Identification of Trigeminal Sensory Neuronal Types Innervating Masseter Muscle

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Abstract

Understanding masseter muscle (MM) innervation is critical for the study of cell-specific mechanisms of pain induced by temporomandibular disorder (TMDs) or after facial surgery. Here, we identified trigeminal (TG) sensory neuronal subtypes (MM TG neurons) innervating MM fibers, masseteric fascia, tendons, and adjusted tissues. A combination of patch clamp electrophysiology and immunohistochemistry (IHC) on TG neurons back-traced from reporter mouse MM found nine distinct subtypes of MM TG neurons. Of these neurons, 24% belonged to non-peptidergic IB-4\(^{-}/\)TRPA1\(^{-}\) or IB-4\(^{+}/\)TRPA1\(^{+}\) groups, while two TRPV1\(^{+}\) small-sized neuronal groups were classified as peptidergic/CGRP\(^{+}\). One small-sized CGRP\(^{+}\) neuronal group had a unique electrophysiological profile and were recorded from Nav1.8\(^{-}\) or trkC\(^{-}\) neurons. The remaining CGRP\(^{+}\) neurons were medium-sized, could be divided into Nav1.8\(^{-}\)/trkC\(^{-}\) and Nav1.8\(^{+}\)/trkC\(^{-}\) clusters, and showed large 5HT-induced current. The final two MM TG neuronal groups were trkC\(^{-}\) and had no Nav1.8 and CGRP. Among MM TG neurons, TRPV1\(^{+}/\)CGRP\(^{-}\) (somatostatin\(^{+}\)), tyrosine hydroxylase (TH\(^{+}\)) (C-LTMR), TRPM8\(^{+}\), MrgprA3\(^{+}\), or trkB\(^{+}\) (A\(^{+}\)-LTMR) subtypes have not been detected. Masseteric muscle fibers, tendons and masseteric fascia in mice and the common marmoset, a new world monkey, were exclusively innervated by either CGRP\(^{-}/\)NFH\(^{+}\) or CGRP\(^{+}/\)NFH\(^{-}\) medium-to-large neurons, which we found using a Nav1.8-YFP reporter, and labeling with CGRP, TRPV1, neurofilament heavy chain (NFH) and pgp9.5 antibodies. These nerves were mainly distributed in tendon and at junctions of deep-middle-superficial parts of MM. Overall, the data presented here

Significance Statement

Identification of sensory neuron subtypes innervating masseter muscle (MM) will enable the study of cell-specific mechanisms of masticatory myofascial pain, including temporomandibular disorder (TMD) and after restorative surgeries involving MM. Combining back tracing from MM, patch-clamp electrophysiology, and immunohistochemistry (IHC) with sensory neuronal markers on mouse and nonhuman primate tissues, we identified trigeminal (TG) neuronal groups innervating MM (MM TG neurons). MM and adjacent tissues are innervated by 9 distinct types of TG neurons, some of which are substantially different from L3–L5 DRG neurons. However, MM fibers, tendon, muscle-tendon junction, and fascia in mice and common marmosets are exclusively innervated by medium-to-large neurons. These neurofilament heavy chain (NFH)-positive sensory nerve fibers are mainly distributed in tendon and at junctions between deep, middle, and superficial parts of MM.
demonstrates that MM is innervated by a distinct subset of TG neurons, which have unique characteristics and innervation patterns.

Key words: masseter muscle; myogenous temporomandibular disorders; orofacial pain; sensory neurons; trigeminal

Introduction

Myofascial pain is widely prevalent in the general population (Cimmino et al., 2011). Pain related to mastication muscles, which are comprised of the temporalis, medial pterygoid, lateral pterygoid, and masseter muscle (MM), is referred to as masticatory myofascial pain. Masticatory myofascial pain may manifest as facial pain, headache, soreness, and fatigue of the masticatory muscles (Gerwin, 2001). This pain type is detected among patients with temporomandibular disorders (TMDs; Lobbezoo et al., 1996; Slade et al., 2016), myofascial pain syndrome (Desai et al., 2013; Galasso et al., 2020), and after craniofacial surgeries done to repair facial paralysis, facelift procedures, or tissue trauma affecting MM (Wieckiewicz et al., 2015; Lassus et al., 2018).

MM consists of three heads (superficial, middle, and deep) of muscle fibers, masseteric fascia, and tendons; and is innervated by the masseteric nerve, a branch of the anterior portion of the mandibular division (V3) of the trigeminal (TG) nerve (Fig. 13A,B). Anatomical studies have revealed myofascial trigger points, which represent tender muscle areas that can elicit pain whenever stimulated during normal or pathologic condition. These myofascial trigger points coincide with neuromuscular junctions at the innervation zone between MM heads (Procopio Pinheiro et al., 2020). Previous observations of the head and neck area showed the presence of nociceptive unmyelinated C- and myelinated Aβ fibers (Strassman et al., 1996; Strassman and Raymond, 1999), as well as low-threshold mechanoreceptors (LTMRs) consisting of myelinated Aδ and Aβ fibers (Lobbezoo et al., 2002). Similarly, it has been shown that MM has C-fibers, two types of A-δ high-threshold mechanoreceptors (Aδ-HTMR) as well as Aβ-LTMR fibers (Nishimori et al., 1986; Cuccu et al., 1989; Connor et al., 2005; Wong et al., 2014; Sato et al., 2018). MM is predominantly innervated by Aβ-HTMR, with cell bodies observed to be 27% small, 49% medium, and 24% large-sized neurons (Sato et al., 2018). However, the precise composition of MM innervating TG ganglion neuronal groups or their properties is unknown.

The understanding of molecular mechanisms governing masticatory myofascial pain is hindered by the limited information on the neuronal circuits undergoing plasticity during TMD and after surgical procedures involving MM and/or masseteric nerve. The aim of the present study has been to identify and comprehensively characterize TG sensory neuronal groups innervating MM (MM TG neurons), and to examine the innervation patterns of MM by afferent sensory fibers. To do so, back-tracing of sensory neurons from MM, patch-clamp electrophysiology, Ca imaging, anatomic studies, sensory neuronal reporter mice and tissues from a nonhuman primate species (common marmosets) have been used (Herrity et al., 2014; Patil et al., 2018).

Materials and Methods

Animals

All animal experiments conformed to American Pain Society (APS)’s Guiding Principles in the Care and Use of Vertebrate Animals in Research and Training. We also followed guidelines issued by the National Institutes of Health (NIH) and the Society for Neuroscience (SfN) to minimize the numbers of animals used and their suffering. Protocols used in this study (201901114AR for mice and 20200021AR for nonhuman primates) are approved by the Institution Animal Care and Use Committee (IACUC).

