Prostanoid-dependent spontaneous pain and PAR₂-dependent mechanical allodynia following oral mucosal trauma: Involvement of TRPV1, TRPA1, and TRPV4

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Abstract
During dental treatments, intraoral appliances frequently induce traumatic ulcers in the oral mucosa. Such mucosal injury-induced mucositis leads to severe pain, resulting in poor quality of life and decreased cooperation in the therapy. To elucidate mucosal pain mechanisms, we developed a new rat model of intraoral wire-induced mucositis and investigated pain mechanisms using our proprietary assay system for conscious rats. A thick metal wire was installed in the rats between the inferior incisors for one day. In the mucosa of the mandibular labial fornix region, which was touched with a free end of the wire, traumatic ulcer and submucosal abscess were induced on day 1. The ulcer was quickly cured until next day and abscess formation was gradually disappeared until five days. Spontaneous nociceptive behavior was induced on day 1 only, and mechanical allodynia persisted over day 3. Antibiotic pretreatment did not affect pain induction. Spontaneous nociceptive behavior was sensitive to indomethacin (cyclooxygenase inhibitor), ONO-8711 (prostanoid receptor EP₁ antagonist), SB-366791, and HC-030031 (TRPV1 and TRPA1 antagonists, respectively). Prostaglandin E₂ and 15-deoxyΔ¹²,₁₄-prostaglandin J₂ were upregulated only on day 1. In contrast, mechanical allodynia was sensitive to FSLRY-NH₂ (protease-activated receptor PAR₂ antagonist) and RN-1734 (TRPV4 antagonist). Neutrophil elastase, which is known as a biased agonist for PAR₂, was upregulated on days 1 to 2. These results suggest that prostanoids and PAR₂ activation elicit TRPV1- and TRPA1-mediated spontaneous pain and TRPV4-mediated mechanical allodynia, respectively, independently of bacterial infection, following oral mucosal trauma. The pathophysiological pain mechanism suggests effective analgesic approaches for dental patients suffering from mucosal trauma-induced pain.

Keywords
Oral traumatic mucositis, oral abscess, TRPV1, TRPA1, TRPV4, PAR₂

Introduction
During dental treatments, various metal and plastic appliances are installed on the tooth or implanted in the alveolar bone. Sometimes these appliances iatrogenically induce traumatic ulcer due to mechanical stimulation and subsequently lead to painful mucositis despite considerable managements by dentists. The incidence of mucositis has been reported to be 92% in denture patients and 63%–95% in general orthodontic patients.¹–⁴ Implantation of miniscrews in alveolar
bone for orthodontic loading has been described to frequently cause traumatic lesions in the area in contact with the miniscrew. Additionally, a clinical review has described abscess formation in orthodontic wire-induced trauma. Even after removing the injuring parts of the appliance, the previously induced pain is maintained for a few days until healing, particularly during meals and conversations. Some clinical studies have reported that mucosal trauma-associated pain leads to poor quality of life and decreased cooperation in therapy. However, no studies have investigated the pathological properties and pain mechanisms responsible for mechanical trauma to the oral mucosa.

Recently, a few studies have demonstrated pain mechanisms in the oral mucosa using preclinical models of rats; the involvement of the heat-sensitive TRPV1 channel and the cold/mechanical-sensitive TRPA1 channel in nociceptive trigeminal neuronal fibers. Incision of the facial skin and the cold/mechanical-sensitive TRPA1 channel in rats; the involvement of the heat-sensitive TRPV1 channel, TRPV4, has been reported to be involved in inflammatory pain. Oral ulcerative mucositis treated with acetic acid following the administration of 5-fluorouracil, a representative chemotherapeutic drug for head and neck cancer, induces severe spontaneous pain and mechanical allodynia due to TRPV1 activation and TRPA1 sensitization, respectively, which is dependent on bacterial infection. Some differences in the pain-producing mechanisms induced by the nociceptive channels between preclinical models are probably dependent on different degrees of tissue damage and levels of infection in the oral mucosal region. In addition to TRPV1 and TRPA1, another mechano-sensitive TRP channel, TRPV4, has been reported to be involved in inflammatory pain. Therefore, these channels may contribute to pain in oral traumatic mucositis.

The present study was designed to reveal cellular and molecular mechanisms associated with traumatic mucositis-induced pain in the oral mucosal region. First, we prepared a preclinical rat model for intraoral appliance-induced mucositis and evaluated the pathophysiological status. Next, we investigated spontaneous nociceptive behavior and mechanical allodynia in the preclinical model using our proprietary methods and evaluated effects of various drugs to reveal involvements of TRPV1, TRPA1, and TRPV4.

Materials and methods

**Experimental animals**

Male Wistar rats (eight to nine weeks of age; Kyudo, Saga, Japan) were used in this study. The rats were housed in pairs in clear cages with wood chips under specific pathogen-free conditions and maintained on a light-dark cycle (L:D, 12:12 h) in a temperature- and humidity-controlled room (21°C–23°C and 40%–60%, respectively) with food pellets and water provided ad libitum. All experiments were conducted in accordance with the National Institutes of Health guidelines (Guide for the Care and Use of Laboratory Animals) and were approved by the Animal Experiment Committee of Kyushu Dental University. All efforts were made to minimize animal suffering. Rats were randomly selected for each experiment. Measurements of all pain tests were performed in a blinded manner to the drug applications.

