Sex-dependent Differences in the Genomic Profile of Lingual Sensory Neurons in Naïve and Tongue-Tumor Bearing Mice

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Abbreviated Title: Transcriptomic Changes in Lingual sensory neurons in males and females in oral cancer.

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Abstract

While sex-specific prevalence of orofacial pain is established, mechanisms of sex-dependent orofacial pain are widely understudied. To this end, a significant gap in knowledge exists about comprehensive regulation of tissuespecific trigeminal sensory neurons in diseased state of males and females. Using RNA sequencing of FACS sorted retro-labeled sensory neurons innervating tongue tissue, we determined changes in transcriptomic profiles in males and female mice under naïve as well as tongue-tumor bearing conditions Our data revealed the following interesting findings: 1) Tongue tissue of female mice was innervated with higher number of trigeminal neurons compared to males; 2) Naïve female neurons innervating the tongue exclusively expressed immune cell markers such as Csf1R. C1ga and others, that weren't expressed in males. This was validated by Immunohistochemistry. 3) Male neurons were more tightly regulated than female neurons upon tumor growth; 4) While very few differentially expressed genes (DEGs) overlapped between males and females post-tumor growth, several biological processes (BPs) were similar between two sexes. However, additional distinct processes were sex-specific: 5) Post-tumor growth, male DEGs contained an equal mix of transcription factors. ligands, growth factors, receptors and channels, whereas female DEGs predominantly contained channels/receptors, enzymes, cytokines and chemokines. Taken together, this is the first study to characterize the effect of sex as well as of tongue-tumor on global gene expression, biological pathways and molecular function of tongue-innervating sensory neurons.

Keywords: Oral cancer, pain, Trigeminal Sensory neurons, Sex-differences, Transcriptomic profile

Introduction

Sex-dependent differences in orofacial pain has been clinically well- established with higher prevalence in women in several different pathological conditions(1-6); although the mechanism of sex-differences remain elusive. It is known that sensory neurons including trigeminal neurons, that regulate pain in the orofacial region, are genomically different in males and females even under naïve conditions (7-10). These studies have primarily been conducted using the entire neuronal population of the ganglionic tissue. However, several reports reveal that sensory innervation can be different with each tissue type (7, 11-14). Accordingly, we have previously identified subsets of sensory neurons expressed in mouse tongue(13) that are varied from those innervating the masseter muscle(15). Therefore, it is vital in delineating tissue-specific sex-dependent differences in trigeminal sensory neurons. Moreover, a significant gap-in knowledge exists for sex-specific changes of trigeminal neurons that specifically innervate diseased-tissues. Such studies can provide crucial information about the regulation of trigeminal sensory neurons in tissue-specific pathologies.

The tongue is among the vital organs of the orofacial region as it controls many essential daily activities such as speech, licking, taste, chewing and swallowing. Importantly, many lingual ailments cause acute and chronic pain leading to very distressing and debilitating quality of life (16-29). In fact, among all chronic orofacial pain conditions that pose a challenge in management, three are known to primarily affect the tongue. These include tongue cancer, oral mucositis and burning mouth syndrome(23-26, 30-42). Interestingly, sex-dependent differences in the manifestation of pain have been reported for each of these conditions(30, 31, 35, 36, 43-48). Therefore, the current study identified the differences in tongue-innervating sensory neurons between males and females under naïve and diseased state. We used tongue cancer as our disease model as approximately 50% of oral cancer patients report pain throughout the course of the disease and of these, the prevalence of pain is highest in tongue cancer patients(23-26) (49). Pain from tongue cancer is extremely weakening and significantly deteriorates patient quality of life in addition to having cancer due to very limited treatment options available. Therefore, using the orthotopic tongue cancer xenograft model in mice, we performed bulk-RNA sequencing of isolated tongue-innervated sensory neurons from males and females, to identify changes in genes, biological processes and molecular function between sexes under normal and tumor-bearing conditions.

Materials and Methods

<u>Animals</u>

Six- to eight-week-old adult Balb/c male and female athymic nude mice (Jackson Labs, Bar Harbor, ME, USA) were used for all experiments. All animal handling and procedures were performed according to approved UTHSCSA IACUC protocols and conformed to the guidelines of International Association for the Study of Pain (IASP). Animals were housed in the UTHSCSA laboratory of Animal Resources (LAR) for at least 4 days prior to start of experiments.

In Vivo Orthotopic Xenograft Tongue Tumor Model

Using human oral squamous cell carcinoma cell lines: HSC3, tongue tumors were induced in mice as described by us previously(27, 36, 37). Animals were anesthetized with isoflurane inhalation and 50ul of 3.5 x 10[^] HSC3 cells were injected unilaterally in the ventral side of the tongues using insulin syringes. Animals were then allowed to recover in their cages. Animals were used for experiments at day 15 post-cell inoculation.

Retro-labeling of Tongue-Innervating TG neurons

Tongue-innervated sensory neurons were labeled as described by us previously(13). Briefly, animals were anesthetized with isoflurane inhalation and 10uls of 1% wheat germ agglutinin (WGA)-AF488 (Promega), diluted in 1%DMSO, was injected bilaterally twice in tongue of each animal. The first injection was given in the superficial epithelial layer and 4 hours later, a second injection was given in the deeper muscular layers of the tongue. Animals were allowed to recover in their cages for 2 days before harvesting trigeminal ganglia (TG) tissues.

Preparation of single-cell suspension and flow sorting

Tongue-innervating sensory neurons were isolated from 4 groups: Male normal (MN), Female normal (FN), Male tumor (MT) and Female tumor (FT). Neurons were isolated by preparation of single-cell suspensions of TG tissues followed by flow sorting of WGA+ cells. Each sample was prepared by pooling 4 TGs from 2 animals for normal groups and 4 animals for tumor-bearing groups as only ipsilateral TGs were collected from this group. A total of 3 samples per group was prepared. TG tissues were dissected and collected in cold 1X HBSS buffer, washed three times with 1X HBSS and incubated with 5ul of 50ng/ml dispase (type 2, Sigma) and 75ul of 2.5mg/ml liberase (Roche) for 60 mins at 37°C for enzymatic digestion. Following this, tissues were centrifuged at 2min at 1000rpm and washed with 5mls of DMEM containing 5% FBS and resuspended in 1.5ml DMEM with 5% FBS and triturated with a Pasteur pipette to breakdown the tissues and prepare a homogenous solution. The solution was then strained with a 100 um strainer to remove all debris, supernatant collected in an eppendorf tube and subjected to flow sorting. Flow sorting was performed using FACSAria III (BD Biosciences; San Jose, CA) using 130 µm nozzle. Consecutive gates were used to isolate WGA-AF488 labeled TG neurons. First, debris was excluded by forward scatter area (FSC-A) and side scatter area (SSC-A) gating. Second, duplets and clumps were excluded by side scatter width (SSC-W) and side scatter area gate (SSC-A) gate. Third, WGA-AF488+ bright cells were gated compared to unstained TG control and sorted directly to RLT buffer (Qiagen) containing 1% 2-mercaptoethanol (Sigma) to be able to use for RNA extraction.