Experiments were performed on 10- to 18-week-old male mice. B6.Cg-Gt(Rosa26)Sortm9(CAG-tdTomato)Hze/J (Ai9; tdTomato; stock 007909), B6.Cg-Gt(Rosa)26Sor232(CAG-COP4* H134R/EGFP)Hze/J (Ai32; stock 024109), B6;129P2-Pvalbtm1(cre)Arbr/J (PV-cre; stock 008069) and Mrgrpdr1creERT2(Wol/J (MrgrpD-cre-ER; stock 031286) mouse lines were obtained from The Jackson Laboratory. 5HT3a-GFP (also known as Htr3a-EGFP) and TRPV1-GFP transgenic mouse lines were purchased from the GENSAT program (MMPRC services; UNC, NC and UC Davis, CA, respectively). The CGRPcreERT2 mouse line was kindly provided by Pao-Tien Chuang (University of California San Francisco, San Francisco, CA). Nav1.8creERT2 mouse line was kindly provided by John Wood (University College London, London, United Kingdom). The trkCcreERT2 mouse line was generated in David Ginty’s laboratory (Harvard...
Medical School, Boston, MA) and kindly provided by Yu Shin Kim (The University of Texas Health Science Center at San Antonio). In inducible cre-carrying mouse lines, cre-recombinase was induced in six- to eight-week-old mice by three consecutives (every second day) intraperitoneal injections of 100 mg/kg tamoxifen (dissolved in corn oil). Cre-recombination occurs within two to three weeks after tamoxifen injection. Two aged (11- and 15-year-old) male common marmosets (Callithrix jacchus) were used for collection of MM. IACUC and veterinary oversight regularly monitored marmoset housing and animal conditions to ensure all guidelines for the health and safety of the animals were met and research was conducted in compliance with the United States Public Health Service's Policy on Humane Care and Use of Laboratory Animals and the Guide for the Care and Use of Laboratory Animals and adhered to the American Society of Primatologists (ASP) principles for the ethical treatment of non-human primates. Marmosets were euthanized for humane reasons following veterinary consult and tissue samples were collected immediately following determination of death.

Primary TG neuronal culture

To visualize MM TG neurons, wild-type or reporter mice expressing GFP, YFP (Ai32), or tdTomato (Ai9) gene were injected into the right and left MM closer to tendon with 10 μl of WGA-488 or WGA-555 (50 μg in 0.5% DMSO) back-tracer. Neuronal culture for electrophysiology recordings was always generated from one mouse. TG was dissected 24–36 h post-WGA injections, and sensory neurons were cultured as previously described (Belugin et al., 2013). Cells were maintained in DMEM with 2% fetal
bovine serum, 2 mM L-glutamine, 100 U/ml penicillin, and 100 μg/ml streptomycin. No growth factor was added to the media. The experiments were performed within 24 h after TG neuron plating. This culture conditions changes in sensory neurons (Patil et al., 2018).

Electrophysiology: recording

Before patch clamp recording, cultured TG cells from WGA-488-injected wild-type mice were stained for 0.5–4 h with IB-4 Alexa Fluor 594 (1:1000; Thermo-Fisher Scientific). WGA+/IB-4+ TG neurons were selected for recording (Fig. 1A). Along with WGA+/IB-4+ TG neurons from wild-type mice, WGA+/CGRP-cre-ER+, WGA+/TRPV1-GFP+, WGA+/5HT3a-GFP+, WGA+/PV-cre+, WGA+/trkC-cre-ER+, WGA+/Nav1.8-cre+, and WGA+/Nav1.8-cre− MM TG neurons were selected for recordings. Overall, we have chosen reporters that are not affected by early developmental changes (Patil et al., 2018).

Recordings were made in patch clamp whole-cell voltage [holding potential (Vh)] of –60 mV) or current clamp configurations at room temperature. Data were acquired using an Axopatch 200B amplifier and analyzed with pCLAMP10.6 software (Molecular Devices). Recording data were filtered at 0.5–5 kHz and sampled at 2–20 kHz depending on current kinetics. Borosilicate pipettes (Sutter) were polished to resistances of <5 MΩ. If required, access resistance (R0) was compensated (40–80%) to the value of ≪6–8 MΩ. Data were rejected when R0 changed >20% during recording, leak currents were >100 pA, or input resistance was <300 MΩ. Liquid junction potential (LJP) was not corrected using the equation: Vmeter = Vcell + LJP, since large anions were not included in extracellular or pipette recording solutions. Recordings were made 3–4 min after establishing whole-cell configuration to allow for equilibrium. Currents were considered positive when their amplitudes were fivefold bigger than displayed noise (in root mean square).

Standard external solution (SES) contained the following: 140 mM NaCl, 5 mM KCl, 2 mM CaCl2, 1 mM MgCl2, 10 mM D-glucose, and 10 mM HEPES, pH 7.4. The standard pipette solution (SIS) contained the following: 140 mM KCl, 1 mM MgCl2, 1 mM CaCl2, 10 mM EGTA, 10 mM D-glucose, and 10 mM HEPES, pH 7.3, 2.5 ATP and 0.2 GTP. Drugs were applied by a fast, pressure-driven and computer controlled four-channel system (ValveLink8; AutoMate Scientific) with quartz application pipettes. Recordings were independently performed by two investigators. Data were accumulated from five to eight independent MM TG neuronal cultures for each mouse line. Six to 15 neurons were recorded from each MM-TG neuronal culture.

Electrophysiology: recording protocols and data analysis

On the selected for recording MM TG neurons, we used a sequence of protocols after fast and slow capacitance compensations: (1) single action potential (AP) in current-clamp configuration was generated with 1 nA (2 nA for >40-pF cells) 0.5-ms current pulse (Fig. 1B, C; Petruska et al., 2000; Patil et al., 2018); (2) AP train was induced by applying step currents of 50–550 pA with 100-pA increment for 1 s (Fig. 1D) or 200–2000 pA with 300-pA increment for medium-to-large neurons; (3) after current clamp recordings the electronics were switched to voltage-clamp configuration (Vh = –60 mV) and ATP (30 μM) current was recorded by applying drug for 5 s; (4) after ATP-gated current, 5HT (30 μM)-gated current was recorded by 5-s-long drug application; (5) the next protocol in a voltage-clamp configuration was a step down from Vh to –80 mV kept for 500 ms, and then 200-ms depolarizing command steps (20 mV) were applied from –40 mV to a final potential of +20 mV (Fig. 1E; Petruska et al., 2000). Time gap between these successive one to five protocols was 1–2 min. In some sets of experiments, mustard oil (MO; 25 μM) and menthol (Ment; 100 μM) responses in MM TG neurons of reporter mice were evaluated using Ca2⁺ imaging system as previously described (Salas et al., 2009; Patil et al., 2018).

Data on sensory neuronal marker expression in MM TG neurons, capacitance (in pF) and resting membrane potential (RMP; Vm in mV) values were collected prior recording and after application of protocol-1 (Fig. 1A, B). Cells were considered as a spherical, and therefore, diameter (d in μm) of cells was calculated from capacitance (Cm in pF) values using the following:

\[ d = 5 \times \sqrt{(Cm/4 \pi)} \]

AP duration at the base (dB; time from Vm starting point to Vm levels at falling phase of AP), after-hyperpolarization (AHP) peak as a distance from RMP to a lowest point of (AHP) peak as a distance from Rm to Vm levels at the base of the command steps (20 mV) were applied from –40 mV to a final potential of +20 mV (Fig. 1E; Patruska et al., 2000). Time gap between these successive one to five protocols was 1–2 min. In some sets of experiments, mustard oil (MO; 25 μM) and menthol (Ment; 100 μM) responses in MM TG neurons of reporter mice were evaluated using Ca2⁺ imaging system as previously described (Salas et al., 2009; Patil et al., 2018).