**Model of wire-induced oral mucositis**

First, in preliminary experiments, we attempted to induce traumatic mucositis by incision or mechanical scratching in the oral mucosa using a needle tip. However, the transient mechanical injuries rapidly healed by the following day, which is inconsistent with clinical pain complaints. Hence, we selected to install a thick wire in the oral cavity for one day to generate a traumatic ulcer. An orthodontic 10-mm-length wire (E00097, TRU-CHROME, 0.018 × 0.025 inch [0.457 × 0.635 mm]; Rocky Mountain Orthodontics, Denver, CO) was manually bent into the shape shown in Figure 1(a) and soldered to a ligature wire (506-02, preformed ligature wire, 0.25 mm in diameter; Tomy International Inc., Tokyo, Japan) at the first 50°-bended angle using a spot welder (Type 660; Rocky Mountain Orthodontics). Under pentobarbital anesthesia (50 mg/kg, intraperitoneal), the inferior incisors were grooved at the top edge of the incisor gingiva (~5 mm from the tip face of the incisors) using a round diamond bur (616001-K1; Morita, Osaka, Japan). The bent wire was installed between the incisors using the ligature wire (Figure 1(b)), and the ligation portion was completely covered with dental adhesive resin cement (Super-Bond, Sun Medical, Shiga, Japan) to avoid mucosal injury at the ligation site. The free end of the thick wire, with a sharp tip generated by a wire niper, touched the labial fornix region of the oral mucosa. As a control, a shorter thick wire (4 mm) was installed similarly without any injurious contact with the oral mucosa. To avoid detachment of the thick wires, the rats were provided with powdered food (not pellet type) during the wire installation. On the following day (day 1), the wire was removed under 2% isoflurane anesthesia. In rare cases, removal of the wire from the incisors revealed a pus discharge and bleeding from the ulcer area. Such additionally injured rats were excluded from this study.

**Evaluation of oral mucositis severity**

To evaluate the severity of the oral mucositis, we used the following visual oral ulcerative mucositis score,
Figure 1. A rat model of wire-induced mucositis (WiM) and the pathological properties. (a) Design of the thick metal wire as an intraoral appliance in the WiM model. A 10-mm-length wire was bent three times at 50°, 30°, and 25°. (b) Procedure for installing the intraoral appliance. The inferior incisors were grooved 5 mm from the edge (horizontal dashed line) using a round diamond bur; and the bent wire was installed using the ligature wire soldered at the first angle, as indicated by a black arrow. The ligation portion is covered with dental resin cement, and the free end of the thick wire touches the labial fornix region of the oral mucosa, as shown in the photograph of the rat mouth on the right side. (c) Visual appearance of the oral mucosa of sham rats and the WiM model. For the sham, a shorter thick wire (4 mm) was similarly installed. (d) The pH of the surface of the oral mucosa in the sham and WiM model (each group, n = 6). (e) Daily changes in body weight following thick wire installation in the sham and WiM model (each group, n = 5). An arrow and horizontal bar indicate a procedure point, installation of the thick wire, and powdered food loading, respectively. (f) Daily changes in oral mucositis scores following the procedure in the sham and WiM model and effect of antibiotic pretreatment (+ antibiotic) on day 1 (each group, n = 6). For antibiotic pretreatment, the drinking water was supplemented with sulfamethoxazole and trimethoprim at 800 and 400 µg/mL, respectively, beginning two days prior to the procedure. * and ** indicate *P < 0.05 and 0.01, respectively, compared with day 1; Dunnett post hoc test following one-way repeated-measures analysis of variance. †*P < 0.05 compared with day 1 without antibiotic pretreatment; Student t test. (g) The oral mucositis score in the WiM model on day 1 following indomethacin (Indo) pretreatment or vehicle (Veh; 0.1 M Tris-buffered saline) (each group, n = 9). †*P < 0.05 compared with Veh; Student t test. (h) Activity of the neutrophil-specific enzyme MPO in the sham and WiM model (each group, n = 4). †*P < 0.05 compared with the sham; Student t test. (i) Representative hematoxylin and eosin-stained microphotographs of the oral mucosa of sham and WiM model rats at days 1 and 3 after the procedure. Scale bar, 500 µm. Compared with the sham (n = 5), the oral mucosal region was largely destroyed (traumatic ulcer, left side) and formed an abscess in the submucosal layer (right side; 0.6 mm for the section shown on the left side) in WiM model rats on day 1 (n = 5). On day 3 (n = 5), the abscess trended to be reduced compared with day 1. However, we failed to detect abscess formation in any tissue sections on days 4 (n = 2) and 5 (n = 2 of 3), except for sections from a WiM model rat on day 5 with a tiny abscess (data not shown). (j) PAS staining of the abscess in the WiM model on day 1. The abscess was negative for PAS (n = 2), indicating that it was not a mucus cyst. The lower microphotograph is an expansion of the center of the abscess in the upper microphotograph. PAS-positive cell wall-like images. Scale bars in the upper and lower microphotographs, 300 and 100 µm, respectively.
which was modified from a previously reported score using an acetic acid-induced oral mucositis model;[10] normal; 0.5, doubtful of the presence/absence of redness; 1, slight but definite redness; 2, severe redness; 3, focal pseudomembrane, without a break in the epithelium; 4, pseudomembrane, with a clear traumatic ulcer; 5, broad pseudomembrane with yellow mucus; and 6, severe swelling of the lower lip with a score level of 5. The evaluation was performed transiently under 2% isoflurane anesthesia.

**Measurements of pH and temperature in the oral mucosa**

Using a portable pH meter (PH-201 Z; Chemical Instruments, Tokyo, Japan) with a small electrode (CMS-30D; Chemical Instruments) and an infrared thermometer (IT-550 S; HORIBA, Kyoto, Japan), the pH and temperature on the surface of the oral mucosa was measured on day 1 under 2% isoflurane anesthesia.