RNA isolation

RNA was extracted using Qiagen RNeasy micro kit (Qiagen) with on column DNase I digestion according to manufacturer's instructions. RNA quantity was evaluated using Agilent 2100 Bioanalyzer RNA 6000 Nano chip (Agilent Technologies, Santa Clara, CA).

Bulk RNA sequencing

Bulk-RNA sequencing of samples was conducted at the UTHSCSA Genome Sequencing Facility. RNA integrity was determined using Fragment Analyzer (Agilent, Santa Clara, CA) prior to library preparation. All samples were ensured to have RIN values >6.5 to proceed with library preparations. RIN values for each sample is listed in **Supplementary Table 1**. RNA-seq libraries were prepared according to SMART-seq2 protocol (50), with the following modifications: PCR preamplification to 10-12 cycles, two rounds beads cleanup with 1:1 ratio after cDNA synthesis, and 0.6-0.8 dual beads cleanup for Nextera XT DNA-seq library purification. RNA-seq libraries were sequenced using Illumina HiSeq 3000 system (Illumina, San Diego, CA) with 50bp single-read sequencing module. Upon sequencing completion, short read sequences from RNAseq were first aligned to UCSC mm9 genome build using TopHat2 aligner and then quantified for gene expression by HTSeq to obtain raw read counts per gene, and then converted to RPKM (Read Per Kilobase of gene length per Million reads of the library).

Analyses of RNA sequencing Data

Sequencing data was analyzed as previously described(51). Briefly, differential expression analysis was performed using DESeq (52) algorithm (R/Bioconductor) to estimate the differential expression in read counts and their statistical significance. Significantly differentially expressed genes were selected based on following criterion: 1) RPKM > 5 in one of the comparison group, 2) fold-change > 1.5, and 3) differential expression p-value < 0.05 along with False-discovery rate (FDR) for each gene. When comparing female tumor samples with male tumor samples, all genes coded in sex-chromosomes (chrX and chrY) were excluded. Functional assessment of DEGs were performed by using over-representation statistic (Fisher's Exact test) for Gene Ontology using Panther platform.

Immunohistochemistry

Protocol for immunostaining is described previously by us(13, 36). Mice were anesthetized with ketamine (75mg/kg)/ dexmedotomidine (1mg/kg) solution and perfused with 4% paraformaldehyde (PFA) and trigeminal ganglia was dissected bilaterally from each mouse. Tissues were post-fixed in PFA, washed in 0.1M phosphate buffer and cryosectioned in Neg-50 (Richard Allan, Kalamazoo, MI, USA) at 20uM thickness. Sections were then subjected to immunostaining with specific primary antibodies as described in **Table 1**. Donkey anti- secondary antibodies were purchased from Molecular Probes, Eugene, OR, USA and were used at a dilution of 1:200 for all experiments. Following staining, tissue sections were mounted in Vectashield and imaged using Nikon Eclipse

90i microscope equipped with a C1si laser scanning confocal imaging system. Z-stack images were acquired of the V3 region of TGs from at least 2 animals per group and a total of at least 6-10 images were taken for each antibody combination per group. All images were obtained with a 20x objective at fixed acquisition parameters across all groups and were unaltered from that initially taken. Laser gain settings were determined such that no-primary control did not show any positive staining. Quantitation was achieved using Adobe Photoshop 2023 for number of neurons above threshold, in the V3 region of the TG tissue in each image. A total of 200-500 neurons per analyses were counted.

Marker	Company	Catalog Number	Dilution	
TRPV1	Neuromics, Edina,	GP14100	1:200	
	MN			
CSF1R	Thermofisher	14-1152-85	1:50	
	Scientific, Waltham,			
	MA, USA			
C1qa	Novus Biologicals,	NB200-539	1:50	
	Centennial, CO,			
	USA			
IL1-Beta	Novus Biologicals	NB600-633	1:50	
HSP70	Novus Biologicals	NBP1-77456SS	1:50	
Table 1. List of Primary Antibodies used for				
Immunohistochemistry				

Statistical Analyses

Sample sizes were calculated using G-power application to obtain 80% power at a two-sided tail with α error probability of 0.05. All animals were allocated to groups using simple randomization. All statistical analyses and graphical representations were performed in GraphPad Prism 9.0. Data are presented as mean ± standard error of the mean (SEM) or as heatmaps and volcano plots for RNA sequencing data. Statistical significance was determined using either Paired Student T-test, Unpaired Student's T -test, One-way ANOVA or Two-way ANOVA with Sidak's post-hoc test and p<0.05.

Results

Isolation of Lingual TG neurons revealed differences in number of innervating neurons between males and females.

To assess transcriptomic changes in tongue-innervating TG neurons in males and females under naïve and tumor-bearing conditions, we performed bulk-RNA sequencing of lingual TG neurons. To achieve this, we first isolated retro-labeled TG neurons innervating mouse tongue using WGA-488 and subsequently flow-sorted fluorescently labeled neurons from four groups: Male normal (MN), Male tumor (MT), Female Normal (FN) and Female Tumor (FT). Representative gating strategy for the sorting protocol is shown in **Fig 1A**. Flow sorting obtained an average of approximately 10K to 23K neurons per sample for all four groups (**Fig 1B and Suppl Table 1**). Surprisingly, we found that the number of neurons isolated from all of female samples from both groups (i.e. FN and FT,) were higher (approx. 22000 cells) compared to male groups (i.e MN and MT, approx. 12000 cells) (**Fig 1B**). In potting the percentage of WGA+ neurons sorted over all live events in the samples, we found that percentage of WGA+ neurons isolated from female samples were significantly higher (approx. 3 fold, one-way ANOVA, p<0.05) than in males (**Fig 1C**) suggesting that female mice may have increased number of sensory neurons innervating the tongue tissue compared to males in mice. To further investigate this finding, we

employed immunohistochemistry to evaluate nociceptive and non-nociceptive WGA+ neurons in naïve male and female mice. We used TRPV1 to distinguish between the two neuronal classes and found that males had a higher percentage of TRPV1+ tongue innervating neurons compared to females **Fig 1D and E**) (27.6% males vs 19.31% females, two-way ANOVA, p=0.0077). Accordingly, females had a higher percentage of TRPV1 negative tongue innervating neurons than males (Fig 1D and E) (72.38% males vs 80.6% females, two-way ANOVA, p=0.0077).

