The neurotrophin receptor p75 regulates gustatory axon branching and promotes innervation of the tongue during development

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Abstract

Background: Brain-derived neurotrophic factor (BDNF) and neurotrophin-4 (NT4) regulate the survival of gustatory neurons, axon growth and branching, and innervation of taste buds during development. These actions are largely, but not completely, mediated through the tyrosine kinase receptor, TrkB. Here, we investigated the role of p75, the other major receptor for BDNF and NT4, in the development of the taste system.

Results: We found that p75−/− mice showed delayed axon outgrowth and reduced branching of gustatory axons at embryonic day (E)13.5. From E14.5 to E18.5, gustatory neurons innervated fewer papillae and completely failed to innervate the mid-region of the tongue in p75−/− mice. These early effects of the p75 mutation on gustatory axons preceded the loss of geniculate ganglion neurons starting at E14.5 and also contributed to a loss of taste buds at and after birth. Because knockouts for the TrkB receptor (TrkB−/−) do not lose as many taste buds as hybrid knockouts for its two ligands (BDNF and NT4), we asked if p75 maintains those additional taste buds in the absence of TrkB. It does not; hybrid TrkB−/−/p75−/− mice had more taste buds than TrkB−/− mice; these additional taste buds were not due to an increase in neurons or innervation.

Conclusions: p75 regulates gustatory neuron axon branching and tongue innervation patterns during taste system development. This function is likely accomplished independently of BDNF, NT4, and TrkB. In addition, p75 does not support the remaining neurons or taste buds in TrkB−/− mice.

Keywords: Taste, Gustatory, p75, Neurotrophins, TrkB

Background

During development, axons from gustatory neurons of the geniculate ganglion grow toward and innervate specific regions of the tongue. These axons follow defined pathways, suggesting that a series of molecular cues in the environment regulate their guidance [1]. Upon arrival in the tongue, these gustatory fibers innervate specialized structures (for example, fungiform placodes) but not the surrounding epithelia. The fibers defasciculate upon entering the epithelium thereby forming a neural bud [2]. In addition to regulating the location of innervation within the tongue, molecular factors such as neurotrophins also control the number of neurons that innervate each taste bud [3].

Brain-derived neurotrophic factor (BDNF), a neurotrophin produced in developing taste papillae, regulates the survival of gustatory neurons [4,5]. Acting as a chemottractant [6], BDNF also guides gustatory axons to their final targets [7-9]. Neurotrophin-4 (NT4) also regulates the survival of gustatory neurons, albeit earlier in development than BDNF [10,11], but has no role in axon guidance [8]. Therefore, these two neurotrophins control different aspects of taste system development. These two factors function differently primarily because of their locations and timing of expression and not due to differential signaling [12,13]. They accomplish their diverse functions by acting through the same receptors - the tyrosine kinase receptor TrkB and the pan-neurotrophic receptor p75 [14-16]. Recently, we found that neurotrophins guide a small number of TrkB-negative gustatory neurons to fungiform placodes [17]. As p75 also binds BDNF and NT4, it is possible that this receptor could mediate BDNF and NT4 actions.
NT4 function independently of TrkB. Observations that adult p75 mutant mice have fewer taste buds and geniculate ganglion neurons [18] suggest that p75 regulates the development of the taste system. However, the specific role of p75 in taste system development is difficult to predict, as p75 is known to interact with several molecular factors in a context- or system-specific manner [19-26].

The goal of the present study was to determine the role of p75 in taste system development. In particular, we examined whether p75 is required for geniculate ganglion axon guidance and neuron survival as well as the maintenance of taste buds during embryonic development by comparing tongue innervation patterns and numbers of geniculate ganglion neurons and taste buds between wild-type and mutant (p75−/−) mice. We found that p75 regulates geniculate ganglion axon branching and promotes innervation of the tongue but does not play a prominent role in gustatory neuron survival. Furthermore, p75 does not appear to mediate BDNF or NT4 function.

