Proprioception from masticatory apparatus and periodontal ligaments comes through the trigeminal mesencephalic nucleus (Vmes). We evaluated the effects of tooth loss on neurodegeneration of the Vmes and trigeminal motor nucleus (Vmo). Bilateral maxillary molars of 2-month-old C57BL/6J mice were extracted under anesthesia. Neural projections of the Vmes to the periodontium were confirmed by injecting Fluoro-Gold (FG) retrogradely into the extraction sockets, and for the anterograde labeling adeno-associated virus encoding green fluorescent protein (AAV-GFP) was applied. For immunohistochemistry, Piezo2, ATF3, Caspase 3, ChAT and TDP-43 antibodies were used. At 1 month after tooth extraction, the number of Piezo2-immunoreactive (IR) Vmes neurons were decreased significantly. ATF3-IR neurons were detected on day 5 after tooth extraction. Dead cleaved caspase-3-IR neurons were found among Vmes neurons on days 7 and 12. In the Vmo, neuronal cytoplasmic inclusions (NCIs) formation type of TDP-43 increased at 1 and 2 months after extraction. These indicate the existence of neural projections from the Vmes to the periodontium in mice and that tooth loss induces the death of Vmes neurons followed by TDP-43 pathology in the Vmo. Therefore, tooth loss induces Vmes neuronal cell death, causing Vmo neurodegeneration and presumably affecting masticatory function.

Key words: trigeminal mesencephalic nucleus, trigeminal motor nucleus, degeneration, TDP-43, tooth extraction

I. Introduction

The mesencephalic trigeminal nucleus (Vmes) neuron is unique in the trigeminal nervous system, and the cell bodies of primary afferent neurons are in the brainstem, not in the trigeminal ganglion [16]. Vmes neurons innervate muscle spindles of the masticatory muscle [1], Ruffini endings of the periodontal ligament (PDL) [7], and the dental pulp [2]. These projections implicate the Vmes in the control of mastication [3, 31, 32] and periodontal proprioception [19]. Approximately 20% of the Vmes neurons in cats [12, 27] and approximately 10–15% in monkeys [14] have afferent connections to the periodontal mechanoreceptors; the remainder project to the jaw-closing muscles.

The functions of the teeth, including mastication and pronunciation, are crucial for life. The neurodegeneration caused by tooth loss is not well-understood. Loss of teeth causes damage to the sensory receptors of the pulp and the PDL [23]. Additionally, degeneration and death of Vmes neurons occurred after peripheral axotomy in rats [30] and following tooth extraction in cats [20]. Also, some Vmes neurons died after tooth extraction in mice with induced Alzheimer’s disease (AD) [11]. However, whether death
of Vmes neurons occurs after tooth extraction in wild-type mice is unknown. Since the Vmes directly controls the function of trigeminal motor neurons, neurodegeneration of Vmes neurons caused by tooth extraction may affect the trigeminal motor nucleus (Vmo). To clarify the neurodegenerative effects on Vmo neurons we focused on the transactivation responsive region-DNA-binding protein of 43 kDa (TDP-43). TDP-43 is present in the nucleus and binds to the TG (U/G) repeat sequence of DNA and RNA under normal condition [4]. In pathological conditions such as amyotrophic lateral sclerosis (ALS) and frontotemporal dementia, TDP-43 reportedly translocates from the nucleus into the cytoplasm of motor neurons or forms neuronal cytoplasmic inclusions (NCIs) [4, 26]. In this study, we investigated whether TDP-43 neuropathology in Vmo neurons is caused by neurodegeneration of the Vmes after tooth extraction.

Here, we investigated the neuronal connection between Vmes neurons and Ruffini endings in the PDL in mice and evaluated the neurodegeneration and death of Vmes neurons following extraction of maxillary molars in C57BL/6J mice. We also examined TDP-43 pathology in Vmo neurons after tooth extraction. The findings indicate that tooth loss induces Vmes neuronal cell death, causing Vmo neurodegeneration and presumably affecting masticatory function.