Sex-dependent differences in gene expression in lingual neurons of naïve mice

Bulk-RNA sequencing was conducted of flow-sorted WGA+ TG neurons from all four groups: MN, MT, FN and FT. Data were analyzed to identify differentially expressed genes (DEGs) using the criteria: RPKM>5, FC> 1.5 and p<0.05. In comparing FN vs MN groups, we identified a total of 81 DEGs between both sexes after excluding all sex-linked genes. Of these, 30 genes were exclusively expressed in females and only 2 genes exclusively expressed in males (Fig 2A and B.). The remaining 49 genes, while expressed in both sexes, were significantly upregulated in females compared to males (Fig 2C and D). Details of RPKM and FC of the top 10 genes is listed in Supplementary Table 2. Two genes, that were higher in males than females were Fam23a(transmembrane protein 236) and Ddx3y (DEAD box helicase). In further assessing the expression of female-specific genes, we tested the expression of selected two genes: Csf1R and C1qa using immunohistochemistry and confirmed their higher expression in females than in males. Csf1R was expressed in ~ 42% of all WGA+ neurons in females compared to 5.5% of WGA+neurons in males (Unpaired Student T test, p<0.0001) (Fig 2E). Of all Csf1R positive neurons innervating the tongue in females, 18% were found to be TRPV1 positive (Fig 2F and G) (Paired Student's T test, p=0.0049). Similarly, ~53% of all tongue innervating neurons expressed C1qa in females whereas almost no expression of this gene was found in males (Fig 2H) (Unpaired Student T test, p<0.0001). Majority of C1ga positive neurons in the tongue of females were TRPV1 negative (94%) with only a very small proportion of C1qa expression in TRPV1+ nociceptors (6%) (Fig 2I and J) (Paired Student's T Test, p<0.001).

Changes in Transcriptomic profile of lingual neurons upon tongue tumor growth in males and females.

We next compared MT vs MN to elucidate the changes in transcriptome of tongue-innervating neurons upon HSC3-induced tongue tumor in male mice. As shown in **Fig 3A and B**, we observed 83 DEGs, of which 75 genes were upregulated and 8 genes were down-regulated in tumor-bearing group compared to control. Of the 75 upregulated genes, two were sex-linked genes (**Fig 3B**) (i.e Timp1 and Rbm3). The Top 3 genes that were upregulated upon tumor growth were found to be Sprr1a (small proline-rich protein 1A, FC=19.09, p<0.01), Gal (Galanin, FC=5.11, p<0.0001) and FGF23 (fibroblast growth factor receptor 23, FC=4.22, p<0.001) (**Fig 3C**). The top three genes downregulated in the tumor-bearing group of male mice were Gm2058 (ubiquitin-conjugating enzyme E2H pseudogene, FC=0.211, p<0.05), Sst (somatostatin, FC=0.313, p<0.001), Ms4a3 (membrane-spanning 4-domains, subfamily A, member 3, FC=0.327, p<0.001) (**Fig 3C**).

Unlike in males, the number of DEGs in females upon tumor growth were considerably higher (**Fig 3D**). Comparing FT vs FN, we observed a total of 382 DEGs (**Fig 3D and E**), out of which 283 genes were upregulated and 99 genes were downregulated. Four upregulated (i.e. Timp1, Arxes1, Timm17b and Rnf113a1) and five downregulated (i.e. Bmx, Gyk, Map3k15, Tmem47 and Slitrk4) genes were sex-linked (**Fig 3E**). The top three upregulated genes in FT vs FN comparison were found to be Snord49b (small nucleolar RNA C/D box 49B, FC =86.79, p=0.028), Gm1568 (coiled-coil domain containing 177, FC=13.196, p=0.0023) and Npy (neuropeptide Y, FC=5.776, p<0.0001) (**Fig 3F**). The top three downregulated genes in tumor-bearing group of female mice were Ctgf (cellular communication network factor 2, FC=0.276, p=0.0007), Cav2 (caveolin 2, FC=0.305, p<0.0014) and Tek (TEK receptor tyrosine kinase, FC=0.32, p<0.0021) (**Fig 3F**).

Between DEGs observed from MT vs MN and FT vs FN, we found 18 genes that were common to both comparisons whereas majority of the genes were regulated in a sex-specific manner (**Fig 3G**). Interestingly though, out of the 18 common DEGs, 11 were upregulated in both comparisons whereas 3 genes that were upregulated in MT vs MN were significantly downregulated in FT vs FN (Suppl Table 3). In contrast 4 genes that were downregulated in MT vs MN were upregulated in FT vs FN (**Suppl Table 3**).

Collectively, these data suggests that the regulation of lingual neurons by the tongue tumor may be considerably different in males and females.