**Histology of the oral mucosa tissue**

Under deep anesthesia with pentobarbital (100 mg/kg, intraperitoneal), the lower lip of the sham and wire-induced oral mucositis (WiM) model rats was removed and fixed in formalin. After paraffin embedding, 10-μm-thick sagittal sections of the oral mucosa were stained with hematoxylin and eosin. Periodic acid-Schiff reagent (PAS) and Alcian blue staining were performed to distinguish abscesses and mucus cysts in the submucosal layer. The sections were also stained with toluidine blue to evaluate mast cell degranulation.

To assess extravasation in the oral mucositis, WiM model rats on day 1 received Evans blue (1% in saline; E-2129; Sigma–Aldrich, St. Louis, MO) via the tail vain. The isolated lip was trimmed to construct a block with a mucosal surface of approximately 3 × 3 mm and a thickness of 1–2 mm. The block was then bisected onto Brain Heart Infusion agar (Nissui, Tokyo, Japan), followed by overnight incubation at 37°C. The colony-forming units (CFUs) of the duplicated bacterial cultures were then manually counted. The number of CFUs is presented as the number per wet tissue weight.

**Myeloperoxidase activity measurements and enzyme-linked immunospecific assays**

Mucosal tissue in the labial fornix region of the inferior incisors, which includes the area with the most severe mucositis, was extracted from rats that were deeply anesthetized with pentobarbital (100 mg/kg, intraperitoneal) in sham and WiM model rats on day 1 or 2. The tissue was stored at −80°C for use within three months for biochemical assays. The preserved oral mucosal tissues were melted and homogenized in PBS with 10 μM indomethacin and a protease inhibitor cocktail. Following centrifugation, the supernatants were collected, and total protein concentrations were measured using a BCA protein assay kit (Thermo Fisher Scientific, Waltham, MA). Myeloperoxidase (MPO) activity was measured using the MPO activity colorimetric assay kit (Bio Vision, Milpitas, CA). The concentrations of cyclooxygenase-2 (COX-2), prostaglandin E2 (PGE2), 15-deoxy-Δ12,14-PGJ2 (15d-PGJ2), interleukin-1β (IL-1β), IL-6, tumor necrosis factor-α (TNF-α), bradykinin, and neutrophil elastase in the supernatants were measured with the following ELISA kits: Rat COX-2 Assay Kit (27187; IBL International, Hamburg, Germany), Prostaglandin E2 ELISA Kit (ab133021; Abcam, Cambridge, MA), 15-deoxy-delta12,14-PGJ2 ELISA Kit (ab133031; Abcam), Rat IL-1 beta, IL-6 and TNFα ELISA Kits (ER2IL1B, ER3IL6, and ER3TNFA, respectively; Thermo Fisher Scientific), Bradykinin ELISA Kit (ab136936; Abcam) and Rat HNE/Neutrophil Elastase ELISA Kit (LS-F20775; LifeSpan BioScience, Inc., Seattle, WA). The assays were performed in duplicate, and the concentrations and enzyme activities were standardized based on the total protein concentrations.

**Quantification of bacterial counts**

On day 1, the mucosal tissues of the sham and the WiM model with and without antibiotic pretreatment were extracted from rats that were deeply anesthetized with pentobarbital (100 mg/kg, intraperitoneal). All instruments and the entire lower lip skin were sterilized with 70% ethanol before each preparation, and the superficial skin was carefully removed with scissors before removing the lower lip. The isolated lip was trimmed to construct a block with a mucosal surface of approximately 3 × 3 mm and a thickness of 1–2 mm. The block was then bisected with a surgical scalpel. The divided mucosal tissue was placed into a pre-weighed 1.5-mL plastic tube filled with 1 mL of sterilized PBS and ultra-sonicated for 30 s (38 kHz, US-2; SOND, Nagano, Japan) to leach out oral bacteria. Each sample (50 μL) was plated in duplicate onto Brain Heart Infusion agar (Nissui, Tokyo, Japan), not exceeding 500 counts/90-mm-diameter dish. Anaerobic incubation was performed in an airtight container with AnaeroPack-Anaero, an O2-absorbing and CO2-generating agent (Mitsubishi Gas Chemical, Saga, Japan), followed by overnight incubation at 37°C. The colony-forming units (CFUs) of the duplicated bacterial culture plates were then manually counted. The number of CFUs is presented as the number per wet tissue weight.

**Drug applications**

In this study, four kinds of drug applications were performed: (1) administration of fluids, (2) intraperitoneal
administration, (3) swab application, and (4) submucosal injection.

As antibiotic pretreatment for the WiM model, sulfamethoxazole (Wako, Osaka Japan) and trimethoprim (MP Biomedicals, Illkirch, France) were dissolved in drinking water at 800 and 400 μg/mL, respectively. The pretreatment was performed beginning at two days prior to the wire installation and continued for five days. The pretreatment revealed a significant efficacy for blocking bacterial loading in oral ulcer models of rats.\[^{10}\]

As an anti-inflammatory pretreatment, the nonselective COX inhibitor indomethacin (Wako) was intraperitoneally administered at 5 mg/5 mL/kg. The drug administration was performed once per day on days 0 and 1 (4–6 hr prior to the pain tests). As a control, the same volume of a vehicle solution (0.1 M Tris-buffered saline) was administered. The pretreatments revealed a significant efficacy for blocking mechanical nociceptive behavior.\[^{22}\]

To examine whether oxidation, ATP and TRP channel activation are involved in the local pain mechanism in the oral mucosa of rats, we used the stable intraoral opening (SIO) method.\[^{22}\] Before the measurements, the rats were acclimated over a period of three days (once per day) to the plastic cage (30 × 30 × 30 cm) used for the observations. To evaluate spontaneous pain, the behavioral observation was performed for 10 min.