II. Materials and Methods

Animals

Forty-seven 2-month-old male C57BL/6J mice (25–30 g) were used in this study. The mice were housed in cages at a constant temperature of 25°C under a 12-hr light-dark cycle (light from 6.00 to 18.00). This study was conducted in accordance with the appropriate animal protection guidelines and was approved by the Committee for Animal Experiments of Kagoshima University (approval number, D 20007, D200016). All procedures are in accordance with the ARRIVE guidelines and are carried out in accordance with the EC Directive 86/609/EEC for animal experiments. For tooth extraction, mice were deeply anesthetized with a combination anesthetic of medetomidine (0.3 mg/kg; Kobayashi Kako, Fukui, Japan), midazolam (4.0 mg/kg; Astellas Pharma, Tokyo, Japan), and butorphanol (2.5 mg/kg; Meiji Seika Pharma Co., Ltd., Tokyo, Japan) [15]. The mice were euthanized by deep anesthesia with sodium pentobarbital (75 mg/kg body weight) and transcardially perfused with 50 mL of 10 mM phosphate-buffered saline (PBS; pH 7.3), followed by 100 mL of 4% paraformaldehyde, 75% saturated picric acid, and 0.1 M NaHPO₄ (adjusted to pH 7.0 with NaOH). The brains were next removed from the mice and postfixed with the same fixative for 4 hr at 24°C. After cryoprotection in 30% sucrose in PBS, the brainstems were cut on a cryostat, divided into 13 blocks containing six adjacent series of 50 μm sections from the rostral side to the caudal side of the Vmes, and stored at 4°C in PBS.

Retrograde and anterograde tracers

After tooth extraction, a 4% solution of FG (2-hydroxy-4,4′-diamidinostilbene; Fluorochrome Inc., Denver, CO) in 0.9% saline was injected into the extracted tooth socket for retrograde tracing. For anterograde tracing, AAV2/1-CAG-Flex-eGFP (AV-1-ALL854) and AAV2/1-hSyn-Cre (AV-1-PV2676) were purchased from the Penn Vector Core, University of Pennsylvania. The two virus solutions were mixed and diluted to a final titer of 1 × 10¹¹ genome copies/mL (GC/mL) for injection.

Surgery

The animals were deeply anesthetized with medetomidine (0.3 mg/kg), butorphanol (5.0 mg/kg), midazolam (4.0 mg/kg). The anesthetized animals were mounted onto a stereotaxic apparatus (SR-5R; Narishige, Tokyo, Japan). For anterograde labeling of Vmes neurons, a mixture of AAV2/1-CAG-Flex-eGFP and AAV2/1-hSyn-Cre in 0.2 μL of 10 mM PBS (pH 7.4) was pressure injected into the Vmes (5.4 mm posterior to the bregma, 1.0 mm lateral to the midline, and 2.6 mm deep from the brain surface) through a glass micropipette attached to a Pneumatic Pico Pump PV830 (World Precision Instruments, Sarasota, FL). Virus-injected mice survived for 2 weeks after the injection.

The bi-maxillary first to third molars in anesthetized mice were extracted using a dental probe. After the extrac-
tion, the mice were provided nutrient-rich food. Measurements of body weight were continued until 1 month after tooth extraction, when the mice began to appear normal.

**Immunostaining**

The list of antibodies used in this experiment is shown in Table 1. Brain sections were blocked with 3% donkey serum and 1% bovine serum albumin in 50 mM Tris-buffered saline containing 0.1% TritonX-100. The primary antibodies were directed against ATF3, Piezo2, active cleaved caspase-3, and TDP-43.

For immunofluorescence, the sections were incubated overnight at 4°C with primary antibodies in PBS containing 0.3% Triton X-100, 0.12% lambda-carrageenan, and 1% donkey serum (PBS-XCD) and subsequently, for 4 hr with 1 mg/mL Alexa Fluor 555-, or 488-conjugated goat anti-mouse, anti-rabbit, and anti-goat antibody in PBS-XCD. All incubations were performed at 24°C and were followed by rinsing with PBS containing 0.3% Triton X-100. The sections were observed under a confocal laser scanning microscope (LSM700; Zeiss, Oberkochen, Germany) or a Nikon Eclipse E800M microscope (Nikon, Tokyo, Japan) equipped with a COMOS MP-102300A digital camera (Bio Tools Inc., Gunma, Japan) using ToupView software (ToupTek Photonics, Hangzhou, China).

For immunoperoxidase staining, the sections were incubated with an anti-cleaved caspase-3 antibody overnight, biotinylated goat anti-rabbit immunoglobulin G in PBS-XCD for 3 hr, and avidin-biotin-peroxidase complex (1:100, elite variety; Vector Labs) in PBS-XCD for 1 hr. Finally, the sections were incubated in 0.02% diaminobenzidine tetrahydrochloride and 0.005% H₂O₂ in 0.05 M Tris-HCl buffer (pH 7.6) for 10 min. The sections were mounted on glass slides and observed under a BX51 microscope (Olympus, Tokyo, Japan) equipped with a DP25 camera and cellSens software (Olympus).