Biological Processes and Molecular Function Regulated Upon Tumor Growth in Males and Females. We next conducted gene ontology analyses to identify key biological processes in males and females upon tumor growth. Fourteen distinct biological processes were identified with upregulated DEGs in males upon tumor growth (Fig 4A). These processes pertained to signaling mechanisms, immune process and inflammation, metabolic processes as well as some others including response to glucocorticoid stimulus, circadian regulation glial cell proliferation and apoptosis. Majority of DEGs in males were involved in signaling pathways and immune and inflammatory processes with predominant processes being second messenger signaling, response to growth factors, inflammatory response or leucocyte migration (Fig 4A and Suppl Table 4). Additionally, considerable number of DEGs involved in cell differentiation and cell death were found to be upregulated in males upon tumor growth. Similar to males, sixteen processes were identified in females (Fig 4B), although all were classified under regulation of signaling pathways, immune and inflammatory processes as well as other processes such as cell death, transport regulation and positive regulation of angiogenesis. DEGs contributing to metabolic processes and cell proliferation and differentiation were not found in females, unlike in males (Fig 4B). Five specific processes were found to be common in both sexes. These included response to interferon-gamma, IL-1 signaling, leucocyte migration, inflammatory response and apoptosis (Fig 4A and B). However, the number and type of DEGs involved in each of these pathways were different between males and females (Suppl Table 4 and 5). While no specific processes were identified for downregulated DEGs in males due to very few numbers of genes, downregulated DEGs in females were found to be associated with processes such as negative regulation of angiogenesis, locomotion, cell adhesion and cytoskeletal organization (Fig 4C). List of downregulated DEGs associated with each of the BPs in females in given in Suppl. Table 6. Additionally, we also identified specific transcription factors (TFs), ligands, peptides, growth factors, receptors, channels, enzymes, chemokines and cytokines that were differentially regulated in males and females upon tumor growth. Of all DEGs between both comparisons, we found 11 TFs, 19 ligands, peptides and growth factors, 24 channels and receptors, 65 enzymes and 14 chemokines/cytokines (Fig 4D). While the number of TFs were higher in male neurons upon tumor growth, female neurons had higher number of channels and receptors, enzymes and chemokines and cytokines, regulated post-tumor growth (Fig 4E and F). Taken together, these data indicate significant differences in regulation of genes, processes and pathways by tongue tumor, between males and females.

Discussion

It is well established that chronic orofacial pain is sexually dimorphic with a higher prevalence in women than men for various pain conditions including temporomandibular joint disorders(53), apical periodontitis(4), oral mucositis(32, 33, 43), burning mouth syndrome(31) as well as oral cancer(42, 46, 47). Yet, there is a large gap in knowledge about the mechanisms that lead to sex-dependent differences in orofacial pain. Moreover, sexdependent global genomic changes in trigeminal neurons during disease is entirely unexplored. One prior study that reported changes in transcriptomic profiles of trigeminal ganglia in conditions of masseter muscle inflammation in rats, only included males (54). On the other hand, another study that investigated changes in gene expression of TG following neuropathic pain in males and females, did not evaluate the whole genome (55). Furthermore, both these studies employed whole ganglionic tissues that primarily represent non-neuronal population as recently confirmed by *Mecklenberg et al* (7). To address this drawback, a third study employed single-cell sequencing of human and mouse TG tissues to implicate genes and cell types in migraine. However, sequencing of TG tissue in this study only evaluated sex-differences under naïve conditions and genomic data obtained from mouse migraine models was not separated by sex(10). Therefore, no study till date has investigated changes in transcriptomic profiles of the trigeminal sensory neurons during disease in both sexes. Furthermore, it is critical to study neurons-innervating specific tissues over all ganglionic neurons as it is indicated that type of sensory innervation is tissue-specific. For example, it has been reported that dental pulp innervating neuronal soma primarily are large diameter myelinated neurons unlike other tissues in the orofacial region such as the skin or mucosa(56, 57). Accordingly, percentage of TRPA1 expressing neurons are higher in the oral mucosal tissues than in dental pulp(57). Similarly, we have shown that about 20% of tongue innervating sensory neurons are CGRP positive (13) whereas almost 50% of masseter muscle innervating neurons are CGRP positive(15).

Therefore, in the current study, we explored changes in genomic profile of isolated sensory neurons innervating mouse tongue tissue and identified sex-dependent differences. We employed the approach of bulk-RNA sequencing of FACS sorted retro-labeled mouse tongue innervating TG neurons, to be able to concentrate neuronal population from whole ganglionic tissue as well as explore tissue-specific differences in neurons

between sexes. Additionally, we utilized the preclinical mouse tongue cancer model as our disease model, as it has been shown that patients with tongue cancer are more commonly in pain than other oral cavity cancers and it is widely included in clinical and preclinical studies of oral cancer pain (35-40). However, till date, there is only a couple of studies investigating sexually dimorphic mechanism for tongue cancer pain (42, 46). While these studies demonstrated the role of the immune response in sex-dependent differences in oral cancer pain, differential gene expression in sensory neurons upon tumor growth cannot be excluded, as we and others have reported that sensory neuronal activity is altered upon oral tumor growth(27, 36, 37) (58-60).

FACS sorting of lingual TG neurons was performed from normal and tumor-bearing animals for both sexes and our data indicated that female mice have higher number of tongue-innervating sensory neurons than males as observed by significant increased percentages of sorted cells from female samples than male samples. In further exploring this result, we found that female mice had higher percentage of TRPV1- and lower percentage of TRPV1+ neurons innervating the tongue than in male mice. However, recently *Scheff et al*, demonstrated no difference in capsaicin sensitivity in the tongue between males and females under naïve conditions (59), indicating that while the percentage of TRPV1+ neurons are different between the two sexes, the total number of TRPV1+ lingual neurons may not be different between males and females. Therefore, given that the total number of tongue-innervating neurons are higher in females, our data suggests that the total number of non-nociceptive sensory neurons in mouse tongue may be higher in females than in males. To our knowledge, this is the very first evidence of differences in innervation density within orofacial tissues between sexes. Whether or not the increased innervation of non-nociceptive neurons contributes to sex-dependent pain observed in any of the chronic lingual diseased states is yet to be determined. Besides, whether this finding is specific to mice or exists in tongue tissues of higher order species such as human and non-human primates is unknown.