Analysis of protocol-2 gave AP activation threshold (in pA) and decay tau (τ; ms) calculation was performed using pCLAMP10.6 software (Fig. 1E). Shape of current (I), including presence or absence of “spike-like” feature at steps to 0 and +20 mV, was an important clustering variable (Figs. 3F, 4C). Clustering is based on at least two unique features for a specified cluster (Table 2). An approach for generation of clustering parameters described in detail in Results and in the previous publication (Patil et al., 2018).

Immunohistochemistry (IHC)

Wild-type male C57/B1 mice were injected into the right and left MM close to the tendon with 10 μl of WGA-488
Table 1: Properties of MM TG neuronal groups

| Gr  | Marker | N  | Size, μm² | RMP, mV | dP, ms | Shape ¹ | AHPm, ms | AHP peak² | AP Thr., pA | AP train, Hz | ATP | SHT, pA | Tau, ms | Shape Cur ² |
|-----|--------|----|-----------|---------|--------|---------|----------|------------|-------------|-------------|------|---------|---------|-------------|
| S1  | IB4    | 16 | 20.5 ± 0.8 | -40.4 ± 2.1 | 11.9 ± 1.1 | Bow     | 35.2 ± 4.9 | -14.5 ± 1.2 | 405 ± 177   | 10 ± 2.2    | 0    | 0       | 19.2 ± 6.5 | Smooth    |
|     | Nav1.8 |    |           |         |        |         |          |            |             |             |      |         |         |             |
| S2  | IB4    | 52 | 19.9 ± 0.4 | -41.3 ± 1.0 | 11.5 ± 0.6 | Deflection | 67.9 ± 4.0 | -19.0 ± 0.8 | 156.6 ± 20.3 | 8.3 ± 0.8    | 0    | 0       | 26.3 ± 4.2 | Smooth², ³ |
|     | Nav1.8 |    |           |         |        |         |          |            |             |             |      |         |         |             |
| S3  | trpV1  | 47 | 21.4 ± 0.4 | -44.7 ± 1.0 | 8.7 ± 0.3 | Hump    | 76.3 ± 17.2 | -13.7 ± 0.9 | 109.1 ± 23.4 | 11.8 ± 2.2   | 0    | 0       | 20.6 ± 2.6 | Smooth⁷ |
|     | CGRP   |    |           |         |        |         |          |            |             |             |      |         |         |             |
|     | Nav1.8 |    |           |         |        |         |          |            |             |             |      |         |         |             |
| S4  | trpV1  | 24 | 21.4 ± 0.9 | -43.0 ± 1.1 | 8.5 ± 0.5 | Deflection | 63.5 ± 20.5 | -10.3 ± 1.4 | 64.3 ± 14.3 | 6.3 ± 1.3    | 0    | 0       | 16.9 ± 2.1 | Smooth⁶ |
|     | CGRP   |    |           |         |        |         |          |            |             |             |      |         |         |             |
|     | Nav1.8 |    |           |         |        |         |          |            |             |             |      |         |         |             |
| S5  | CGRP   | 30 | 21.5 ± 1.0 | -44.7 ± 1.5 | 7.7 ± 0.5 | Straight | 65.2 ± 16.7 | -13.7 ± 1.3 | 50.0 ± 0     | 22 ± 3.8     | 0    | 195 ± 18 | 24 ± 4.6 | Box like⁶  |
|     | 5HT3a  |    |           |         |        |         |          |            |             |             |      |         |         |             |
|     | trkC    |    |           |         |        |         |          |            |             |             |      |         |         |             |
| M1  | CGRP   | 34 | 29.5 ± 0.9 | -52.1 ± 1.1 | 6.3 ± 0.2 | Deflection | 92.4 ± 11.7 | -13.2 ± 1.2 | 316 ± 107   | 10 ± 3.7     | 0    | 818 ± 100 | 3.5 ± 0.6 | Box like⁶  |
|     | 5HT3a  |    |           |         |        |         |          |            |             |             |      |         |         |             |
|     | trkC²  | 66 | 30.7 ± 0.5 | -51.6 ± 1.6 | 4.8 ± 0.2 | Straight | 73.6 ± 9.1  | -13.3 ± 0.5 | 580 ± 128   | 13.1 ± 3.3   | 0    | 774 ± 88 | 22.0 ± 2.3 | Spike    |
|     | 5HT3a  |    |           |         |        |         |          |            |             |             |      |         |         |             |
|     | trkC⁵  |    |           |         |        |         |          |            |             |             |      |         |         |             |
|     | PV     |    |           |         |        |         |          |            |             |             |      |         |         |             |
| M3  | trkC⁶  | 56 | 33.5 ± 0.6 | -55.2 ± 0.6 | 2.8 ± 0.1 | Straight | 86.3 ± 12.0 | -10.3 ± 0.6 | 1280 ± 152  | 7 ± 2.3      | 0    | 0       | 22.3 ± 1.9 | Spike¹⁰ |
|     | PV     |    |           |         |        |         |          |            |             |             |      |         |         |             |
| M4  | trkC⁷  | 40 | 30.5 ± 0.8 | -54.0 ± 0.7 | 1.9 ± 0.06 | Straight | 6.9 ± 0.6  | -10.7 ± 0.8 | 0           | 0           | 0    | 0       | 16.0 ± 2.7 | Box like²  |
|     | PV     |    |           |         |        |         |          |            |             |             |      |         |         |             |

¹ Green font is non-peptiderigic; red font is peptiderigic, and blue font is mechanoreceptors.
² Size in μm is calculated from pF (see Materials and Methods).
³ Characteristic feature on AP downstroke: bow, hump, deflection, and straight (Figs. 1B, C, 2A).
⁴ AHPm is a lowest point of AP, and it is measured in negative mV.
⁵ Characteristic feature of the last (generated by stepping to +20 mV) outward current: “smooth,” ”box like,” and “spike” (Figs. 1E, 2D, E, 3E, F, 4C, D).
⁶ Currents do not have inward component.
⁷ Neurons are responsive to MO.
⁸ 5HT3a is encoded by Htr3a gene.
⁹ Contain low level Nav1.8.
¹⁰ Spike is sharper in M3 than M2.
¹¹ Some (~15%) M3 neurons respond to 5HT (30 μM).