**Statistical analysis**

Numerical values and error bars represent the means ± standard deviations. Mean values were compared using the unpaired Student’s *t*-test or two-way analysis of variance followed by the Tukey post hoc test with R software (R Core Development Team, 2020).

**III. Results**

**Characteristics of Vmes neurons projecting to the periodontal ligament**

We investigated whether Vmes neurons project to the PDL of the upper molars by means of retrograde Fluoro-Gold (FG) and anterograde adeno-associated virus encoding green fluorescent protein (AAV-GFP) injection. Immediately following extraction of the upper molars and socket hemostasis, FG was dripped into the sockets. The mice were anesthetized, their heads were fixed to the device, and AAV-GFP was injected anterogradely at the location of the Vmes (Fig. 1A). FG-labelled Vmes neurons could be recognized at 15 days after FG injection (Fig. 1B). At 10 days after its injection into the Vmes, the localization of AAV-GFP was examined. AAV-GFP labeling was confirmed at a position neuroanatomically considered to be the Vmes in a brainstem tissue section (Fig. 1C). Moreover, AAV-GFP labeling was detected in the axon, and a structure thought to be a Ruffini ending was evident in the PDL.
of a mouse with confirmed AAV-GFP labeling of the Vmes (Fig. 1D, E).

There are morphological differences between Vmes neurons projecting to the PDL and those projecting to the muscle spindle of the masticatory muscle in cats [31, 32]. We injected FG into the extraction socket and masseter muscle and assessed the diameter of the central cross-section of the Vmes neurons (Fig. 2A, B). We also measured the diameter of the central cross-section using confocal microscopy. Vmes neurons projecting to the PDL were significantly smaller than those projecting to the muscle spindle of the masticatory muscles (Fig. 2C).

**Loss of Vmes neurons after tooth extraction**

At 1 month after tooth extraction, we enumerated Piezo2-immunoreactive (IR) Vmes neurons in a section from each block (Fig. 3A). Control specimens were obtained from age-matched (3-month-old) mice that did not undergo tooth extraction. The Vmes was divided into 13 equal-size blocks along the rostro-caudal axis, and the number of Piezo2-IR Vmes neurons in one section of each block was determined (Fig. 3A–C). There were 24% fewer Piezo2-IR Vmes neurons in blocks 9–11 (Fig. 3D), and the number of Vmes neurons differed significantly between the control and extraction groups (unpaired t-test, p = 0.005; Fig. 3E).

**Neurodegeneration of Vmes neurons after tooth extraction**

Next, we investigated the post-extraction timings of Vmes neuronal damage and death using ATF3 for one section from block 11 and cleaved caspase-3 for all blocks. ATF3-IR nuclei became evident in Vmes neurons at 5 days after tooth extraction (Fig. 4A), and the number of ATF3-IR Vmes neurons decreased at 10 and 15 days after tooth extraction (Fig. 4B, C). Also, the number of
ATF3-IR Vmes neurons was significantly reduced in the 5-day group compared to the 10-day group (Tukey post hoc test, p = 0.008, Fig. 4D). To confirm that neuronal death occurred at 5 to 10 days after tooth extraction, the number of cleaved caspase-3-IR Vmes neurons was examined. Cleaved caspase-3-IR Vmes neurons were detected in rostral and caudal sections (Fig. 4E, F). The number of cleaved caspase-3-IR Vmes neurons was greater on day 7 after tooth extraction than on day 12, and a large number of cleaved caspase-3-IR neurons were observed in blocks 7–11 (Fig. 4G).