**Withdrawal threshold to von Frey stimulation**

Using the stable intraoral opening (SIO) method, the rats were pierced with a magnetized ring in the lower lip to stably expose the oral mucosa in the labial fornix region of the inferior incisors, as described in detail in our previous study.\[^{22}\] First, in rats anesthetized with pentobarbital (50 mg/kg, intraperitoneal), a magnetized needle (the equivalent of 22 gauge, Daiso Sangyo, Hiroshima, Japan) was used to pierce the mentum skin below the lower lip of the rat. The needle was then bent into a ring, and the tip of the needle was removed. One week after piercing, the rats were trained for 10 min per day to stably protrude their perioral regions through a hole in a hand-made restrainer. After demonstrating stable behavior in the restrainer (~three to four days after the initiation of training), the rats were further trained to expose saline in WiM model rats under 2% isoflurane anesthesia immediately after removing the thick wire (1 hr prior to the pain tests), using a 30-G needle with a 50-μL Hamilton syringe (Hamilton, Reno, NV). RN-1734 was similarly injected into the same region in WiM model rats on day 2 after measuring pre-value of mechanical threshold. The concentrations of α-LA, A-317491, and RuR were determined according to previous studies,\[^{10,18,19}\] and that of RN-1734 was the same as the swab application. As a control, a vehicle solution (saline) was applied.

To examine whether the interaction of PAR2 and TRPV4 in the oral mucosa, elastase (Roche, Basel, Switzerland) and GSK1016790A (Cayman Chemical) were similarly injected at 10 μg/mL and 5 μg/μL, respectively, in 10 μL saline in naïve rats under 2% isoflurane anesthesia, after measuring pre-value of mechanical threshold. The concentrations of GSK1016790A and elastase were determined according to previous studies.\[^{20,21}\] GSK1016790A was diluted 100-times in ethanol, while the other drugs were diluted 10-times in DMSO. RN-1734 and FSLLRY-NH3 were coinjected with elastase to block TRPV4 and PAR2 activation, respectively. The concentrations of RN-1734 and FSLLRY-NH3 were the same as the swab application. As a control, a vehicle solution (1% ethanol- and 10% DMSO-containing saline) was applied.

**Behavioral observation of mouth rubbing**

Mouth rubbing with both forelimbs was measured over a constant time period (between 4 and 6 p.m.) as an oro-facial nociceptive behavior.\[^{22}\] Prior to the measurements, all rats were acclimated over a period of three days (once per day) to the plastic cage (30 × 30 × 30 cm) used for the observations. To evaluate spontaneous pain, the behavioral observation was performed for 10 min.
the oral mucosa. A small neodymium magnet (3 mm in diameter; Waki, Osaka, Japan), which was attached to a 4-g weight by a string, was attached to the pierced ring, resulting in a constant vertical pressure. When the rats flicked away and backed up, the magnet was released from the ring without damage to the pierced region. The following intraoral behavioral test was performed after confirming that each rat had adapted to the experimental conditions (~two to three weeks after piercing).

The head withdrawal threshold to mechanical stimulation was measured in the oral mucosa by applying von Frey filaments (0.02–6 g filaments, North Coast Medical, Morgan Hill, CA, and 0.2 and 0.3 g handmade filaments). These mechanical stimulations were applied to the oral mucosa in the labial fornix region of the inferior incisors, which involved oral mucositis in the WiM model. The head withdrawal threshold was defined as the minimum pressure required to evoke an escape. Each measurement was repeated five times at 3–4-min intervals, and the measurements were averaged following exclusion of the maximum and minimum values.\textsuperscript{13,23–25}

**Real-time reverse transcription-polymerase chain reaction**

Bilateral trigeminal ganglia were extracted from sham and WiM model rats on day 1 under deep anesthesia with pentobarbital (100 mg/kg, intraperitoneal), immediately transferred to ISOGEN (Wako) and frozen at −80°C. After homogenization of the trigeminal ganglia tissues of the third branch, which innervates the lower lip, total RNA was extracted using the RNeasy Mini Kit (Qiagen, Tokyo, Japan) and reverse-transcribed with random hexamers (Applied Biosystems, Foster, CA). Four replicates from four rats were assessed for each group. A 1.5-μl cDNA aliquot was amplified in a 7300 Real Time PCR System (Thermo Fisher Scientific) according to the manufacturer’s recommendations with THUNDERBIRD SYBR qPCR Mix (Toyobo, Osaka, Japan). The following primers were used: PTGER1 (EP1 gene), 5’-GGGCACTCATGGTGTCCC-3’ and 5’-GC CAACACCAATACCA-3’; F2RL1 (PAR2 gene), 5’-GGCTGCTGGGAGGTATCAC-3’ and 5’-CGTGT CGTGT-CCCG-3’; TRPV1, 5’-CAACAGGA AGGGGCTCAC-3’ and 5’-TCTGGAGAATGTAGGC CAAGAC-3’; TRPA1, 5’-ATTTGCAGCCCTGAG TTTT-3’ and 5’-GCGTACATCCATCTTGTCC-3’; TRPV4, 5’-CCACCCAGTGACAAACAG-3’ and 5’-GAGCTTTGGGGCTGTG-3’; β-actin, 5’-CCCC CGAGTACAACTTCT-3’ and 5’-CGTCTACCATG GCAGAAT-3’. The assays were performed in duplicate, and the ΔΔCT method was used to quantify the relative expression. There were no differences in the CT values (the cycle number when the fluorescent signal crosses the detection threshold) for β-actin between the two animal groups (data not shown). Hence, the given relative amount for each sample was standardized to that of β-actin, and the ratios were calculated relative to the average value for the naive group.