(50 μg in 0.5% DMSO) back-tracer. TG was dissected at 2 d post-WGA injection from 4% paraformaldehyde-perfused mice. MM was dissected from WGA un-injected wild-type mice. TG and MM were collected from two aged marmosets (11 and 15 years of age) that were not injected with WGA. TG and MM from both species were additionally fixed with 4% paraformaldehyde for 15 min, cryo-protected overnight with 30% sucrose in phosphate buffer, embedded in Neg 50 (Richard Allan Scientific), and 25-μm transverse cryo-sections were generated from both mouse and marmoset TG and MM as previously described (Belugin et al., 2013). IHC was conducted as described (Belugin et al., 2013). IHC was simultaneously performed on 6–12 sections generated from three or four mice; or three to four sections generated from the marmoset tissue. The following previously characterized primary antibodies were used for mouse sections: anti-neurofilament heavy chain (NFH) chicken polyclonal antibodies (BioLegend, catalog #PCK-592P, 1:400; Zappulo et al., 2017); anti-pgp9.5 (Millipore-Sigma, catalog #AB1761-1, 1:1000; Roy et al., 2020); anti–TRPV1 guinea pig polyclonal (Neuromics, catalog #GP14100, 1:700; Patil et al., 2014); anti–CGRP rabbit polyclonal (Sigma, C8198, 1:300; Ruparel et al., 2008; Neeb et al., 2011; Lorenzo et al., 2014); anti-tyrosine hydroxylase (TH) rabbit polyclonal (Pel-Freez, P40101, 1:400; Dunkley et al., 2004; Ma et al., 2004); anti-mrgD rabbit polyclonal (Alamone Lab, AMR-061, 1:200; Chang et al., 2016; Patil et al., 2018); anti-trkC goat polyclonal (R&D Systems, AF1404, 1:200; Arcourt et al., 2017; Patil et al., 2018); anti-trkB goat polyclonal (R&D Systems, AF1494, 1:200; Ninkina et al., 1997; Arcourt et al., 2017); rabbit anti-parvalbumin (Swant, PV25, 1:500; Schurmans et al., 1997; Schwaller et al., 1999; Arcourt et al., 2017); and rabbit anti-Calbindin D28k (Swant, CB-38a, 1:500; Schurmans et al., 1997; Arcourt et al., 2017). Antibodies for MrgprD and PV produced weaker signals compare to other antibodies and GFP/YFP/tdTomato signals, hence their intensities were increased, and backgrounds were reduced. Anti-NFH and anti-pgp9.5, but not anti-CGRP produced clear IHC signals in marmoset sections (Resnikoff et al., 2019). Sections were incubated with species appropriate Alexa Fluor secondary antibodies (1:200; Invitrogen).

Images were acquired using a Keyence BZ-X810 All-in-One Fluorescent Microscope (Keyence) or a Nikon Eclipse 90i microscope (Nikon Instruments) equipped with a C1si laser scanning confocal imaging system. Images were processed with NIS-elements (Nikon Instruments) and Adobe Photoshop CC 2019 software. Gain setting was constant during acquisition, and it was established on no primary control slides. Control IHC was performed on tissue sections processed as described but either lacking primary antibodies or lacking primary and secondary antibodies. Cell counts from
IHC images acquired as Z-stuck were performed using ImageJ software. Cells positive for WGA and each marker as well as the combinations of markers were counted. We used three or four independent mice (n = 3–4) to generate sections and counted three to five sections per mouse. Mean values from these three to five sections represented n of 1.

Statistical analysis

GraphPad Prism 8.0 (GraphPad) was used for statistical analyses. Data in the figures are mean ± SEM, with n referring to the number of mice used for IHC and the numbers of analyzed recorded cells. Differences between IHC and electrophysiologically characterized groups were assessed by unpaired t test or regular one-way ANOVA with Tukey’s post hoc tests, each column was compared with all other columns. A difference is accepted as statistically significant when p < 0.05. Interaction F ratios and the associated p values are reported.

Results

Sensory neurons have previously been identified and classified using multiple methods including, back-tracing from the target tissue (da Silva Serra et al., 2016), sensory neuron reporter mice, patch-clamp recording and classification according to AP properties, sensitivity to algic agents and appearances of a variety of voltage-gated currents (Xu et al., 2010; Li et al., 2011; Patil et al., 2018) and IHC (Patil et al., 2018). Described in Materials and Methods, back-tracing from MM will label TG neurons innervating MM fibers, tendon, muscle-tendon junction, and massteric fascia. This approach could have two drawbacks. First, along with these MM structures, diffusion of WGA injected into MM led to labeling of adjusted tissues such as masseteric nerve fibers and the subcutaneous layer of facial skin (Fig. 1A, B). Second, WGA produces nonuniform size-dependent labeling of TG neurons (Robertson, 1990). This nonuniform labeling of TG neurons is unlikely miss entire neuronal group since there is a variation in neuronal sizes within every TG neuronal group. Patch clamp experiments from these WGA-labeled MM TG neurons will yield data on 12 variables: cell size, RMP, AP width (dB), characteristic features of AP shapes, AHP-peak, AHP80, responsiveness to 5HT, ATP, capsicain (CAP), MO, and Ment, τ (tau) from fitting of voltage-gated currents (l), and shape of these I, including presence or absence of a spike-like feature on outward portions of these voltage-gated currents (Table 1).

Figure 2. Recordings from TRPV1⁺ MM TG neurons. A, In MM TG neurons belonging to the group S3, a single AP in WGA⁺/TRPV1⁺ neuron is generated by current pulse (protocol-1), which has characteristic hump on down stroke phase of AP. B, In MM TG neurons belonging to the group S4, a single AP in WGA⁺/TRPV1⁺ neuron has distinctive deflection on down stroke phase of AP. C, Current-evoked AP train from a WGA⁺/TRPV1⁺ TG neuron belonging to the group S4. Depicted AP train is evoked by a 50-pA step lasting 1 s. D, In WGA⁺/TRPV1⁺ TG neurons belonging to the group S3, currents were generated by waveforms from Figure 1E. E, In WGA⁺/TRPV1⁺ TG neurons belonging to the group S4, currents generated by waveforms from Figure 1E do not have inward component.
IB4⁺ MM TG sensory neuronal groups

Cultured TG neurons from wild-type male mice back-traced with WGA-488 from MM were stained with IB-4⁻, Nav1.8⁺, and Nav1.8⁻ MM TG neurons to MO (25 μm) and Ment (100 μm) measured with Ca²⁺ imaging. We selected approximately equal numbers (48 from 62) of small-sized (<30 pF) and medium-to-large-sized (>30 pF) neurons for recording. A majority (48 from 62) of MM TG neurons could be differentiated from S4 because of the following: only S3 neurons had a large inward component (compare Fig. 2D) and τ values (Table 1). We found S3 TRPV1⁺ MM TG neurons were differentiated from S4 because of the following: only S3 neurons were responsive to MO (Fig. 3A); S3 had a hump (more pronounced than deflection) on AP (Fig. 2A), while S4 had a deflection (Fig. 2B); and unlike S4, current (I) in S3 neurons had a large inward component (compare Fig. 2D,E). S3 and S4 are dissimilar from other types of TG neurons back-traced from MM in expression of TRPV1 and responsiveness to CAP (Table 2). In summary, there are two subtypes of TRPV1⁺ small-sized TG neurons labeled by WGA injected into MM.

CGRP⁺ MM TG sensory neuronal groups

A total of 122 CGRP⁺ MM-TG neurons were recorded and analyzed. We selected approximately equal numbers of small-sized (<30 pF) and medium-to-large-sized (>30 pF) neurons for recording. A majority (48 from 62) of small-sized (<30 pF) WGA⁻/CGRP⁺ neurons were CAP responsive and according to their properties, could be classified as either S3 or S4 (Table 1). This indicated that...
CGRP

M3 group neurons recorded either from a MM TG trkC

1.8 (TRPM8 agonist), and inward component (Fig. 3)

generated an AP train even after 50-pA current injection (Fig. 3)

AHP80, presence of spike on the I (Figs. 3, 4C) and requirement of large current to generate AP train were similar for M3 versus M2 neurons (Table 1). An exception was AP dB, which was significantly narrower for M3 compared with M2 neurons (t test; t = 9.622 df = 125; p < 0.0001; Fig. 4A).