Effects on TDP-43 appearances after tooth extraction

Since Vmes neurons are involved in controlling the trigeminal motor nucleus, we examined the effects of tooth extraction on Vmo neurons using TDP-43 appearances as an indicator. Labeling with AAV-GFP confirmed that the axons of Vmes neurons reached the motor nucleus (Fig. 5A). Additional staining for choline acetyltransferase (ChAT), a marker of motor neurons, confirmed that the axons of Vmes neurons were in contact with Vmo neurons (Fig. 5B). TDP-43 pathology in the Vmo neurons after tooth extraction was examined by double staining using anti-ChAT and anti-TDP-43 antibodies (Fig. 5C–K). We identified three types of TDP-43-IR, intranuclear, cytoplasmic, and NCIs formation types (Fig. 5F–K). In the normal type, which TDP-43-IR stain was distributed throughout the nucleus was defined as intranuclear type. As for pathological types, two types of abnormal TDP-43 localization were observed: one in which TDP-43-IR is present only in the cytoplasm and not in the nucleus, and one in which TDP-43-IR NCIs was formed. (Fig. 5H, K). We evaluated the frequency of each type at 12 days, 1 and 2 months after tooth extraction (Fig. 6A). In the 2-month control, approximately 90% of the Vmo neurons were of the normal type, but some were of NCIs formation (6.8%) and cytoplasmic (3.5%) types. At 12 days after tooth extraction, once NCIs formation type disappeared and cytoplasmic type increased comparer with control (Fig. 6C). At 1 month after tooth extraction, the ratio of the cytoplasmic type and NCIs formation types increased (Fig. 6D). At 2 months after tooth extraction, the ratio of the NCIs formation type also increased to 20% but that of cytoplasmic type decreased to 5.7% (Fig. 6E). Time dependent changes of the percentage of cytoplasmic type and ICN type after tooth extraction were summarized. Cytoplasmic type showed the initial increase until 1 month then decrease at 2 months after tooth extraction (Fig. 6F). ICN type was found the temporal disappearance at 12 days after tooth extraction and increase thereafter (Fig. 6G).

IV. Discussion

Neuronal projections of Vmes neurons were associated with Ruffini ending-like structures in the PDL of C57BL/6J mice, and death of Vmes neurons was induced after extraction of maxillary molars. Neurodegeneration of Vmes neurons occurred within 5 days of tooth extraction and was followed by their death. Also, the cell death effects as observed in the Vmes extended to the Vmo, with pathological changes in TDP-43 distribution observed in the Vmo at 1 month after tooth extraction. Therefore, tooth extraction causes pathological changes in the Vmo.

The Vmes regulates jaw movements and is the only
sensory nerve in the trigeminal nervous system that has primary neurons in the brain stem. In rats [6, 8, 17], cats [7] and monkeys [14], Vmes neurons have periodontal mechanoreceptor afferents. In this study, we examined Vmes-PDL mechanoreceptors in C57BL/6J mice using anterograde and retrograde labeling. Although Ruffini nerve endings are present in the mouse PDL [24, 28], whether they contact the Vmes was unclear. From anterograde tracing using AAV-GFP, we found that Vmes neurons project to Ruffini nerve ending-like structures in the PDL of mice. From retrograde tracing, because we extracted the molars without leaving the roots in situ, the Vmes could be accurately marked by dropping FG into the extraction socket.

Because the Vmes innervates at least two major regions—namely, the muscle spindle of the masticatory muscles and the PDL, we examined the size of the Vmes soma. We found that the Vmes is composed of two subpopulations of neurons—smaller Vmes neurons (20.02 ± 3.86 μm) innervate the PDL mechanoreceptor and larger Vmes neurons (23.51 ± 3.48 μm) innervate the masticatory muscle spindle. The Vmes in cats and rats can be divided into two subpopulations according to size. In rats, the larger Vmes has a diameter of 30–55 μm and the smaller, a diameter of ≤ 25 μm [21, 22]. This difference in Vmes size is reportedly due to differences between pseudo-unipolar and multipolar neurons [22]. However, in this study, the size of the Vmes differed depending on whether their neurons innervated muscle spindles or the periodontal membrane.

Death of Vmes neurons after tooth extraction has been
reported in 3xTg-AD model mice [11]. Because intense deposition of amyloid β occurs in the Vmes of 3xTg-AD mice, it is possible that amyloid β affects neuronal death in the Vmes. Using wild-type C57BL/6J mice, we observed similar neuronal death in the Vmes after extraction of the upper molars. Typically, even if a portion of the axons of peripheral neurons is damaged, Wallerian degeneration occurs, but neuronal death is rare. Although we do not know why tooth extraction causes neuronal cell death in the Vmes, the Vmes neuron is the only neuron with a soma localized in the brainstem; thus, damage to the axon terminal may also cause damage in the cell soma in the brainstem. Therefore, tooth extraction could cause degeneration of the entire nerve.