**Ca\textsuperscript{2+} imaging for trigeminal ganglion neurons**

After deep anesthesia with pentobarbital (100 mg/kg, intraperitoneal), the bilateral trigeminal ganglia of naive rats was removed, minced, and incubated in collagenase (2 mg/mL; Wako) and dispase (0.2 mg/mL; Sanko, Tokyo, Japan) in Ca/Mg-free HBSS for 1 h, as previously described.\textsuperscript{31} To remove debris, digested cell suspensions were centrifuged on 30% Percoll (Sigma–Aldrich). After washing, the cell pellet was resuspended in L-15 Lebovitz medium (Thermo Fisher Scientific) with antibiotic/antimycotic solution (0.1%), and the cells were plated on glass bottom dishes that had been coated with poly-ornithine and laminin. The plated cells were incubated at 37°C in a humidified chamber. To avoid the effects of culture and neurotrophin on cell properties,\textsuperscript{26,27} this study was performed using a short duration culture (3–8 h after cell plating) without any neurotrophins.

Before recording, trigeminal ganglion cells were incubated with fura-2AM (10 μM, Dojindo, Kumamoto, Japan) at 37°C for 30 min. The dishes were mounted on the stage of an inverted fluorescence microscope (IX71, Olympus, Tokyo, Japan) and washed with culture medium using a perfusion solution (mM): NaCl 130, KCl 5, MgCl\textsubscript{2} 1, CaCl\textsubscript{2} 2, glucose 10, and HEPES 10 (pH 7.4, adjusted with NaOH). The fura-2 was excited every 2 s by alternate illumination with 340 and 380 nm light using a Ca\textsuperscript{2+} imaging system (Hamamatsu Photonics, Hamamatsu, Japan). GSK1016790A, the TRPA1 agonist allyl isothiocyanate (AITC), and the TRPV1 agonist capsaicin (CPS) were diluted as stock solutions in ethanol and then further diluted 100-fold to 100 nM, 1 mM and 1 μM, respectively, with the perfusion solution. All drugs were applied for 2 min by bath application. To confirm whether the recorded cells were neurons, 50 mM KCl-supplemented perfusion solution was consistently applied at the end of the recordings. All Ca\textsuperscript{2+} responses are expressed as a ratio (F340/ F380), and the recorded cells were deemed to be sensitive to drug if they provided values of more than 0.01 for the Δ ratio after application. In a preliminarily experiment, 1% ethanol-containing solution did not change the ratio compared with the baseline (data not shown).

**Statistical analysis**

Data are presented as the mean ± SEM, and n represents the number of rats tested. An unpaired Student t test was used to compare differences between two different
groups or experimental days. To compare between-group differences in the number of CFUs, the Mann-Whitney U test was used. Following two-way repeated-measures analysis of variance, the Sidak post hoc test was applied to analyze daily or time changes between two different groups. Dunnett post hoc test was applied following one-way repeated-measures analysis of variance to analyze three or more groups. Significance was accepted at $P < 0.05$.

**Results**

**Generation of the WiM model**

On the following day (day 1) from installation of a bent wire, a clear traumatic ulcer was caused at the other free end of the wire-touched mucosal area of the labial fornix, as shown in Figure 1(c). The traumatic ulcer was limited to the region in contact with the wire, which differed visually from the acetic acid-induced ulcerative mucositis to the region in contact with the wire, which differed from the day 1 probably resulted from the wire installation procedure and powder food loading (to avoid detachment of non-host materials, oral bacteria, and food). After tail vein injection of Evans blue, the submucosal area involving the abscess on day 1 was stained with the drug (data not shown), suggesting a high level of extravasation in traumatic mucositis, including the abscess. To examine the involvement of mast cell degranulation in

The mean oral mucositis score of 5.7 on day 1 rapidly decreased to 3 on day 2 (Figure 1(f)), indicating a disappearance of the traumatic ulcer on day 2. Gradually, the mucositis was cured over five days (Figure 1(c)); the oral mucositis scores decreased each day (Dunnett post hoc test, $P < 0.05$ on day 3 and $P < 0.01$ on days 4 and 5, compared with day 1; Figure 1(f)). On day 1, the antibiotic and anti-inflammatory (indomethacin) pretreatments slightly decreased to a score of 5 and 4.3, respectively ($t$ test, $P < 0.05$ for both treatments; Figure 1(f) and (g)). Such limited efficacies of both the antibiotic and indomethacin pretreatments for oral mucositis severity have been similarly observed in the 5-fluorouracil-administered oral ulcerative mucositis model.10

MPO activity was significantly elevated in the WiM model compared with the sham, suggesting leukocyte infiltration into the oral mucosa of the model ($t$ test, $P < 0.01$; Figure 1(h)). In the ELISA assays, IL-1β, IL-6, and TNF-α were upregulated on day 1 in the WiM model relative to the sham ($t$ test, $P < 0.01$ for IL-1β and $P < 0.05$ for IL-6 and TNF-α; Supplementary Figure 1(a)–(c)). However, there was no significant difference in bradykinin, which is a well-known inflammatory pain inducer,11 between the sham and WiM model (Supplementary Figure 1(d)).