The most critical parameter distinguishing M3 from M2 was their responsiveness to 5HT, which was detected exclusively in M2 neurons (Tables 1, 2). M4 neurons had several unique properties distinguishing them from other MM TG neurons. First, no AP train could be generated even with an injection of 2-nA current (protocol-2). Second, M4 neurons were fastest and had the smallest dB value (one-way ANOVA; F(8,279) = 6.398; p < 0.0001; Fig. 4B; Table 1). Third, AHP80 was fastest in M4 neurons (one-way ANOVA; F(8,279) = 6.398; p < 0.0001; Fig. 4B; Table 1). Fourth, the I from M4 neurons had a “box-like” shape but possessed a small spike (Fig. 4D). We also recorded algic currents from medium-to-large sized MM TG neurons. These neurons did not have ATP, MO, and Ment-gated currents (Fig. 3A; Table 1).

Next, using reporter mice, we recorded and analyzed 65 trkC+, 32 PV+, and 35 medium-to-large sized Nav1.8+ MM TG neurons. This information revealed that trkC was expressed by S5, M2–M4 neurons; PV was present in M2–M4 groups; and Nav1.8 was mainly detected in S1–S4.

**Table 2: Clustering parameters for MM TG neuronal groups**

| Gr | Key clustering parameters | Putative function |
|----|---------------------------|-------------------|
| S1 | IB4+/CGRP+; non-responsive to MO and CAP | Non-peptidergic; C-nociceptor |
| S2 | IB4+/CGRP+; responsive to MO but not to CAP | Non-peptidergic; C-nociceptor |
| S3 | TRPV1+/CGRP+; AP hump | Peptidergic; C-nociceptor |
| S4 | TRPV1+/CGRP+; AP deflection; no inward current | Peptidergic; C-nociceptor |
| S5 | CGRP+/SHT3a+/Nav1.8+; AP straight; box-like current; small response to 5HT | ? |
| M1 | CGRP+/Nav1.8+; AP deflection; box-like current; big response to 5HT | Aβ-nociceptors |
| M2 | CGRP+/trkC+/Nav1.8+; AP straight; spike on current; big response to 5HT | Aβ-nociceptors |
| M3 | trkC+/CGRP+/Nav1.8+; AP fast and straight; sharp spike on current; non-response to 5HT | Aβ-mechanoreceptor |
| M4 | trkC+/CGRP+/Nav1.8+; fastest AP; unique AHP; box-like current; non-response to 5HT | Aβ-mechanoreceptor |
S4 groups. Medium-to-large sized Nav1.8+ MM TG neurons had weaker YFP signal than Nav1.8+ small-sized neurons. A bulk majority of medium-to-large sized Nav1.8+ MM TG neurons belonged to the M1 group, but four of 35 were classified into the M2 group. Since M1 and M2 neurons were recorded among Nav1.8+ and Nav1.8- neurons, this suggests that they had weak Nav1.8 expression, which was sufficient to drive Ai32 reporter in some, but not for all M1 and M2 neurons. Figure 13C summarizes clustering of TG neurons labeled by back-tracing from MM. These clusters were generated on this basis of electrophysiological properties obtained from recordings on TG neurons isolated from different reporter mouse lines. Unlike previously characterized L3–L5 DRG neurons, which innervate the skin and muscles of legs, TG neurons labeled by back-tracing from MM did not have small-sized TRPV1+/CGRP- (somatostatin-) neurons (Usoskin et al., 2015; Sharma et al., 2020), TRPM8+ (cold thermoceptors) neurons (Sharma et al., 2020) and strongly responding to ATP MrgrpA3+ neurons (Usoskin et al., 2015; Patil et al., 2018). MM TG neurons also did not have multiple small-sized peptidergic groups (Patil et al., 2018; Zeisel et al., 2018). Expectedly, MM TG lacked neuronal groups associated with innervation of hairs: C-LTMR and Aδ-LTMR, which are Nav1.8+/CGRP-/trkC- (Usoskin et al., 2015) and have distinct AP (Patil et al., 2018). Finally, MM TG neurons do not have proprioceptors (Sharma et al., 2020), which are located in a brain stem region of the TG system. Overall, MM TG neuronal groups were found to be substantially different from the well-characterized L3–L5 DRG neuronal clusters, and S5 and M2 groups have no analogs among L3–L5 DRG neurons (see Discussion).

IHC analysis of MM TG sensory neuronal groups

To further characterize the MM TG sensory neuronal groups and confirm our electrophysiology data, we used IHC to examine expression of sensory neuronal markers in TG neuron sections from naive male mice injected with WGA-488 into MM. Peptidergic MM TG neurons (S3–S5, M1, and M2 groups) labeled with CGRP antibodies composed 51.7 ± 2.1% of WGA+ TG neurons (Figs. 5A,A', 8C). Most 15.3 ± 2.4% of WGA+ TG neurons were TRPV1+ (S3 and S4 groups) and all of these neurons were labeled with CGRP (Figs. 5B,B', 8C, 13C). Consistent with our electrophysiology data, TRPV1+/CGRP+ TG neurons, which have strong TRPV1 labeling (Fig. 5B,B', brown arrows), did not contain WGA.

Small-sized non-peptidergic (i.e., IB4+/CGRP-) DRG neurons express MrgrpD as a marker (Reynolds et al., 2015; Patil et al., 2018). However, despite 24 ± 2.4% of WGA- cultured TG neurons staining for IB-4 (S1 and S2 groups), only a few (3.1 ± 0.7%, n = 3) MM TG neurons expressed MrgrpD (Figs. 6A,A', 8C). Anti-mrgrpD antibodies are weak and generate high background (Fig. 6A,A'); therefore, we used an alternative approach to confirm these data by back-tracing WGA from MM in MrgrpD-cre/ER/tdTomato reporter mice. Analysis of these sections confirmed that the MrgrpD-cre+ signal was only detected in a few TG MM neurons (2.3 ± 0.6%, n = 3; Fig. 6B,B'). However, some of MrgrpD-cre+ signals were registered in weakly labeled WAGA- neurons (Fig. 6B,B'). These data indicate that MrgrpD may not be an appropriate marker for non-peptidergic MM TG neurons.

Single-cell transcriptomic of L3–L5 mouse DRG neurons assigned calbindin D28-positive (Calb+) neurons to group NF2 (Usoskin et al., 2015) or Aβ-Field (Sharma et
Functional studies on L3–L5 DRG Calb− neurons demonstrated that they are a subset of Aβ-LTMR neurons (Aarcourt et al., 2017). TG MM neurons express Calb at low levels (∼2%; Figs. 7A, A′, 8C). Recording from MM TG neurons isolated from PV-cre/tdTomato reporter mice showed that PV could be present in M2–M4 groups. IHC data indicate that PV antibodies had low signal in ∼21% MM TG neurons (Figs. 7B, B′, 8C). Similarly, electrophysiology data on MM TG neurons from trkC/tdTomato mice indicated that trkC could be found in groups S5, M2–M4. TrkC antibodies labeled ∼30% WGA+ TG neurons (Fig. 8A, A′, C). This implies that S5 are ∼5% of MM TG neurons (Fig. 13C). Our patch-clamp recordings did not show the presence of C-LTMR-like neurons.

