the total cells connecting the palate taste buds concentrate rostrally near the GSP exit (“G”) (Fig. 10C,D). Notably, G has only 16% (SD, 12.99; n = 5) of the total cells innervating the front of the tongue, whereas P has only 8% (SD, 9.23) of the total palate cells. The internal and central domains showed no preference and were populated equally by both palatal and lingual neurons.

**Hamster bud injections**
A control for failure of collaterals to label because of possible technical problems encountered in the three bud injection experiments was to label three adjacent buds in a species with ganglion cells known to have collaterals (e.g., the hamster) (Whitehead et al., 1999). The result was a higher incidence (23 cells or 35.3% of total labeled cells) of double-labeled geniculate ganglion cells in hamster, in contrast to the mouse (6 cells or 5.4% of total) (Fig. 12). The numbers of double-labeled cells were significantly increased (p < 0.001 in the hamster (n = 5 ganglia; 4.60 ± 1.67 SD) compared with the mouse (n = 7 ganglia; 0.86 ± 0.9 SD)).
Discussion

Bud units in mouse

The principal finding made after injections of single murine buds with contrasting markers is that each bud, whether on the tongue or palate, is innervated by a small group of ganglion cells that innervate single buds exclusively. This discrete innervation pattern for the mouse, with predominantly unbranched ganglion cells, contrasts with reports of branching nerve fibers in the anterior tongue reaching several buds in rats and hamsters (Miller, 1971, 1974; Whitehead et al., 1999). This discrepancy could reflect true differences between species or technical differences between studies. In the rat, collections of silver-stained nerve fibers were described as branching below the taste bud (Miller, 1974). However, with silver staining it is difficult to resolve individual fibers or their branches. Electrophysiological study in rats found that single fiber responses to chemical stimulation of one papilla could also be obtained by electrical stimulation of nearby papillae (Miller, 1971) (<50% of fibers responded exclusively to one papilla). In cats, single geniculate ganglion cells responded to electrical stimulation of single buds in only one-fifth of tested neurons; four-fifths responded to two to five nearby buds (Boudreau et al., 1971). More recently, a similar result was obtained in sheep using more refined methods (Nagai et al., 1988). Thus, species differences in the selectivity of bud innervation by single ganglion cells seem likely.

Miller (1971) referred to the single afferent taste fiber and the papillae that supply its input as a “peripheral nerve fiber unit.” We have adopted a modification of this phrase, termed here as “bud unit,” to name the predominant recepтонeural arrangement in mouse taste. A bud unit is a taste bud and the small group of ganglion cells that exclusively innervate it. A mouse’s bud units are remarkably discrete; only rarely do geniculate ganglion cells innervate two or more buds. Injecting adjacent buds in a mouse with different color dyes did not yield significant numbers of double- or triple-labeled ganglion cells. Moreover, control experiments in which one bud was injected with one marker, whereas all other surrounding buds (as many as 20–22) were injected with a contrasting marker, did not increase the number of double-labeled cells. Thus, the discrete innervation of mouse buds appears certain. A recent study of hamster fungiform bud innervation using iontophoretic application of tetramethyl rhodamine dextran to single buds labeled nerve fibers and their endings that terminated in three to five buds nearby the injected bud (Whitehead et al., 1999). This finding was interpreted as indicating that the marker labeled collaterals of fibers innervating the injected bud. If so, then the innervation pattern for the hamster would be expected to be more diffuse than that in the mouse. This was confirmed in the present control injections of three adjacent buds in hamsters, a repetition of the experiments done in mice. The marked increase in double-labeled hamster ganglion cells verified the significant incidence of branching of ganglion cell peripheral fibers in this species. Moreover, as a technical control, the increased multiple labeling in hamsters indicates that when collaterals are present they can successfully transport the lipophilic dyes to the ganglion.

Figure 7. Injections of three adjacent rows of papillae with different color markers in attempts to reveal a corresponding topographic pattern in the geniculate ganglion. A, All five to six papillae of rows RIII, RIV, and RV were injected with DiO, DiI, and DiD. B, Merged image of all three colors shows apparently dispersed and random distributions of labeled cells; in this case, only one cell was double labeled (arrow). C–E, Individual images of each marker that were combined in B. The side panels show the confocal z-axis series merged along the y-axis (horizontal) and x-axis (vertical). F–I, Four additional cases showing merged views of all three markers after injections as in A. The few double-labeled cells were either yellow or pink (i.e., all double-labeled cells bore the labels from two adjacent rows only). The facial nerve (VII) is at the bottom of each figure. Scale bars, 100 μm. The asterisk marks the GSP nerve, whereas the + symbol marks the central (toward brain) corner of the ganglia.
Innervation of Mouse Taste Buds

Numbers of ganglion cells innervating single buds

An average of four to six geniculate ganglion cells innervate single taste buds in the mouse, whether on the anterior tongue or palate. This number suggests that fewer ganglion cells innervate fungiform taste buds in mice than in other species [e.g., rats (2–16) (Krimm and Hill, 1998) and hamsters (5–35) (Whitehead et al., 1999)]. Remarkably, however, in hamsters, when the number of geniculate ganglion cells innervating the tongue is divided by the number of fungiform buds, the result is four. Thus, if peripheral fiber branching were not present in hamsters, the ratio of ganglion cells to bud would be the same as in mice. In rats, the numbers of geniculate ganglion cells innervating single buds varies from 2 to 16 depending on the bud volume and the age of the animal (Krimm and Hill, 1998). Notwithstanding the difficulty tracing silver-stained fibers between sections and through purported branch points, all of the drawings of fibers innervating fungiform papillae in rats show, consistent with the present results, three to seven fibers per bud (Miller, 1974).