**Histological appearance**

The histological assessment revealed an obvious traumatic ulcer and mucosal swelling of the oral mucosal tissue in the labial fornix region in the WiM model on day 1 relative to the sham (Figure 1(i)), consistent with the visual oral mucositis score (Figure 1(f)). Furthermore, the histology demonstrated the formation of a clear abscess, which formed with leukocyte accumulation, of more than 1 mm in the submucosal area, in addition to broad inflammatory cell invasion (Figure 1(i)). The histological structure was further confirmed as an abscess based on negative images with PAS staining (Figure 1(j)) and Alcian blue staining (data not shown), which detects the mucus cyst. Such abscess formation has been suggested in a clinical review.6 The abscess was gradually reduced each day (Figure 1(i)), and it difficult to detect on days 4 and 5.

PAS staining revealed PAS-positive cell wall-like particles in the center of the abscess, which were derived from sawdust and powdered foods (inset microphotograph of a high-power field; Figure 1(j)). However, abscess formation was observed even under experimental conditions without sawdust and powdered foods and after antibiotic pretreatment (data not shown), suggesting that it was primarily derived from damaged host cells rather than non-host materials, oral bacteria, and food.
the oral mucositis, the oral mucosal tissue sections of the labial fornix region were stained with toluidine blue. Many toluidine blue-positive granule-containing mast cells were detected in the submucosal layer of the healthy oral mucosa, but such toluidine blue-positive cells were rarely detected around the traumatic ulcer and abscess in the WiM model (Supplementary Figure 1(e)), indicating exhaustion of the mast cell granules.

**Spontaneous pain and mechanical allodynia**

Spontaneous pain was evaluated by quantifying the spontaneous mouth rubbing behaviors for 10-min periods, and mechanical allodynia was evaluated based on the head withdrawal threshold by von Frey filaments using the SIO method, which is a proprietary assay system for conscious rats. Figure 2(a) and (b) shows the daily changes in spontaneous mouth rubbing and the head withdrawal threshold, respectively, in sham and WiM model rats. After removing the thick wire, relative to the sham, WiM model rats showed increased spontaneous mouth rubbing, peaking on day 1 (Sidak post hoc test, \( P < 0.01 \)) followed by a rapid recovery to pretreatment levels on day 2 (Figure 2(a)), and reduced head withdrawal thresholds, peaking on day 1 (Sidak post hoc test, \( P < 0.01 \) on days 1 and 2 and \( P < 0.05 \) on day 3) and gradually recovering to control levels up to day 4 (Figure 2(b)). These results suggest the induction of spontaneous pain and mechanical allodynia in the WiM model.

Importantly, in contrast to the model of acetic acid-induced oral ulcerative mucositis, \(^{10}\) antibiotic pretreatment did not significantly suppress the induction of spontaneous pain and mechanical allodynia in the WiM model (Figure 2(a) and (b)). To examine the impact of bacterial loading, we quantified bacterial infections in the traumatic ulcerative region in the model. The numbers of CFUs under aerobic and anaerobic conditions on day 1 were significantly increased approximately 100-fold compared with the healthy oral mucosa of the sham (Mann-Whitney \( U \) test, \( P < 0.05 \) for both conditions), and the increments were suppressed by antibiotic pretreatment (Mann-Whitney \( U \) test, \( P < 0.05 \) for both conditions) (Figure 2(c)). Together with the decrease in the oral mucositis score following antibiotic pretreatment, these results indicated that the antibiotic pretreatment was effective for the suppression of bacterial infection. In previous studies investigating infectious inflammatory pain, \(^{10,28}\) spontaneous pain, and/or mechanical allodynia were induced from over \( 10^4 \) CFU per milligram of tissue. Therefore, in this study, the number of CFUs in tissue was regarded as the pain-producing level. The increment levels of bacterial loading in the WiM model were insufficient to generate a pain-producing level (Figure 2(c)).

**Spontaneous pain mechanism**

The anti-inflammatory pretreatment using the COX-1/2 inhibitor indomethacin significantly inhibited the

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**Figure 2.** Spontaneous pain and mechanical allodynia induction in the wire-induced mucositis (WiM) model and effect of antibiotic pretreatment. As the antibiotic pretreatment, sulfamethoxazole- and trimethoprim-containing drinking water was supplemented beginning two days prior to the procedure (installation of the thick wire) and continued for five days. (a) Daily changes in spontaneous mouth rubbing periods in the sham and model rats with or without antibiotic pretreatment (\( n = 5, 4, \) and 5, respectively). (b) Daily changes in the head withdrawal threshold by von Frey stimulation in sham and model rats with or without antibiotic pretreatment (\( n = 5, 4, \) and 5, respectively). In (a) and (b), * and ** indicate \( P < 0.05 \) and 0.01, respectively, compared with the sham; Sidak post hoc test following two-way repeated-measures analysis of variance. (c) The numbers of CFU under aerobic and anaerobic conditions in extracted oral mucosal tissues (each group, \( n = 4 \)). The horizontal dashed line in the graph indicates the pain-producing level, according to previous studies investigating bacterial infection-induced pain.\(^{10,28}\) *\( P < 0.05 \) compared with sham; **\( P < 0.05 \) compared without antibiotic pretreatment in the WiM model; Mann-Whitney \( U \) test.
increase in spontaneous mouth rubbing on day 1 in the WiM model (t test, \( P < 0.05 \); Figure 3(a)). Consistently, the inflammation regulatory enzyme, COX-2, was upregulated on day 1 in the WiM model relative to the sham (t test, \( P < 0.05 \); Figure 3(b)). Furthermore, swab application of the PGE2 receptor subtype 1 EP1-selective antagonist, ONO-8711, significantly inhibited the increase in spontaneous mouth rubbing on day 1 in the WiM model (t test, \( P < 0.05 \); Figure 3(c)). To confirm the production of prostanoids, we measured the contents of PGE2 and 15d-PGJ2 in the oral mucosal regions of sham and WiM model rats on day 1 and/or 2 by ELISA. The 15d-PGJ2 is a cyclopentenone PG that is produced by dehydration within the cyclopentane ring of PGD2.29 The two distinctly formed prostanoids were significantly upregulated in the WiM model on day 1 relative to the sham (Dunnett post hoc test, \( P < 0.01 \) in PGE2 and \( P < 0.05 \) in 15d-PGJ2), and the increments were decreased on day 2 to levels mimicking the sham (Figure 3(d) and (e)). Thus, the time course and drug sensitivities of spontaneous pain induction were consistent with those of prostanoid production.