In analyzing RNA sequencing data between naïve male and female neurons, we found that several genes (i.e 30 genes) were exclusively expressed in females and not in males. This was in accordance to previous report showing increased DEGs in naïve female versus male neurons from whole mouse trigeminal ganglia(7). Interestingly, many of these DEGs such as CSF1R (colony- stimulating factor receptor 1)(61, 62), C1qa (complement component 1 q) (63-65), Sh2b3 (lymphocyte adapter protein)(66), Hhex (Hematopoetically expressed homeobox) (67, 68) and Retnlg (Resistin-like gamma)(69), are known to be primarily expressed in immune cells and have shown to play various roles in immune processes(61, 65, 70), myeloid cell differentiation(67, 68), cytokine signaling(66) and inflammation(69, 71). In fact, some of these genes such as Csf1R and C1qa are considered specific markers of mononuclear phagocytic system such as microglia, monocytes, macrophages and dendritic cells(61, 62, 64, 65). Csf1R in microglia and macrophages has been reported to be activated in neuropathic pain via its ligand Csf1 expressed in sensory neurons (70). Similarly, the contribution of microglia-expressed C1g proteins have also been reported in pain including orofacial pain(72-74). Additionally, it has been shown that Csf1R exerts its action in a sex-specific manner by favoring a response in males compared to females(75). However, for the first time, we report the expression of Csf1R in trigeminal sensory neurons, although its expression is specific to female neurons. Our immunohistochemical analyses of Csf1r and C1ga confirmed our RNA sequencing result and revealed that majority of these genes were expressed in non-TRPV1 expressing neurons, albeit Csf1R expressed in the small proportion of nociceptors as well. These data indicate that perhaps female sensory neurons may be functionally different than male neurons innervating the mouse tongue.

Our analyses for RNA sequencing data to identify changes in lingual sensory neurons upon tongue tumor growth revealed altered expression of several DEGs in both sexes. However, the number of DEGs in females was considerably higher than in males (83 DEGs in males vs 382 DEGs in females) suggesting that male neurons are more tightly regulated in spite of the tumor compared to female neurons. It would be interesting to investigate whether this result is specific to tongue tumors or even other lingual pathologies. Moreover, there were only 18 genes that were commonly altered between males and females, indicating that the mechanisms of neuronal regulation in oral cancer may be considerably different between sexes. Since the tumor developed in both sexes was from the same cell line, perhaps the differential regulation of neurons between males and females may be due to stark differences in the tumor microenvironment or alterations occurring at the ganglionic level upon tumor growth. Intraganglionic activation of macrophages (76) and satellite glial cells (77, 78) has been reported in pain. While no studies have yet probed the contribution of these cells for oral cancer pain, a similar response within the TG can be expected that in turn would lead to changes in the neuronal soma. On the other hand, the peripheral tumor microenvironment consists of various cell types including keratinocytes, fibroblasts, endothelial cells, Schwann cells, and immune cells. Schwann cells (79) and immune cells(42, 46) have already been shown to contribute to tongue cancer pain and as mentioned above, immune cells even play a sex-dependent role in tongue cancer pain (42, 46). Therefore, examining the role of these cell types in mediating global changes in the

neurons of males and females would be crucial in better understanding of the impact of the tumor microenvironment in sexually dimorphic tongue cancer pain.

It is noteworthy though that despite only 18 DEGS common to both males and females, 5 biological processes (BPs) were found to be common between males and females post tumor growth. These included apoptosis, response to interferon gamma, inflammatory response, leucocyte migration and IL-1 signaling. Of the 18 common DEGs, four genes were associated with two of the common BPs; i.e inflammatory process and apoptosis. These four genes were Timp 1(tissue inhibitor of metallopeptidase 1) Gal (galanin), Chi3l3 (chitinase-like 3) and Cxcl10 (chemokine-ligand 10). Timp1 is a glycoprotein that is known to promote cell proliferation and anti-apoptosis; is implicated in cancer progression(80); and negatively regulates matrix metalloproteinases and disintegrin-metalloproteinases (ADAMs)(81). Interestingly, ADAM17 is implicated in oral cancer pain(82). Accordingly, Timp1 has been shown to attenuate inflammatory pain in preclinical models(83).

Galanin is a neuropeptide and is considered a potent modulator of inflammation by promoting cytokine production in immune cells(84, 85). It also has been shown to induce cell death in pheochromocytoma cells(86). Importantly, the role of galanin in pain has been reported to have pro- and anti-nociceptive functions and whether or not galanin plays a role in peripheral nociceptive mechanisms is yet to be confirmed(87). Interestingly, galanin release from sensory neurons have been shown to promote oral cancer progression(88).

Chitinase-like proteins belong to the family of glycoside hydrolase and are involved in the regulation of the innate immune response(89). Chi3l3 has been specifically reported to orchestrate recruitment of eosinophils in meningitis and autoimmune neuroinflammation(90, 91). However, its role in pain is not yet defined.

Cxcl10 is an important chemokine for inflammatory processes and its function in pain has been studied in several pain models including neuropathic pain and inflammatory pain(92-94).

Aside from the above-mentioned four genes, many other genes were selectively altered in each sex, yet associated with the common BPs. For example, the cytokine, interleukin-1beta (IL1B) was specifically shown to be induced in female post tumor growth and was not expressed in males. The role of IL1B has been widely reported in pain, with the cytokine mostly produced by immune cells and other non-neuronal cells during injury(95, 96). However, because most of the studies have been conducted using male animals, neuronal IL1B has not been reported previously. To this end, a sex-specific role of IL1B in pain has not been studied till date. Interestingly, while induction of neuronal IL1B was only observed in females, downstream signaling of IL1 pathway was observed in both sexes post tumor growth indicating that perhaps IL1B in males may be increased in the periphery or within the ganglia in non-neuronal cells as demonstrated in other injury models(95-98).

Another example was expression of Hspa1a which encodes for heat-shock protein 70 (Hsp70). Our RNA sequencing data showed that while this gene was expressed in both sexes, it was specifically upregulated only in males post tumor growth. Interestingly, hsp70 has been demonstrated to have a protective role in pain during nerve damage(99, 100), migraine (101) and opioid-induced hyperalgesia(102). Therefore, upregulation of Hsp70 in males post-tumor growth might indicate that male neurons might express an endogenous feedback mechanism to suppress pain that may be lacking in females.

In addition to the common BPs, sex-selective BPs were identified post-tumor growth. Noteworthy femaleselective BPs including cytokine/chemokine production and signaling as well as angiogenesis. The contribution of various cytokine/chemokines in nociception and inflammation(103-105) is widely established and our data showed that the number of cytokine/chemokines altered in female neurons was higher than in males (i.e. 14 in females versus 3 in males). This is an intriguing yet not a surprising observation as it aligns with the finding that female neurons express immune cell markers as described above.

Female DEGs were also associated with angiogenesis. Interestingly, genes associated with positive regulation of angiogenesis (e.g. *Gadd45a, Dcn, Ngp, cxcl10, Hmox1,ccl11,Pgf etc)* were upregulated upon tumor growth and those associated with negative regulation of angiogenesis (e.g. *Cav2, Ptprm, Mmrn2, Xdh*, etc) were downregulated, indicating that sensory neuro-vascular interaction may be sex-specific.