Results

Embryonic p75−/− mice show reduced geniculate ganglion axon branching and a lack of innervation in the mid-region of the tongue

We previously found that BDNF and NT4 guide a small number of neurons to fungiform placodes independently of TrkB during development [17], leading us to speculate that the p75 receptor may be important for the TrkB-independent gustatory axon guidance. To test this possibility, we labeled geniculate ganglion neurons with DiI and examined innervation of the tongue in wild-type and p75−/− mouse embryos. In wild-type mice on embryonic day (E)13.5, chorda tympani fiber bundles branched from the main nerve bundle toward the surface of the lateral parts of the tongue but did not yet reach the dorsal tongue surface (Figure 1A,C). Even at this early stage, p75−/− mice showed less axon branching than wild-type mice (Figure 1B,D). In wild-type mice on E14.5, fiber bundle branches reached the tongue surface across its entire medial-to-lateral extent, where they formed bulb-like terminations called 'neural buds' [2] (Figure 1A). By contrast, p75−/− mice showed less branching and fewer neural buds at the dorsal surface of the tongue, particularly in the mid-region, with only a few neural buds present at the tip and posterior regions of the tongue (Figure 1B-C). Because innervation of the tongue proceeds in a caudal-to-rostral direction during development [27], the presence of neural buds at the tongue tip suggests that axonal growth and targeting is not merely delayed but that branching is also reduced in p75−/− mice.

To determine if this reduced innervation in p75−/− mice continues into development, we examined chorda tympani branching in wild-type and p75−/− mice on E16.5 and E18.5. Among wild-type mice, the pattern of axonal branching on E16.5 was similar to that on E14.5 (Figure 2D). p75−/− mice, however, showed an even more obvious disruption in branching, with most branches not entering the mid-region of the tongue (Figure 2E). At the tongue tip, the general pattern of branching was similar between wild-type and p75−/− mice, but the number of neural buds was noticeably reduced in p75−/− mice (Figure 2F,G). On E18.5, there was still an obvious deficit in innervation of the tongue mid-region in p75−/− mice compared with wild-type mice (Figure 3A-F). By sectioning the tongue and examining innervation of its dorsal surface, we confirmed that branches reached the epithelial surface in p75−/− mice but that innervation did not extend as far caudally as in wild-type mice (Figure 3G,H). Therefore, p75 is required for...
the development of normal chorda tympani branching and innervation of the mid-region of the tongue.

Embryonic p75−/− mice lose geniculate ganglion neurons beginning on embryonic day 14.5
Mice lacking the p75 receptor lose about 25% of geniculate ganglion neurons by adulthood [18], suggesting that our observed reduction in branching in p75−/− mice could be due to a loss of neurons. Geniculate ganglion neuron survival depends on both BDNF and NT4, with a loss of neurons starting at E11.5 in Ntf4−/− mice and at E13.5 in Bdnf−/− mice [5,11] and a major loss of neurons by E13.5 in TrkB−/− and Bdnf−/−/Ntf4−/− mice [15,17]. We reasoned that if the p75 receptor mediates geniculate ganglion neuron survival via BDNF or NT4 binding, we should see neuron loss in p75−/− mice by E13.5. To test this hypothesis, we examined geniculate ganglion neuron number in wild-type and p75−/− mice at E13.5 using anti-β-III tubulin antibody (TUJ-1). Geniculate ganglion neurons were easily identified by their clear nuclei and dark cytoplasm (Figure 4A, B). We found no significant neuron loss in p75−/− mice compared with wild-type mice at E13.5 (P ≤ 0.05 (E14.5), P ≤ 0.01 (E16.5), P ≤ 0.001 (E18.5); Figure 5), indicating that p75−/− mice start to lose geniculate ganglion neurons at E13.5 and this loss continues throughout embryonic development. Because E14.5 is the age that gustatory nerves start to innervate peripheral targets [8], and neuron number is controlled by the
amount of trophic support from these peripheral targets, this loss of neurons may be due to disrupted innervation of targets.