The number of ATF3-IR neurons in the Vmes region increased in a time-dependent manner, and cleaved caspase-3 positivity indicated neuronal death in the Vmes. In rats, ATF3-IR staining was observed in the neurons of the trigeminal ganglion at 3 days after extraction of molars [13]. In this study, we found ATF3-IR Vmes neurons at 2 weeks after the extraction of maxillary molars from AD model mice [11]. ATF3-IR neurons appeared at 5 days after tooth extraction, and their number decreased at 10 and 15 days after tooth extraction. The decreased number of ATF3-IR cells could be due to cell death or recovery from damage. Therefore, we confirmed cell death using an antibody

Fig. 6. Changes in TDP-43 expression type in Vmo after tooth extraction. (A) Illustrations of TDP-43 expression type. TDP-43-IRs in Vmo are divided into three types: Intranuclear type (IN), Cytoplasmic type (Cyto), and NCIs formation type (NCI). Percentage of IN, Cyto, and NCI type in control (B), 12 days- (C), 1 month- (D), and 2 months after tooth extraction (E). (F, G) Time-dependent changes of the percentage of cytoplasmic type and ICN type after tooth extraction. *p < 0.05, **p < 0.01, two-way ANOVA followed by Tukey’s post hoc analysis. Data are presented as the means ± SDs, n = 10.
against cleaved caspase-3. The changes in position and number of cleaved caspase-3-IR neurons were similar to those observed for the cells at each site (Fig. 3D). That is, the decrease in cell number was greater on the caudal side, where there were a large number of cleaved caspase-3-IR neurons. However, some cleaved caspase-3-IR neurons survived. Also, caspase-3 is reportedly activated during apoptosis, indicating a functional shift in Vmes cells rather than their impending death [10, 18]. Vmes neurons relay proprioceptive sensory information from mechanoreceptors in the PDLs and spindles of the masticatory muscle, and this information is relayed primarily to the rostral end of the Vmes, nucleus supratrigeminalis, and trigeminal motor nucleus as well as for parvocellular reticular formation. These projections control the mechanics of mastication [5, 9].

We confirmed the existence of projections from Vmes to Vmo neurons using AAV-GFP. This finding suggests that the death of Vmes neurons causes neurodegeneration in the Vmo. TDP-43 is the major component of the neuronal inclusions characteristic of ALS and frontotemporal lobar dementia (FTD) with ubiquitin inclusions. Although TDP-43 was initially thought to be specific to ALS and frontotemporal lobar degeneration, TDP-43 pathology has been detected in a number of other neurodegenerative diseases. These include diseases associated with tau pathology, including Guam–Parkinson dementia complex and AD [34]. TDP-43-IR NCIs with a rounded, specular, or skein-type appearance were observed in motor neurons of the trigeminal or facial cranial nerve of one patient with FTD and in the spinal cord of three patients with FTD and motor neuron disease. The majority of studies on TDP-43 have focused on its abnormal distribution in disease states—TDP-43-IR are present in up to 97% of patients with ALS [29]. TDP-43 pathology is infrequent (≤ 3%) in neuropologically normal elderly individuals [25]. Interestingly, 10.2% of the motor neurons in C57BL/6J mice exhibited TDP-43 pathology, the frequency of which increased to 18.4% and 14.7% at 1 and 2 months, respectively, after tooth extraction. The more severe TDP-43 pathology in mouse motor neurons compared to human elderly motor neurons under normal conditions might be due to species differences or to effects on responsiveness to neurodegeneration. In C57BL/6J mice, NCIs formation type of TDP-43 was not found in the motor nerve at 12 days after tooth extraction but was found to increase in Vmo neurons at 1–2 months after extraction. The difference between the pathological roles of cytoplasmic type and ICN type is not well understood, but our results show that Vmo neuron after tooth extraction tends to increase cytoplasmic type first and then increase ICN type. However, the pathological changes in TDP-43 after tooth extraction were limited, compared to the pathological changes in TDP-43 in more than 90% of motor neurons in patients with ALS. These indicate that the effects of tooth extraction on Vmo neurons were due to not only muscle atrophy caused by loss of occlusion but also the effect of decreased input from the Vmes. Reversible induction of TDP-43 granules has been reported in traumatic brain injury [33] and our results suggest that TDP-43 pathology is a reversible change. Further studies are needed on the effects of TDP-43 pathology on masticatory movements.

We used C57BL/6J mice to investigate projections from the Vmes to the PDL and evaluated the effects of extraction of maxillary bilateral molars on the Vmes and Vmo. Projections from the Vmes to the PDL were confirmed using FG-retrograde and AVV-GFP-antegrade tracing. Extraction of the maxillary bilateral molars reduced the number of Vmes neurons by 24%, but ATF3-IR neurons were detected in the Vmes within 5 days, followed by cleaved caspase-3-IR staining and cell death. After neuronal death in the Vmes, NCIs formation type of TDP-43 in the Vmo increased. Therefore, after tooth extraction, neurodegeneration spreads from Vmes to Vmo neurons and may affect the function of the masticatory muscles.

V. Conflicts of Interest

The authors declare no conflicts of interest.

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