In contrast, male-specific BPs included growth factor signaling and ion transport. One of the top genes upregulated in males upon tumor growth; fibroblast growth factor 23 (Fgf23) is reported to be associated to both of these processes. It is not only a growth factor that induces downstream signaling via its receptors (106) but also controls phosphate homeostasis (107). Furthermore, it is also considered a bone-derived hormone that is regulated by inflammation and associated with bone pain (108, 109).

Taken together, our data points to significant differences in the regulation of lingual sensory neurons in males and females upon tongue tumor growth. The current study is significant as it is the first to comprehensively characterize the genomic profile tongue-innervating neurons under naïve and tumor-bearing conditions. A few limitations of the study include use of one cell-line to induce the tongue tumor and at this point, the study cannot confirm which genes and processes are common across different tongue tumor types as well as across different

preclinical tongue tumor models. Besides, it would be useful to delineate the temporal effect of the tumor on neuronal genes and processes to further gain insight into the sex-specific mechanisms of oral cancer pain.

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Author Contributions

T.I. performed animal injections for tumor growth, WGA injections and conducted all immunohistochemical experiments, P.W. performed TG dissections and single cell suspensions for Flow sorting. A.T. performed flow sorting and RNA extractions, L.Z and K.W prepared cDNA libraries and performed RNA sequencing. Y.C and W.I analyzed RNA sequencing data along with S.R. S.R. conceptualized, designed and analyzed experiments, as well as wrote and edited the manuscript. All authors edited the manuscript.

Conflict of Interest

Authors declare no Conflict of Interest

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Figure Legends

Fig 1. Isolation and Estimation of Tongue-Innervating Sensory Neurons. A,B and C. Male and Female mice were injected with 3 x 10⁵ HSC3 cells in the tongue and at day 13 post-cell inoculation, tongue tissues were bilaterally injected with 1% WGA-488. Normal group received no HSC3 cells. Two days later, TG tissues were dissected to make single-cell suspension and subjected to flow sorting. Animals were groups into male normal (MN), male tumor (MT), female normal (FN) and female tumor (FT). N=3 per group. **A.** Gating Strategy for flow sorting is shown. **B.** Number of cells sorted for each group. **C.** Percentage of WGA+ neurons of total events in each group. Data are represented as mean ± SEM and analyzed by one-way ANOVA with Sidak's post-hoc test.p<0.05. **D and E.** Naïve male and female mice were injected with 1%WGA-488 and 2 days later, TG tissues were harvested for immunohistochemistry. Images were taken at 20x magnification using the Nikon C1 confocal microscope. N=2 per group. **D.** Percentage of TRPV1+/WGA+ and TRPV1-/WGA+ TG neurons in naïve males and females. Data are represented as mean ± SEM and analyzed by one-way ANOVA with Sidak's post-hoc test. **E.** Representative images of WGA and TRPV1 staining of TG tissues in male and females are shown. Arrows indicate colocalization of TRPV1 and WGA.

Fig 2. Differentially Expressed Genes in Male versus Female Lingual Neurons. A-D. Flow sorted neurons were subjected to RNA sequencing. DEGs were identified in normal male and female mice based on RPKM>5, FC>1.5 and p<0.05. N=3 per group. **A**. Heatmap of genes that were expressed exclusively in tongue-innervating neurons of normal male and female mice. **B**. Number of genes in males and females is depicted as bar graphs. **C**. Heatmap of genes differentially expressed in female normal (FN) vs male normal (MN). **D**. Bar graphs shows number of genes upregulated and downregulated in FN vs MN. **E-J**. Validation of RNA sequencing data in normal male and female by Immunohistochemistry. N=2 per group. Images taken with C1 Nikon Confocal Microscope at 20x magnification. **E**. Percentage of WGA+neurons expressing Csf1R in males and females. Data are represented as mean ± SEM and analyzed by unpaired Student's T Test at p<0.05. **F**. Percentage of Csf1R in lingual TRPV1+ and TRPV1- neurons. Data are represented as mean ± SEM and analyzed by unpaired Student's T Test at p<0.05 **G**. Representative images of immunostaining of Csf1R and TRPV1 in WGA+ neurons in males and females. White arrows indicate colocalization of Csf1R and WGA whereas orange arrows indicate colocalization of Csf1R, WGA and TRPV1. **H**. Percentage of WGA+neurons expressing C1qa in males and females. Data are represented as mean ± SEM and analyzed by unpaired Student's T Test at p<0.05 **F**.

Percentage of C1qa in lingual TRPV1+ and TRPV1- neurons. Data are represented as mean ± SEM and analyzed by paired Student's T Test at p<0.05 **G**. Representative images of immunostaining of C1qa and TRPV1 in WGA+ neurons in males and females. White arrows indicate colocalization of C1qa and WGA whereas orange arrows indicate colocalization of C1qa, WGA and TRPV1.

Fig 3. Effect of tongue tumor on transcriptomic profile of lingual neurons in male and female mice. DEGs were identified by conducting two comparisons. **A**. Volcanic Plots for all genes for MT vs MN comparison. DEGs identified are colored showing downregulated genes on the left and upregulated genes on the right. **B**. Number of upregulated and downregulated genes are plotted as bar graph. **C**. Top three upregulated and downregulated genes in MT vs MN are plotted as heatmap as well as tabulated for values of RPKM, Fold change (FC) and p-value. Data for heatmap plotted as fold change. Similarly, **D**. Volcanic plots for FT vs FN. **E**. Bar graph for upregulated and downregulated genes for FT vs FN. **F**. Top 3 three upregulated and downregulated DEGs in FT vs FN as heatmaps and tabulated for RPKM, FC and p-values. Data in heatmap plotted as fold change. **G**. Venn Diagram indicating number of overlapping and non-overlapping between MT vs MN and FT vs FN.