Figure 6. Expression of MrgprD in MM TG neurons. A, A′, Expression of MrgprD (mrgD; red) neurons in a TG section containing MM TG neurons (WGA; green). Objective is 20×. B, B′, Expression of MrgprD-cre/tdTomato (mrgD-cre; red) neurons in a TG section from mice back-traced from MM with WGA-488 (WGA; green). Objective is 20×. MM TG neurons containing both mrgD or mrgD-cre and WGA are marked with yellow arrows. MM TG neurons containing WGA, but not mrgD or mrgD-cre or trpV1, are marked with cyan arrows. White horizontal bar shows 20-μm scale for each panel.

Figure 7. Expression of calbindin-D28 and parvalbumin mechanoreceptor markers in MM TG neurons. A, A′, Expression of calbindin D28 (Calb; red) in a TG section containing MM TG neurons (WGA; green). B, B′, Expression of parvalbumin (PV; red) in a TG section containing MM TG neurons (WGA; green). MM TG neurons lacking Calb or PV are shown with cyan arrows. MM TG neurons containing Calb or PV are shown with yellow arrows. For all panels, objective is 20×. White horizontal bar shows 20-μm scale for each panel.
nerves that innervate MM fibers, tendon, muscle-tendon junction, and masseteric fascia. Injection into MM could lead to contamination and/or diffusion of WGA into subcutaneous tissue (Fig. 13B). Additionally, such an injection could label sensory afferent fibers in the main trunk of masseteric nerve (Fig. 13A,B). Using IHC, we examined the distribution of different sensory fiber types within MM, as well as innervation of muscle fibers, tendon, muscle-tendon junctions, and fascia surrounding whole MM.

The main trunk of the masseteric nerve branches out from the mandibular nerve and enter the MM in the sigmoid fascia area (Kim et al., 2010; Fig. 13A,B). We found this area of masseteric nerve trunk contained CGRP+/NFH− (yellow arrows; S3–S5 groups), CGRP+/NFH− (M1 and M2 groups; Fig. 9A,A′, cyan arrows) and CGRP+/NFH− (M3 and M4 groups; Fig. 9A,A′, white arrows) sensory nerves. The masseteric nerve trunk then descends into MM between the middle and deep parts of muscle fibers and then divides into many nerve branches in the posterosuperior, posteroinferior, anterosuperior, and anteroinferior directions (Luo et al., 1991; Kim et al., 2010; Fig. 13A,B). We observed that multiple branches of CGRP+ and NFH+ nerve bundles reached the deep and middle layers of the muscle and junctions of muscle fibers and tendon (Fig. 9B). Distribution of CGRP+ and NFH+ nerve bundles and individual fibers occurred throughout MM. However, they mainly concentrate in tendon and at junctions between superficial, middle, and deep parts of MM (Fig. 9B,C,C′). Additionally, these fibers were found to innervate masseteric fascia, which surrounds the whole MM (Fig. 9D).

Analysis of multiple slides showed that MM tendons almost exclusively contained CGRP+/NFH− (M1 and M2 groups; Fig. 10A,A′, cyan arrows) and CGRP+/NFH− nerves (M3 and M4 groups; Fig. 10A,A′, white arrows). Masseteric muscle fibers were also nearly exclusively innervated by CGRP+/NFH+ (cyan arrows) or CGRP+/NFH− (white arrow) MM TG neurons (Fig. 10B,B′). This innervation pattern was detected around masseteric fascia as well (Fig. 9D). Thus, only 1–3 peptidergic CGRP+/NFH− nerves (S3, S4, and maybe S5 groups) were observed in muscle fibers, adjacent tendons of MM and masseteric fascia investigated by 20× objective across 16–24 MM sections generated from three to four mice.

To further investigate this unexpected finding, we examined MM distribution of nerves with peptidergic CGRP+/NFH− markers such as trpV1+; Nav.1.8+, and pgp9.5+; and NFH+ markers such as trkC+ and PV+. Moreover, to evaluate whether these findings are translated across species, MM fiber types were labeled in the marmoset tissue. Pgp9.5 is marker for all types of sensory nerves, and could highlight location of non-peptidergic unmyelinated nerve fibers (Tokushige et al., 2006). Analysis 8–16 slides with MM sections from male mice showed thatpgp9.5+ nerves innervating muscle fibers (Fig. 11A,A′), tendon of MM (Fig. 11B), and masseteric fascia (Fig. 11C) are always co-labeled by NFH (cyan arrows on all panels). Muscle fibers of male marmosets were predominantly innervated by pgp9.5+/NFH+ nerves (Fig. 11D,D′, cyan arrows). However, pgp9.5+/NFH− nerves (Fig. 11D,D′, white arrows), which likely represent

TG sensory afferent nerve types innervating masseteric muscle fibers and tendons

The masseteric nerve enters the deep portion of MM and is divided into many branches within MM (Fig. 13A,B). These branches eventually supply the sensory afferent

Figure 8. Expression of trkC and TH mechanoreceptor markers in MM TG neurons. A, A′: Expression of trkC (trkC; red) in a TG section containing MM TG neurons (WGA; green). B, B′: Expression of TH (red) in a TG section containing MM TG neurons (WGA; green). MM TG neurons lacking a sensory neuronal marker are shown with cyan arrows on the A, B panels. MM TG neurons containing a sensory neuronal marker are shown with yellow arrows on the A, A′ panels. For all panels, objective is 20×. White horizontal bar shows 20-μm scale for each panel.

C, Percentages of WGA+ TG neurons labeled with a variety of indicated sensory neuronal markers. Cell counting is from four animals, three to five sections each.

For TG neuronal groups, we could estimate that 24% of MM TG neurons belong to groups S1 and S2; 15% to S3 and S2; 5% to S5, 24% to M1; 5% to M2; 12% to M3; and 10% to M4 (Fig. 13C).
C-fibers, were seldomly detected on one of two MM sections observed with 10× objective. These findings imply that male mouse MM fibers, tendon and fascia are almost solely innervated by A-fiber myelinated fibers, while C-fiber unmyelinated fibers are rarely encountered in MM fibers of marmosets. TRPV1 marks S3 and S4 groups of MM TG neurons containing peptidergic unmyelinated fibers. TRPV1+ nerves were not detected within masseteric muscle fibers (Fig. 11A,A′), tendon or fascia of MM (data not shown). However, some nerve branches of masseteric nerve located along muscle fibers contain TRPV1+ sensory fibers, which was also labeled by pgp9.5+ (Fig. 11B,B′, cyan arrows). Nav1.8+ fibers were highlighted by isolation of MM from Nav1.8-cre/Ai32-YFP mice. Electrophysiology recording showed that strong Nav1.8-YFP signal is detectable only in small neurons (S1–S4 groups), while M1 has weak Nav1.8 presence and S5 and M2–M4 have none. Hence, Nav1.8+ sensory fibers were not detected within muscle fibers and tendon. However, certain masseteric nerve branches contained Nav1.8+ fibers (Fig. 11C,C′, red arrow). Interestingly, we have detected multiple nerve branches adjusted or inside MM expressing both pgp9.5 and NFH, but not showing Nav1.8 (Fig. 11C,C′, yellow arrows). Expression patterns of CGRP+, pgp9.5+, Nav1.8+, and NFH+ nerves within MM indicate that muscle fibers, tendon and fascia are innervated by M1–M4 and maybe S5 TG MM neurons. In concordance with this, trkC, which is a marker for S5 and M2–M4 MM TG neurons (Fig. 13C), was detected in many NFH+ fibers (Fig. 11D,D′, cyan arrows). However, unlike trkC, anti-PV antibodies produced only weak labeling in masseteric nerve branches (data not shown). Overall, our data suggest that muscle fibers, tendon, and fascia of MM in mice and marmosets are almost exclusively innervated by A-fiber myelinated fibers, while C-fiber unmyelinated fibers are rarely encountered in MM fibers of marmosets.