PGE2 injection induces TRPV1-, TRPA1-, and TRPV4-mediated pain, and cyclopentenone PGs directly activate TRPA1.29–32 Therefore, to examine the involvements of the nociceptive TRP channels in spontaneous pain following mechanical trauma, we investigated the effects of the TRPV1, TRPA1, and TRPV4 antagonists, SB-366791, HC-030031, and RN-1734, respectively, on spontaneous mouth rubbing in the WiM model.

**Figure 3.** Mechanism underlying spontaneous pain in the wire-induced mucositis (WiM) model. (a) Spontaneous mouth rubbing after intraperitoneal (i.p.) indomethacin pretreatment (Indo) and vehicle (Veh; 0.1 M Tris-buffered saline) on day 1 (each group, \( n = 9 \)). \( * \) \( P < 0.05 \) compared with Veh; Student t test. (b) Cyclooxygenase-2 (COX-2) level in the oral mucosa of sham and WiM on day 1 (each group, \( n = 4 \)) by ELISA. \( * \) \( P < 0.05 \) compared with the sham; Student t test. (c) Spontaneous mouth rubbing after swab application of the EP1 antagonist ONO-8711 and Veh (10% dimethylsulfoxide [DMSO] -containing saline) on day 1 (each group, \( n = 6 \)). \( * \) \( P < 0.05 \) compared with Veh; Student t test. (d and e) Prostaglandin E2 and 15-deoxy-\( \Delta_{12,14} \)-PGJ2 (known as a TRPA1 agonist) levels in the oral mucosa of the sham on day 1 and the WiM model on days 1 and 2 (each group, \( n = 4 \)) by ELISA. \( * \) and \( ** \) indicate \( P < 0.05 \) and 0.01, respectively, compared with the sham on day 1; Dunnett post hoc test following one-way repeated-measures analysis of variance (ANOVA). (f) Spontaneous mouth rubbing after i.p. administration of SB-366791 (SB: a TRPV1 antagonist), HC-030031 (HC: a TRPA1 antagonist), RN-1734 (RN: a TRPV4 antagonist), and Veh (10% DMSO-containing saline) in the WiM model on day 1 (each group, \( n = 5 \)). \( ** \) \( P < 0.01 \) compared with Veh; Dunnett post hoc test following one-way repeated-measures ANOVA. (g) Spontaneous mouth rubbing after swab application of SB, HC alone, a mixture of SB and HC (SB + HC) and Veh (10% DMSO-containing saline) in the WiM model on day 1 (each group, \( n = 5 \)). \( * \) and \( ** \) indicate \( P < 0.05 \) and 0.01, respectively, compared with Veh; Dunnett post hoc test following one-way repeated-measures ANOVA.
The increase in spontaneous pain on day 1 was suppressed by intraperitoneal administrations of SB-366791 and HC-030031 (Dunnett post hoc test, \( P < 0.01 \) for both drugs) but not by RN-1734 (Figure 3(i)), suggesting a dependency of TRPV1 and TRPA1 channel opening and TRPV4 channel closing in the WiM model. Furthermore, swab application of either or both SB-366791 and HC-030031 to the traumatic ulcer region for 5 min significantly suppressed the spontaneous pain (Dunnett post hoc test, \( P < 0.05 \) for SB-366791 and HC-030031 alone and \( P < 0.01 \) for a mixture of SB-366791 and HC-030031; Figure 3(g)). The significant efficacies of SB-366791 and HC-030031 in the swab application suggest that TRPV1 and TRPA1 activation are induced peripherally in traumatic mucositis in this model.

**Mechanism of mechanical allodynia**

In contrast to spontaneous pain, mechanical allodynia in the WiM model on day 1 was resistant to pretreatment with indomethacin (Figure 4(a)). Furthermore, the mechanical allodynia was not inhibited by submucosal injections of the reactive oxygen species inhibitor \( \alpha \)-LA and the nociceptive ionotropic ATP receptor P2X3 and P2X2/3 antagonist A-317491, while it was significantly improved by the non-selective TRP channel antagonist RuR (Figure 4(b)). These results suggest that there is no involvement of the COX pathway, reactive oxygen species, or ATP in mechanical allodynia. Based on the lack of efficacy of the antibiotic pretreatment (Figure 2(a) and (b)), bacterial toxins (lipopolysaccharide and N-formyl peptides), which have been identified to induce
mechanical allodynia in a 5-fluorouracil-administered oral ulcerative mucositis model, might not largely contribute to mechanical allodynia in the WiM model. Another possibility was the activation of PAR2 because of its abundant expression in sensory neurons and involvement in inflammatory pain. Swab application of the PAR2 selective antagonist FSLLRY-NH₂ significantly increased the declined head withdrawal threshold in the model compared with the vehicle (Dunnett post hoc test, \( P < 0.05 \); Figure 4(c)). The same application did not alter the increase in spontaneous mouth rubbing (Figure 4(d)).