Fig 4. Biological Processes and Function of tongue-tumor controlled genes in males and females. Panther Pathway Analysis platform was used to elucidate biological processes from **A**. DEGs upregulated in MT vs MN, **B**. DEGs upregulated in FT vs FN and **C**. DEGs downregulated in FT vs FN. For each analysis, the number of DEGs associated with each biological process is plotted as bar graphs. **D**. Additional analyses was conducted to identify molecular function (MF) of all DEGs in males and females. Pie chart of the number of DEGs identified as transcription factors, ligands/receptors/growth factors (GFs), Channels/Receptors, Enzymes and Cytokines/Chemokines is shown. **E**. Heatmap of select DEGs for each of the molecular function indicate the differences in expression between MT vs MN and FT vs FN. Data plotted as fold change. **F**. Number of total DEGs obtained for each MF with MT vs MN and FT vs FN.





WGA

TRPV1

Merge

bioRxiv preprint doi: https://doi.org/10.1101/2023.01.14.52 01 ; bis version poord January 17, 2023. The copyright holder for this preprint (which was not certified by peer review) is the author/funder who is grane bioRxiv a license to display the preprint in perpetuity. It is made **Genes Exclusively Expressed** -ND 4.0 International Exclusive in FN vs MN



Validation of Genes by Immunohistochemistry



Male Merge CSF1R WGA TRPV1 Female

CSF1R



TRPV1

Female



C1qa

WGA

WGA

Merge

Merge



Biological Processes for Α **Upregulated DEGS in MT vs MN**



Biological Processes for Biological Processes for С **Upregulated DEGs in FT vs FN**

Downregulated DEGs in FT vs FN



DEGs by Molecular Function

В







Transcription Factors Ligands/Peptides/GFs Channels/Receptors Enzymes

Cytokines/Chemokines

Sample	Sorted Cells Collected (Mean± SEM)	RIN value of RNA (Mean± SEM)
Male Normal(MN)	10333±2667	7.567± 0.088
Male Tumor (MT)	12667± 2963	7.267± 0.49
Female Normal (FN)	23333±3383	7.233± 0.466
Female Tumor-1 (FT1)	20667± 5364	7.100± 0.23

Supplementary Table 1. Number of sorted cells and RIN values of RNA of all samples within four groups. Data represented as mean ± SEM.MN: Male Normal. MT: Male Tumor. FN: Female Normal, FT: Female Tumor. N=3 per group.

Gene	Description	MN (RPKM)	FN (RPKM)	FC (FN vs MN)	P-value	
Upregulated						
Tsix	X (inactive)- specific transcript, opposite strand	0.011	8.547	777	0.014	
Sprr1a	small proline-rich protein 1A	3.15	39.293	12.474	0.018	
C1qa	complement component 1, q subcomponent, alpha polypeptide	2.318	28.027	12.091	0.023	
Csf1r	colony stimulating factor 1 receptor	1.03	12.28	11.922	0.014	
C1qc	complement component 1, q subcomponent, C chain	2.935	25.993	8.856	0.0014	
S100a9	S100 calcium binding protein A9 (calgranulin B)	5.096	18.113	3.554	0.005	
Retnlg	resistin like gamma	1.862	6.106	3.279	0.012	
Dnahc1	dynein, axonemal, heavy chain 1	4.994	15.93	3.19	0.004	
Downregulated						
Fam23a	transmembrane protein 236	6.654	0.685	0.103	0.025	
Ddx3y	DEAD box helicase 3, Y- linked	12.885	0.03	0.002	0.028	

Supplementary Table 2. Top 10 DEGs of FN vs MN.

Gene	Description	MT (RPKM)	MN (RPKM)	FT (RPKM)	FN (RPKM)	FC (MT vs MN)	FC (FT vs FN)	P value (MT vs MN)	P value (FT vs FN)
Genes U	pregulated in MT vs	MN and FT	vs FN						
Sprr1a	small proline-rich	60 133	3 15	03 055	30 203	10.00	2 368	0.00	0.00
Зрпта	protein IA	00.133	5.15	93.033	39.293	19.09	2.300	0.00	0.00
Gal	galanin and GMAP	402.83	78 825	697 683	349.2	5 11	1 998	0.00	0.02
Chi3l3	chitinase-like 3	30.434	10.121	45.54	12.853	3.007	3.543	0.00	0.02
ernolo	chemokine (C-X-C	001101	101121	10101	121000	0.001	01010	0.01	0.00
Cxcl10	motif) ligand 10	10.051	4.189	6.454	2.582	2.399	2.5	0.04	0.01
Apod	apolipoprotein D	93.495	39.85	315.912	120.196	2.346	2.628	0.00	0.00
	tissue inhibitor of								
Timp1	metalloproteinase 1	25.59	11.59	50.654	26.675	2.208	1.899	0.00	0.02
	tumor necrosis factor (ligand) superfamily,								
Tnfsf9	member 9	5.612	2.692	10.624	4.364	2.085	2.434	0.04	0.01
Musto 1	musculoskeletal, embryonic nuclear	9 656	4 671	14 756	9 275	1 952	1 702	0.00	0.04
Mustri	myeloid leukemia	0.000	4.071	14.750	0.275	1.000	1.703	0.00	0.04
Mlf1	factor 1	7.528	4.19	8.119	4.21	1.797	1.929	0.01	0.04
Godd45b	growth arrest and DNA-damage-	22.007	12 005	22.240	19 622	1 710	1 730	0.00	0.04
Gauu450	regulated in MT vs N	N and ET v	12.905	52.249	10.022	1.712	1.732	0.00	0.04
Genes op			3 F N						
Fam107a	family with sequence similarity 107 member A	6 914	3 697	3 615	8 328	1 87	0 434	0.03	0.03
i unitoru		0.011	0.001	0.010	0.020	1.07	0.101	0.00	0.00
Sgk1	serum/glucocorticoi d regulated kinase 1	26.42	17.516	12.38	28.117	1.508	0.44	0.00	0.04
Ctgf	cellular communication network factor 2	57.387	29.272	20.413	74.014	1.96	0.276	0.04	0.00
Genes Do	wnregulated in MT v	s MN and U	pregulated F	T vs FN					
_									
Scob1a1	secretoglobin, family 1A, member 1 (uteroglobin)	3.968	8.572	24.357	11.736	0.463	2.075	0.02	0.02
Fabp7	fatty acid binding protein 7, brain	154.989	322.27	847.997	391.936	0.481	2.164	0.00	0.02