**p75<sup>−/−</sup> mice lack taste buds and taste bud innervation at birth**

Neurotrophins are required for the normal development of taste buds by birth [28-30] and p75 is important for maintaining a normal number of taste buds in adulthood [18]. Given the disrupted innervation pattern observed in p75<sup>−/−</sup> mouse embryos, we hypothesized that they would show a lack of taste buds at birth. We used anti-cytokeratin-8 and anti-P2X3 to label taste buds and their innervation in wild-type and p75<sup>−/−</sup> mice at birth. We found that p75<sup>−/−</sup> mice had significantly fewer taste buds than wild-type mice (P < 0.001; Figure 6D). Whereas all taste buds were innervated in wild-type mice, only 80% of taste buds were innervated in p75<sup>−/−</sup> mice (Figure 6A-C). This finding is consistent with our earlier observation that embryonic p75<sup>−/−</sup> mice had un-innervated regions of the tongue, particularly the mid-region. When we measured the volume of un-innervated (Figure 6B) and innervated (Figure 6C) taste buds in p75<sup>−/−</sup> mice, we found that innervated taste buds were significantly larger than un-innervated taste buds (Figure 6E, P ≤ 0.01). Therefore, p75 receptor mutation disrupted innervation of the tongue during development, reducing both the number and volume of taste buds.

It is well established that innervation is required to support the existence of taste buds during postnatal development [31,32], suggesting that un-innervated taste buds could be lost at later ages. Therefore, we quantified taste bud number in wild-type and p75<sup>−/−</sup> mice at postnatal day (P)10. Whereas wild-type mice had 116 ± 7 taste buds, p75<sup>−/−</sup> mice had only 75 ± 2 taste buds. However, no additional taste buds were lost in p75<sup>−/−</sup> mice at P10 (35%) compared with birth (42%), indicating that un-innervated taste buds may continue to be supported postnatally.

![Figure 3](http://www.neuraldevelopment.com/content/9/1/15)
p75 is not responsible for supporting the existence of taste buds in TrkB$^{-/-}$ mice

We previously showed that TrkB$^{-/-}$ mice possess more taste buds than hybrid Bdnf$^{-/-}$/Nft4$^{-/-}$ mice [17], perhaps because BDNF or NT4 function through other types of receptor in the absence of TrkB. One likely possibility is the p75 receptor, as we showed earlier p75 is required for normal innervation of the tongue surface. If BDNF and NT4 function via p75 to support the additional taste buds in TrkB$^{-/-}$ mice, the number of taste buds should be reduced in hybrid TrkB$^{-/-}$/p75$^{-/-}$ mice. To test this possibility, we quantified taste buds in TrkB$^{-/-}$/p75$^{-/-}$, TrkB$^{-/-}$, p75$^{-/-}$, and wild-type mice at E13.5. Surprisingly, we observed an increase in taste bud number in TrkB$^{-/-}$/p75$^{-/-}$ mice (53 ± 11) compared with TrkB$^{-/-}$ mice (31 ± 8; $P < 0.01$). The number of taste buds in TrkB$^{-/-}$/p75$^{-/-}$ mice was similar to that in p75$^{-/-}$ mice (60 ± 9) but still significantly less than in wild-type mice (103 ± 14; $P < 0.001$). Therefore, p75 does not appear to support the remaining taste buds in TrkB$^{-/-}$ mice.

Interestingly, in other systems, p75 causes the death of some neurons in the absence of Trk receptors [33,34]. Therefore, we speculated that this might also be true for gustatory neurons, with rescued gustatory neurons supporting the additional taste buds in TrkB$^{-/-}$/p75$^{-/-}$ mice. To test this idea, we quantified geniculate ganglion neuron...
number in \(TrkB^{-/-}\) mice on E13.5, at which time a deficit in neuron number is observed in \(TrkB^{-/-}\) mice. We found that \(TrkB^{-/-}\) mice also lacked geniculate ganglion neurons at E13.5, with no significant difference in the number of remaining neurons between \(TrkB^{-/-}\) (97 ± 18) and \(TrkB^{-/-}/p75^{-/-}\) mice (107 ± 10). Therefore, the deletion of p75 does not appear to increase the number of taste buds in \(TrkB^{-/-}\) mice by rescuing neurons, which is consistent with our observation that the additional taste buds in \(TrkB^{-/-}/p75^{-/-}\) mice did not appear to be innervated (Figure 7A-F). These results indicate that p75 may promote apoptosis in the taste bud in the absence of innervation.