The small number of ganglion cells that innervate mouse taste buds has several implications for taste information coding and transmission to the brain. First, because the number of receptor cells in a bud (50–150) (Kinnamon, 1987) outnumber the innervating neurons by a factor of 10–30, the fiber of each ganglion cell either branches to innervate many cells within the bud, or it innervates only a small subset of available cells, at least at any time point. The latter possibility is supported by ultrastructural analysis of serially sectioned taste bud cells (Kinnamon et al., 1993; Royer and Kinnamon, 1994). Second, whatever the synaptic partners of the small group of innervating ganglion cells, the scarcity of collaterals as presently defined in mice requires that the electrophysiological responses of single primary neurons will usually reflect the activation of a single bud. Finally, the small number of innervating neurons may bear some relationship to the variety and small number of specific types of receptor cells in a single bud, some of which express a distinct heterodimer of two receptor proteins for sweet or umami taste, or a family of bitter receptors for bitter taste (Chandrashekar et al., 2000; Nelson et al., 2001). For example, cells expressing bitter receptors are distinct from those expressing sweet and number only a few per bud (Nelson et al., 2001). A caveat, however, is that the receptors for salt and sour tastes are not fully understood. Neither can coexpression of these latter receptors with bitter, sweet, and umami receptors be ruled out (Caicedo et al., 2002). Moreover, some single geniculate neurons are broadly sensitive to sodium salts, nonsodium salts, and acids. Nevertheless, in light of the presently reported relationship of four to five ganglion cells to one bud, we can speculate that each neuron may transmit information related to only one of the four to five commonly recognized taste qualities. Although the present data on the bud–ganglion cell wiring does not bear directly on taste quality coding, it is consistent with such a hypothesis and with the specificity of responses recorded from some ganglion cells and their peripheral fibers. On the other hand, the present results cannot rule out the possibility that branches of single fibers may synapse with several different types of receptor cells within a single bud, accounting for broad sensitivity (Boudreau, 1971; Frank, 1973; Pfaffmann et al., 1979; Frank et al., 1988; Lundy and Contreras, 1999; Breza et al., 2005; Sollars and Hill, 2005).

Taste bud–geniculate ganglion topography

Many patterns of buds on the tongue were labeled to test for a possible spatial representation of fungiform buds in the geniculate ganglion; none was apparent. Specific regions of buds on the tongue do not appear connected to cells in certain regions of the ganglion. Rather, in every peripheral pattern attempted, the ganglion cells appear randomly scattered. This result is consistent with previous attempts to discern such a topography in other species [in rat (Krimm and Hill, 1998), in hamster (Whitehead et al., 1999)]. However, although a topography of buds in the ganglion is not apparent, it remains conceivable that there may be a functional topography of ganglion cells based on the quality of
receptor cells innervated (i.e., sweet, bitter, sour, salty, and umami). Such an arrangement would be similar to that in the olfactory system (Mombaerts et al., 1996; Gao et al., 2000; Vossshall et al., 2000). Moreover, as with olfaction, a pattern may be evident in the projections to the brain (i.e., within the NTS) for the different types of ganglion cells. Establishing this for taste is at the limits of our current technology. It would require point-to-point mapping by labeling all of the same quality receptor cells with a trans-synaptic marker or by engineering mice that express trans-synaptic proteins in all cells that respond to a single quality with a trans-synaptic marker or by engineering mice that express point mapping by labeling all of the same quality receptor cells the limits of our current technology. It would require point-to-the different types of ganglion cells. Establishing this for taste is at

Anatomically, however, on a larger scale, the ganglion cell populations innervating the two taste receptor fields are arranged topographically; cells innervating the tongue are segregated from those innervating the palate. Some correspondence between the topography of cells in the ganglion and taste quality can be inferred in light of the differential representation of palate and anterior tongue seen here, and the different functional types of neurons that innervate these regions. For example, palatal neurons, unlike those innervating the tongue, include a subset that responds with high frequency to sucrose stimulation [in rat (Sollars and Hill, 2005)]. Conceivably, sucrose is represented rostrally on the periphery of the geniculate ganglion. That broadly different receptor areas are represented by neurons in different regions of the ganglion is consistent with previous differential location of ganglion cells responsive to palate, anterior tongue, and ear stimulation in cat (Boudreau et al., 1971). Moreover, the unbranched nature of geniculate ganglion cell peripheral fibers is preserved in the representations of entire peripheral fields in the ganglion; geniculate ganglion cells innervate fungiform or palatal taste buds, but not both.

References


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Injecting three adjacent buds in the hamster reveals a significant incidence of ganglion peripheral fiber branching among taste buds. **A**, Populations of single- and double-labeled ganglion cells in the hamster have a different profile than that for mice. The percentage of cells double labeled through fiber collaterals, known for the hamster (Whitehead et al., 1999), is markedly greater (35.3%) than in the mouse (5.4%). Error bars represent SDs. M, Single labeled; D, double labeled; T, triple labeled. **B**, Three adjacent buds, injected with different color markers in the same approximate sector of the tongue tip in hamster, multiply labels a notable subset of ganglion cells (4 cases; C–G, arrows, double labeled; circles, triple labeled) (compare with Fig. 3 for mouse). The asterisk marks the GSP nerve, whereas the + symbol marks the central (toward brain) corner of the ganglia. Viln marks the seventh cranial nerve. Scale bars, 100 μm.