Supplementary Table 3. DEGs common to MT vs MN and FT vs FN

Biological Process	DEGs
Second Messenger Signaling	Gal, Mt1, Avpr1a, Irgm1, Igtp, Mt2
IL1-Signaling	Irak3, Rps6ka5, Egr1
Pattern Recognition Signaling	Tifa, Irgm1, Irak3, Igtp
Response to Interferon-Gamma	Irgm1, Igtp
Inflammatory Response	Fas, Timp1, Csf1, chi3l3,Gal, Cxcl10, Serpinb1a,Rps6ka5,Anxa1
Immune Process	Fas, Fst, Tifa, Myc, csf1, Zfp36, Rps19, Nts, cxcl10, Gbp, Fas, Mif1, Irgm1, Igtp, Dll4, tnfsf9,Cebpd,Fos, Hes, Egr1,Anxa1,
Leucocyte Differentiation	Fasn, Csf1, Fas, Dll4, tnfsf9, Fos, Egr1, Anxa1
Leucocyte Migration	Csf1, Rps19, cxcl10, Anxa1
Regulation of Ion Transport	Gem, Fgf23, Usp2, Cckar, Gal, cxcl10, Avpr1a, Sgk1, Per1, Hes1
Autophagosome Assembly	lrgm1
Glial Cell Proliferation	Igtp
Response to Glucocorticoid Stimulus	Zfp36, Fam107a, Fas, Sgk1, Anxa1
Circadian Regulation	Fas, gm129, Per1, Bhlhe40, Egr1
Apoptosis	Timp1, Myc, Phlda1, Zfp36, Mt1, Cxcl10, Ier3, fas, Socs3, Sgk1, Irak3, Gadd45b, Egr1, Anxa1, Spink2, hspa1a

Supplementary Table 4. Biological Processes associated with DEGs upregulated in MT vs MN

Biological Process	DEGs
Chemokine Signaling Pathway	Cxcl2, Cxcl12, Ccl12, ccl11, Pf4, cxcl1, Cxcl10, Ccl7, Ccl4, Ccl3,
MAPK Signaling	Gadd45a, IL1b, Sfpi1, Gadd45b
Lipopolysaccharide Signaling Pathway	Ccl12, Ly86, IL1b, cd14, Sfpi
Interferon Gamma Signaling	Ccl12, cd74, Ccl11, Ccl7, ifitm1, H2-Eb1, ccl4, Rab20, h2Ab1, H2-Aa, Ccl3
IL-1 Signaling	Ccl12, Rbmxrt, II1b, Ccl11, Ccl7, Ccl4, Ccl3
Immune Process	Cd83, Gm8909, C1qc, ccl12, cd74, casp6, Kcnn4, Fcgr2b, Pgf, II1b, Icam1, cxcl1, Scgb1a1,Gal,Cacl10,Hmox1,cd14, ccl7, npy,Sfpi1, BC013712, Apod,Fcer1g, H2-Eb1,tnfsf9,H2-Q6, Lgals1, H2-Ab1, H2-Aa, Ccl3 cmtm3
Humoral Immune Response	Cxcl2. ccl12, Pf4, Cxcl1, Cxcl10, Npy, camp,S100a9,
Leucocyte Migration	Cxcl2, xxl12, ll1b, lcam1, ccl11, Pf4, Spp1, cxcl1, cxcl10,ccl7, Fcer1g, ch25h,ccl4, S100a8,S100a9,ccl3
Antigen Processing and Presentation	Gm8909,cf74, Fcgr2b, Icam1, Fcer1g,H2-Eb1,H2-Q6, H2-Ab1, H2-Aa, Ctss
Inflammatory Response	Timp1, cxcl2, ccl12, Chi3l1, Ly86, Casp6, Il1b,chi3l3,Icam1, ccl11, pf4, cacl1,gal, cxcl10, Hmox1,ccl7, BC013712, CampFcer1g, Ccl4,S100a8,S10a9, Ccl3
Regulation of Cytokine Production	Cd83, chi3l1, cd74, IL1b, chi3l3, Lum, cd14, camp, Fcer1g, Tnfsf9, ccl4,ccl3
RNA Splicing	Rnf113a2, rnf113a1, Exosc5
Apoptosis	Gadd45a, Txndc12, timp1, Ccl12, Cd74, Casp6, Fcgr2b, Glrx, IL1b, Icam1, Spp1, Gal,Rps7, Cxcl10, Hmox1,Nme5, Bag1, Npy, Cryab, Gadd45b, Lgals1,Hspb6,S100a8,Atp5g1,S100a9,Ccl3,Plekhf1
Cell Activation	Cd83, timp1, cd74, Fcgr2b, IL1b, Icam1, Pf4,Scgb1a1, Ptgds, Cxcl10,Npy,Sfpi1, Camp,Pcer1g,Tnfsf9, Lgals1, H2-Ab1
Transport Regulation	Ccl12, cf74, Casp6, Kcnn4, Fcgr2b, Glrx, IL1b, Icam1, Rab13, Spp1, cxcl1,gal,cxcl10, Hmox1, cd14,Car7, Sfpi1,Apod, Aqp2,Cryab, Dcn,Fcer1g,S100a8,Vtn,Fxyd3, Apoc1,Ctss
Positive Regulation of Angiogenesis	Gadd45a, Chi3l1, Dcn, Ngp, Pgf, IL1b, Ccl11, cxcl10,Hmox1, Camp,Hspb6

Supplementary Table 5. Biological Processes associated with DEGs upregulated in FT vs FN

Biological Process	DEGs
Negative Regulation of Angiogenesis	Tspan12, tek, Ctgf, Col18a1, Robo4, Mmrn2, Calcrl,Lef1,Pecam1
Cytoskeleton Organization	Fli1, Gmfb, Tek, Ppm1f, Spry4, Ctgf, Fam107a, Sgk1,Cdh5, Arfip1, Scin, Pecam1
Locomotion	Tek, Ppm1f, Sema3g, Col18a1, Ptprm, Fam107a, Nedd9, Robo4,mmrn2, F2r, Cdh5, Net1,Arrdc3, Mctp1,Patz1,Lef1,Pecam1
Cell Adhesion	Tek, Thsd1,Bmx, Ctgf, Col18a1, Ptprm,Nedd9, Tmem47, Robo4, Mmrn2, Cdh5, net1,Lef1, pecam1,

Supplementary Table 6. Biological Processes associated with DEGs downregulated in FT vs FN