**Discussion**

Our goal was to determine the role of the neurotrophin receptor p75 in taste system development. In particular, because we previously found that a small subpopulation of TrkB-negative gustatory axons require BDNF to navigate to taste buds through an unknown receptor [17], we tested whether p75 could be this receptor and regulate axon guidance and innervation of targets in a
manner similar to BDNF. We found that deletion of p75 disrupted innervation of the tongue surface during development. Specifically, peripheral axons branched less and did not innervate the mid-region of the tongue in embryonic p75−/− mice. As this pattern of innervation was unlike that previously observed in BDNF mutant mice [8], p75 probably does not mediate the role of BDNF in axon guidance. We also found that geniculate ganglion neurons were lost after alterations in axonal branching in p75−/− mice, suggesting that disrupted innervation influences neuron survival in p75−/− mice, and not that neuron death disrupts innervation patterns. The reduced innervation to the tongue surface resulted in taste bud loss, with only 80% of remaining taste buds being innervated in p75−/− mice.

In p75−/− mouse embryos, the chorda tympani nerve branched less and failed to innervate the mid-region of the tongue. Because innervation progresses in a caudal/lateral-to-medial/caudal direction as nerve fibers grow into the tongue [2,35-37], this lack of innervation could be due to delayed or inhibited nerve outgrowth. This possibility is consistent with the finding that blocking p75 function in vitro reduces BDNF-stimulated outgrowth of geniculate ganglion neurons [38]. In the whisker pad, the absence of p75 delays trigeminal innervation, decreases axon branching, and is associated with delayed glial development [39]. Unlike trigeminal axons, however, which eventually innervate their targets, p75−/− mice show a consistent deficit in innervation of the tongue through E18.5, leading to a permanent reduction in the number of taste buds [18].

At all ages observed, most gustatory nerve endings were missing in the middle and caudal areas of the tongue in p75−/− mice. This pattern of innervation is different from that of Bdnf−/− mice, in which branching is increased and fibers extend to all regions of the tongue but fail to innervate fungiform papillae [8]. Thus, although p75 is a receptor for BDNF, it does not appear to mediate the role of BDNF in axon guidance, although it could partially mediate the role of BDNF in axon outgrowth [38]. Rather, it is more likely that p75 functions independently of TrkB ligands and instead interacts with a different factor to influence axon guidance. p75 is known to interact with several different guidance molecules, including ephrins [40]. Also, p75 modulates activity of the chemorepellent semaphorin3A (Sema3A). Dorsal root ganglion neurons in p75−/− mice are hypersensitive to Sema3A [41], and Sema3A acts as a chemorepellent against geniculate ganglion axons in vitro [37], with its repellent effect being neurotrophin-dependent [42]. Sema3A is expressed in the tongue epithelium early in development, and its expression slowly decreases in a lateral-to-medial direction as the tongue becomes innervated [43], thereby preventing premature innervation. Furthermore, Sema3A mutant mice show increased innervation of the tongue mid-line [37]. Thus, our finding of decreased innervation of the mid-region of the tongue in p75−/− mice is consistent with a hypersensitivity to Sema3A. In general, however, interactions of the p75 receptor with many different signaling factors could be important for axon guidance and targeting.

We found that p75−/− mice lose geniculate neurons by E14.5 and that this loss continues through E18.5. Because this loss occurs after neurons become dependent on BDNF and NT4 [5,11,44] it is not likely that p75 functions by mediating the effects of neurotrophins on neuron survival, at least not initially. Normally, gustatory nerves reach the tongue epithelium and innervate fungiform pla-codes by E14.5 [2,8]. In p75−/− mouse embryos, fewer fungiform papillae were innervated, indicating that neurons failed to reach their targets. Therefore, these neurons could die starting at E14.5 due to a lack of survival factors.