Figure 9. Location of CGRP+ and NFH+ TG sensory neuronal afferent fibers in MM. A, A′, Expression of CGRP (green) and NFH (blue) fibers in a main trunk of the masseteric nerve located near MM. Objective is 40×. CGRP+/NFH+ nerves are marked with yellow, CGRP+/NFH+ with cyan, and CGRP+/NFH+ with white arrows. Red arrow shows MM fibers. B, Expression of CGRP (red) and NFH (green) nerves in muscle fibers and MM tendon. Objective is 10×. Red arrow shows muscle fibers and blue points on tendons. Nerve bundles are marked with yellow arrows and individual nerves with cyan arrows. C, Expression of CGRP (red) and NFH (green) nerves in MM. Objective is 2×. A red arrow shows muscle fibers. C′, High magnification (objective is 20×) of the image within a white rectangle on the panel C. On the panels C, C′, nerve bundle is marked with a yellow arrow and individual nerves with cyan arrows. D, Expression of Nav1.8-Ai32-YFP (green), CGRP (red), and NFH (blue) nerves in MM fascia area, which is indicated by white double arrowed line. Objective is 10×. White horizontal bar shows 20- or 100-μm scale for each panel.
Mechanisms underlying masticatory myofascial pain are largely unknown. Masticatory myofascial pain involves MM in several conditions, including TMD and after some restorative surgeries in the head and neck. In this respect, understanding jaw muscle and particularly MM innervation is critical for dissecting cell-specific mechanisms controlling development of chronic masticatory myofascial pain. This understanding will eventually allow for the effective management of TMD pain (Dinsdale et al., 2021).

In addition, masseteric nerve is affected during certain restorative surgeries and during transplantations/transfers, which are the most often performed cranial nerve manipulation technique used to treat patients with facial paralysis (Corcoran and Goldman, 2021). In this study, using back-tracking from MM with WGA, reporter mice driven by promoters of well-characterized sensory neuronal markers, patch clamp electrophysiology and IHC, we identified and characterized types of TG sensory neurons innervating MM. This approach was successfully employed to identify and characterize neuronal groups in L3–L5 DRG and TG (Petruska et al., 2000; Xu et al., 2010; Patil et al., 2018).

We clustered most recorded MM TG neurons into nine clusters (Fig. 13; Table 2). One to two percent of recorded neurons did not fit these nine clusters. For example, MM TG neurons did not respond to ATP (Table 1), but from >350 analyzed neurons, seven responded to ATP.

There is not clear consensus on how the sensory neurons between DRG and TG differ. There is also not enough data on the biochemical differences between sensory neurons innervating the MM. Nevertheless, there is agreement that somatosensory neurons in different sensory ganglia have distinct anatomic, functional, physiological, and pathophysiological characteristics (Li, 2007; Belmonte and Viana, 2008; Gambeta et al., 2020). Thus, DRG are located in the intervertebral foramina at spinal levels that mainly innervate tissues within the trunk, legs and feet, while TG rest in Meckel’s cave and innervate the head and neck area (Vermeiren et al., 2020). Another anatomic distinction is proprioceptors with Aα-fibers are located in DRG but are outside TG in the mesencephalic TG nucleus (Jerge, 1963; Dubner, 1986). Since physiological function of tissues in the trunk, legs and feet are quite distinct from tissues in head and neck area, it is presumed that DRG and TG sensory neurons have distinct, specialized, and unique sets of proteins/mRNA. With the advent of RNA-seq, data from sorted sensory neurons and single-cells described some of these differences in recent years (Usoskin et al., 2015; Lopes et al., 2017; Nguyen et al., 2019; Zheng et al., 2019; Mecklenburg et al., 2020; Renthal et al., 2020). However, these sequencing data indicate that the differences between sensory neurons in DRG and TG are not dramatic and lack clear functional distinctions (Lopes et al., 2017; Mecklenburg et al., 2020; Sharma et al., 2020). These differences between DRG and TG neurons could be because of analysis of sensory neuronal profiles in whole ganglia, but not in specific subsets of sensory neurons innervating defined tissues (such as muscle, skin, tooth, dura, etc.). Considering that neuronal subsets may have distinct origins and may therefore be guided to different targets, innervating different tissues (Hockley et al., 2019), we focused our study on a specific subset of TG neurons innervating MM.

Ideally, our data need to be compared with DRG neurons innervating hind/fore limb muscle. However, this information is not available. Single-cell data generated by several independent studies imply that difference between neuronal groups innervating limb skin and muscle
maybe not substantial (Usoskin et al., 2015; Sharma et al., 2020). Thus, our data indicate that 20–25% of L3–L5 DRG neurons were labeled by WGA injected into limb muscles (Mecklenburg et al., 2020). Single-cell sequencing of thousands of sensory neurons clustered major phenotypic differences of DRG groups innervating the skin and muscle, for example, proprioceptors were clustered as a separate group (Usoskin et al., 2015; Sharma et al., 2020). Taken all these points into consideration, we compared our results with published data from whole L3–L5 DRG neurons (Lou et al., 2015), MM TG IB-4+ neurons were not MrgrpD+. Functional studies on DRG neurons distinguished two types of skin innervating IB-4+/MrgrpD+ neurons: C-polymodal nociceptors (PMN) and C-mechano-nociceptors (C-MN; Liu et al., 2012; Lou et al., 2015). For MM TG neurons, it is not clear what functions could be assigned to S1 and TRPA1 containing S2 groups. S3 and S4 groups containing CGRP, TRPV1 and Nav1.8 are likely C-PMN-like with similarities to DRG neurons (Usoskin et al., 2015; Patil et al., 2018; Sharma et al., 2020). The last small-sized MM TG neuronal group characterized was S5. It had unique electrophysiological properties (Table 1) and we observed expression of CGRP and trkC, but not Nav1.8 in S5 group MM TG neurons. There is no analog for S5 neurons in L3–L5 DRG, and it is unknown whether S5 express NFH and belong to nociceptors or LTMR. Unlike DRG neurons innervating skin, MrgrpA3+ (C-PMN; Han et al., 2013), CGRP–/TRPV1+ (also known as somatostatin+; Usoskin et al., 2015), and TH+ (also known as C-LTMR; Seal et al., 2009) groups were not found among MM TG neurons.