**Figure 7** Hybrid TrkB−/−/p75−/− mice have many un-innervated taste buds at birth. Keratin 8-labeled (green) and P2X3-labeled (red) tongue sections from (A) a p75−/− and (B) a TrkB−/−/p75−/− mouse. Whereas the p75−/− mouse had both innervated (D) and un-innervated (C) taste buds, the TrkB−/−/p75−/− mouse had mostly un-innervated taste buds (E,F). Scale bar = 100 μm for A and B, scale bar = 10 μm for C-F.
received from the targets. Consistent with this idea, we observed only a 30% loss in geniculate ganglion neuron number at E14.5, and no neuron loss at E13.5 even though axon branching was already disrupted. Given that alterations in innervation patterns were more severe and occurred earlier than neuron loss, it is more likely that disrupted innervation caused neuron loss rather than vice versa. Neuron loss in other sensory ganglia could also be due to failed targeting [39,45], which could explain why sensory neurons tend to be lost in the absence of p75 [33].

The p75 receptor can promote neuronal apoptosis in the absence of Trk receptors [33,34,48]; for example, nerve growth factor causes the death of retinal neurons that express p75 but not TrkA [48]. Also, BDNF activates p75 to induce apoptosis of sympathetic neurons [33], which do not express TrkB. Based on these findings, we examined whether p75 causes the death of geniculate ganglion neurons in the absence of TrkB and whether the additional removal of p75 can rescue these neurons. Surprisingly, TrkB−/−/p75−/− mice had similar numbers of neurons as TrkB−/− mice. This finding suggests that p75 plays a different role in the development of the taste system compared with other systems [33,48,49], perhaps due to differences among systems in the role of p75 receptor function. In the retina [48], sympathetic ganglia [33], and dorsal root ganglia [49], p75 induces apoptosis specifically in the absence of TrkA. Thus, TrkB may regulate cell death differently than TrkA [49], and the actions of p75 may differ in the absence of these two receptor types. Regardless of the underlying reason, our results indicate that p75 does not cause neuronal apoptosis when TrkB is absent during taste system development.

Because taste buds require support from nerve fibers to retain their normal numbers and size [50-53], taste buds are lost in the absence of neurotrophins [28-30,54]. Likewise, since innervation to the tongue was disrupted, p75 mice also had a loss of taste buds. In all cases with neurotrophin knockout mice some un-innervated taste buds, as defined by keratin 8 labeling, remain [15,54]. These taste bud remnants also remain following nerve section [50,55]. Consistent with this literature we found un-innervated taste buds that remained in p75−/− mice. Because nerve section results in a greater loss of taste buds during development than it does in adulthood [53,56,57], we thought that perhaps these additional taste buds would be lost postnatally, but that was not the case; they still remained in P10 p75−/− mice. It seems likely that these smaller remaining un-innervated collections of K8-positive cells simply do not require innervation to be maintained.

Not all neurotrophin knockout and receptor mice lose the same number of taste buds; for example, TrkB−/− mice possess more taste buds than hybrid *Bdnf−/−/Nt4−/−* mice at birth [15,17,58]. The reason for this is unclear but we speculated that the additional taste buds might be supported by BDNF or NT4 functioning via p75. However, mice with both TrkB and p75 mutations possessed more taste buds than TrkB−/− mice, suggesting that p75 is not required for the additional taste buds in TrkB−/− mice. Although this increase in number of taste buds was surprising, p75 has been found to cause the death of neurons in the absence of Trk expression [33,34,48]. When we investigated whether the increase in taste bud number in TrkB−/−/p75−/− mice might be supported by a larger number of neurons, we found that geniculate ganglion neuron number was unaffected by mutation of both TrkB and p75, and few of the taste buds in TrkB−/−/p75−/− mice appeared to be innervated. Therefore, p75 does not cause the death of neurons in the absence of Trk signaling in the taste system, and the additional taste buds in TrkB−/−/p75−/− mice relative to TrkB−/− mice are not rescued by a greater number of neurons. Together, these findings suggest that, although p75 regulates innervation of the tongue surface, it does not mediate the role of BDNF in axon guidance or the role of TrkB ligands in neuron survival.

Conclusions

The pan-neurotrophic factor receptor, p75, regulates gustatory innervation patterns and branching of gustatory fibers in the tongue. Although gustatory neurons are lost in p75−/− mice this appears to be downstream of the innervation deficits, and is not likely due to a direct role of p75 in mediating BDNF- and/or NT4-regulated survival. The taste buds previously shown to be remaining in TrkB−/− mice, but not Bdnf−/−/Nt4−/− mice, are not maintained by p75 and leave open the possibility that the two factors function through an additional receptor other than TrkB and p75 to regulate gustatory development.

Methods

Animals

Heterozygous *TrkB+/* (stock no. 002544; Jackson Laboratories, Bar Harbor, ME, USA) and p75+/* (stock no. 002213; Jackson Laboratories) mice were bred to produce *p75−/−* and *TrkB−/−/p75−/−* mice. The day that a sperm plug was positively identified was designated E0.5. Mice were genotyped using polymerase chain reaction. All procedures were approved by the University of Louisville Institutional Animal Care and Use Committee committee and are in accordance with the guidelines of the US Public Health Service Policy on Humane Care and Use of..
Laboratory Animals and the National Institutes of Health Guide for the Care and Use of Laboratory Animals.

**Quantification of geniculate ganglion neurons**

E13.5 ($p75^{-/-} n=3$), $TrkB^{-/-}/p75^{-/-} n=2$, and wild-type $n=3$), E14.5 ($p75^{-/-} n=3$, and wild-type $n=3$), E16.5 ($p75^{-/-} n=3$, and wild-type $n=3$), and E18.5 ($p75^{-/-} n=3$, and wild-type $n=3$) embryos were heat-dried overnight, rehydrated, placed into citrate buffer (pH 6.0), heated for 15 minutes in a boiling water bath, and incubated for 10 minutes at room temperature.

Primary rat anti-cytokeratin 8 antibody (1:50; Developmental Studies Hybridoma Bank, Iowa City, IA, USA) and rabbit anti-P2X3 antibody (#AB5895; 1:500, Millipore, Billerica, MA, USA) were used to label gustatory nerves and taste buds. Cytokeratin 8 is a marker of simple epithelia that labels most columnar taste cells [60,61], and P2X3 is an ATP channel that is required for taste function [62] and expressed in most geniculate ganglion neurons innervating the tongue [63]. Secondary anti-rat Alexa 488 (green) and anti-rabbit Alexa 555 (red) antibodies (1:500, Molecular Probes, Grand Island, NY, USA) were used to visualize cytokeratin 8- and P2X3-containing nerve fibers, respectively. Sections were examined in order, and individual taste buds were followed across sections so that each taste bud was only counted once.

To assess innervation of taste buds, confocal stacks of optical sections with a Z-step of 0.5 μm were imaged for three to five taste buds from each mouse ($p75^{-/-} n=5$, and wild-type $n=4$) and analyzed using ImageJ software (http://rsbweb.nih.gov/ij/). The area of the taste bud in each optical section was measured, and these areas were summed and multiplied by section thickness to calculate taste bud volume. The area of P2X3-positive staining within outlined taste buds was also measured in each optical section, and these areas were summed and multiplied by section thickness (0.5 μm) to calculate the volume of P2X3 labeling within taste buds. The percentage of taste bud volume occupied by innervation was calculated by dividing the volume of P2X3 labeling by the volume of cytokeratin 8 labeling.

**Dil-labeling of the geniculate ganglia**

E14.5 ($p75^{-/-} n=3$, wild-type $n=3$), E16.5 ($p75^{-/-} n=6$, wild-type $n=5$), and E18.5 ($p75^{-/-} n=4$, wild-type $n=6$) mice were anesthetized and transcardially perfused in ice-cold 4% PFA. E13.5 mice were immersion-fixed. Dil labeling was performed as previously described [64]. Briefly, brains were removed and Dil crystals were placed on the geniculate ganglion and facial nerve. Embryos were incubated at 37°C for 2 to 8 weeks depending on their age. The tongue was then dissected, examined, and photographed using a fluorescent dissecting microscope (Leica Microsystems, Wetzlar, Germany) equipped with a camera (QImaging, 19535 56th Avenue, Suite 101 Surrey, BC.).
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