As it was previously reported (Sato et al., 2018), medium-sized neurons are most abundant among MM TG neurons (Fig. 13C). Moreover, in L3–L5 DRG innervating either skin or muscle, all CGRP+ neurons were found inside of the Nav1.8+ subset (Patil et al., 2018), while S5, M1, and M2 MM TG neuronal groups expressed CGRP but little-to-no Nav1.8 (Fig. 13C). M1 and M2 medium-sized MM TG peptidergic neurons produced large inward currents on application of 5HT. This property distinguished M1 and M2 from other MM TG neurons. M1 neurons express high levels of CGRP and trkC, as well as MrgrpA3 and TH, but not MrgrpD+. Functional studies on DRG neurons distinguished two types of skin innervating IB-4+/MrgrpD+ neurons: C-polymodal nociceptors (PMN) and C-mechano-nociceptors (C-MN; Liu et al., 2012; Lou et al., 2015). For MM TG neurons, it is not clear what functions could be assigned to S1 and TRPA1 containing S2 groups. S3 and S4 groups containing CGRP, TRPV1 and Nav1.8 are likely C-PMN-like with similarities to DRG neurons (Usoskin et al., 2015; Patil et al., 2018; Sharma et al., 2020). The last small-sized MM TG neuronal group characterized was S5. It had unique electrophysiological properties (Table 1) and we observed expression of CGRP and trkC, but not Nav1.8 in S5 group MM TG neurons. There is no analog for S5 neurons in L3–L5 DRG; and it is unknown whether S5 express NFH and belong to nociceptors or LTMR. Unlike DRG neurons innervating skin, MrgrpA3+ (C-PMN; Han et al., 2013), CGRP–/TRPV1+ (also known as somatostatin+; Usoskin et al., 2015), and TH+ (also known as C-LTMR; Seal et al., 2009) groups were not found among MM TG neurons.

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contains CGRP, a no-to-low level of Nav1.8, and no trkC. Considering previous research on DRG (Usoskin et al., 2015; Patil et al., 2018), M1 could be a peptidergic myelinated A-fiber nociceptor (Aδ-HTMR). However, it is unclear whether previously described A-fiber nociceptors are distinct in limb skin versus muscle (Usoskin et al., 2015; Sharma et al., 2020). L3–L5 DRG contains two types of Aδ-HTMR, which express CGRP, Nav1.8, but not trkC, can be distinguished by the presence of NPY2R marker (Patil et al., 2018; Sharma et al., 2020). In contrast, TG MM M2 neurons have CGRP, trkC, and a no-to-low level of Nav1.8. Additionally, NPY2R-ttdTomato expression have not been detected in TG neurons (Wu et al., 2018). Such expression patterns in M2 MM TG neurons complicates their functional assignment. However, the presence of two A-fiber HTMR groups innervating MM has been reported (Wong et al., 2014); therefore, M2 could be a separate group of A-fiber HTMR, which are sometimes designated as Aβ-HTMR (Djouhri et al., 1998). MM is innervated by two TG neuronal groups, M3 and M4, that are not revealed in CGRPcre-ER1 and Nav1.8cre1 reporter mice. However, M3 and M4 have been recorded among WGA1/Nav1.8-cre+, WGA1/trkC-cre+, and WGA1/PVcre-+ TG neurons. In L3–L5 DRG innervating limb skin and muscle, functional studies suggested that trkC+/PV+ are either skin Aβ-LTMRs or muscle proprioceptors (Bai et

Figure 12. Expression of trpV1+, Nav1.8+, and trkC+ sensory neuronal afferent fibers in MM. A, A’, Expression of pgp9.5 (green) and TRPV1 (red) nerves in masseteric muscle fibers. Objective is 10×. TRPV1+/pgp9.5+ nerves are marked with white arrows. B, B’, Expression of pgp9.5 (green) and TRPV1 (red) nerves in nerve bundles located within masseteric muscle fibers. Objective is 20×. TRPV1+/pgp9.5+ nerves are marked with cyan arrows. Yellow arrows show nerve bundles. C, C’, Expression of Nav1.8-Ai32 (green), pgp9.5 (red), and NFH (blue) fibers in masseteric nerve branches. Objective is 40×. A red arrow shows a branch with Nav1.8-Ai32+ fibers, and yellow arrows mark Nav1.8-Ai32+ branches. D, D’, Expression of trkC (red) in NFH+ (green) nerves within masseteric muscle fibers. Objective is 20×. TrkC+/NFH+ nerves are marked with cyan arrows. White horizontal bar shows 20-μm scale for each panel.
al., 2015). Since, TG do not have proprioceptors (Jerge, 1963), it could be presumed that M3 and M4 may be MM Aβ-LTMRs neurons. Subgroup of skin Aβ-LTMRs, called Aβ-Field, express calbindin D28 (Calb) as a marker (Arcourt et al., 2017; Sharma et al., 2020). Since Calb is not in MM TG neurons, it could be speculated that M3 and M4 are likely rapidly adapting Aβ-LTMRs (Aβ-RA-LTMRs). Interestingly, electrophysiological properties of M3 is similar to DRG skin innervating Aβ-RA-LTMRs nerves, but M4 strongly resembles DRG muscle innervating proprioceptors (Patil et al., 2018). Thus, their AP, a unique fast AHP, unresponsiveness to algesics and their outward I parameters are almost identical (Fig. 4B,D; Table 1) versus DRG group M7 (Patil et al., 2018). Our data shows that unmyelinated and myelinated fibers travel together in a main trunk of the masseteric nerve (Fig. 9A,A). Similarly, nerve branches of the masseteric nerve contain either both unmyelinated and myelinated fibers or only myelinated fibers (Fig. 12C,C'). It was reported that distribution of different types of fibers occur throughout MM and are independent of fiber type (Herring et al., 1989). First, the data indicate that afferent fibers are concentrated in tendons and at junctions between deep, middle, and superficial parts of MM (Fig. 9B,C,C'). Second, importantly, male mouse muscle fibers, tendons, and masseteric fascia of MM were almost exclusively innervated by myelinated NFH1 afferent nerves (Figs. 9B,C,C'). Likewise, male marmoset MM were predominantly innervated by myelinated NFH+ afferent nerves (Figs. 9B,C,C'.D). Mouse MM TG neurons with myelinated fibers belong to M1–M4 groups, and maybe group S5 as well. IHC could detect unmyelinated MM TG neurons, which could be identified as Nav1.8+/NFH−, TRPV1+, and/or pgp9.5+/NFH− afferent nerves, only within a subset of masseteric nerve branches (Fig. 12A–D'). It is not clear where these unmyelinated fibers terminate, since they were not detected by IHC in MM fibers, MM tendons and masseteric fascia. Accordingly, WGA injection into MM could label small TG neurons (S1–S4) found in the main trunk or sub-branches of the masseteric

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nerve, and/or adjacent to MM subcutaneous tissues and skin via WGA diffusion (Fig. 13A, B).

In conclusion, our data show that MM is almost exclusively innervated by myelinated nerves that can be divided into S5, M1–M4 neuronal groups (Fig. 13A–C). WGA injections into MM also revealed small sized TG neurons (S1–S4), nerve fibers from which are within the masseteric nerve or terminate in adjusted to MM structures (Fig. 13B, C). Properties of these MM TG neuronal groups have similarities and many distinctions from the well-described and characterized L3–L5 DRG neurons innervating limb skin and muscles (Petruska et al., 2000; Usoskin et al., 2015; Patil et al., 2018; Sharma et al., 2020). These differences could imply distinct functional consequences during naive and pathologic conditions affecting MM. It also suggests that TG MM neurons may contain a discrete subset of receptors and channels, which could uniquely regulate sensitization of MM TG neurons during masticatory myofascial pain. Finally, our novel report on the innervation of MM in a non-human primate species, the common marmoset, is important for many reasons. Because of the close relatedness of nonhuman primates and humans, delineation of the MM in the marmoset is important to support its use as a potential animal model MM-related pathology. Further, identification of similarities between marmoset and mouse MM further supports the use of both species as preclinical models for development of intervention approaches in this area.

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