Due to the granule exhaustion in mast cells on day 1, there was very little involvement of mast cell-derived serine protease tryptase, potentially causing PAR2 activation. Recently, neutrophil elastase has been reported as a biased PAR2 agonist. It is possible that the enzyme is abundant in the traumatic mucositis of the WiM model because of the accumulation of neutrophils (abscess formation) for a few days (Figure 1(i)), consistent with the duration of mechanical allodynia induction (Figure 2(b)). As expected, a significant upregulation of neutrophil elastase was detected in the WiM model, not only on day 1 but also day 2 (Dunnett post hoc test, \( P < 0.05 \) for both days; Figure 4(e)).

To examine the involvements of the nociceptive TRP channels in mechanical allodynia following mechanical trauma, we investigated the effects of the TRP channel subtype specific antagonists, SB-366791, HC-030031, and RN-1734, on head withdrawal threshold in the WiM model on day 1 and/or 2. The declined head withdrawal threshold on day 1 was suppressed by systemic administration of RN-1734 (Dunnett post hoc test, \( P < 0.05 \)) but not by those of SB-366791 and HC-030031 (Figure 4(f)). Furthermore, swab application of RN-1734 to the traumatic ulcer region for 5 min significantly suppressed mechanical allodynia on day 1 (Dunnett post hoc test, \( P < 0.01 \)) but not on day 2 (Figure 4(g)). Because the ulcer in the model quickly disappeared by day 2 (Figure 1(f)), the drug in the swab might be insufficient to permeate the deeper mucosal area on that day. To exert sufficient efficacy of the drug in the target tissue, we submucosally injected RN-1734 on day 2. The submucosally injected RN-1734 significantly increased the head withdrawal threshold 1 and 2 h compared with vehicle injection (Sidak post hoc test, \( P < 0.05 \) for both time points; Figure 4(g)).

**Elastase-induced TRPV4-mediated mechanical allodynia in naive rats**

In naive rats, submucosal injection of the TRPV4-selective agonist GSK1016790A significantly reduced the head withdrawal threshold 1 h after the injection compared with the vehicle injection (Sidak post hoc test, \( P < 0.01 \); Figure 5(a)). Submucosal injection of elastase significantly diminished the head withdrawal threshold 1 h after the injection compared with vehicle injection (Sidak post hoc test, \( P < 0.01 \)), and this response was blocked by the coinjection of RN-1734 (Sidak post hoc test, \( P < 0.05 \) for 1 and 2 h; Figure 5(a)). Furthermore, the reduced head withdrawal threshold following elastase injection was also blocked by coinjection of FSLLRY-NH₂ \( (n = 5, P < 0.01\text{ for }1\text{ h with elastase alone; data not shown}) \). Submucosal injection of FSLLRY-NH₂ or RN-1734 alone did not change head withdrawal threshold \( (n = 5\text{ in each group; data not shown}) \), indicating no anesthetic effects of the drug themselves. These results suggest the occurrence of PAR2-induced TRPV4-mediated mechanical allodynia following elastase injection into the oral mucosa.

There were no significant differences in the mRNA expression levels of EP₁, PAR₃, TRPV1, TRPA1, and TRPV4 in the trigeminal ganglion between the sham and WiM model (Figure 5(b)), which indicated that expression of the receptors and channels in the WiM model was equivalent to those in naive animals.

**Coexpression of TRPV1, TRPA1, and TRPV4**

QX-314 is a membrane-impermeable anesthetic, but it passes into the intracellular space via the channel pores of TRPV1 and TRPA1. Swab application of 1% QX-314 for 5 min significantly suppressed spontaneous pain in the WiM model (Sidak post hoc test, \( P < 0.01 \), compared with vehicle; Figure 4(d)), supporting the contributions of continuous TRPV1 and TRPA1 opening to the spontaneous pain mechanism. Despite possible TRPV4 closure during resting conditions (Figure 3(f)), the same application also suppressed the mechanical allodynia (Sidak post hoc test, \( P < 0.01 \), compared with vehicle; Figure 4(c)). The suppression of mechanical allodynia by QX-314 application was abrogated by the blockade of TRPV1 and TRPA1 following systemic administration of SB-366791 and HC-030031 (Figure 5(c)). These results suggest that TRPV4-mediated mechanical allodynia is suppressed by QX-314 passing through TRPV1 and TRPA1 pore routes rather than the TRPV4 pore.

To confirm the coexpression of TRPV1, TRPA1, and TRPV4 in peripheral nerve fibers, we investigated the Ca²⁺ response to GSK1016790A, AITC, and CPS in dissociated trigeminal ganglion neurons of naive rats. Of 218 neurons from 3 rats that were responsive to high K⁺ solution, 164 (75%) showed slowly activated long-lasting Ca²⁺ responses to GSK1016790A, 89 (41%) showed transient Ca²⁺ responses to AITC, and 113 (52%) showed long-lasting Ca²⁺ responses to CPS, as shown in Figure 5(d). In the GSK1016790A-sensitive neurons, 98 (60%) neurons responded to either AITC and/or CPS (Figure 5(e)), indicating that a significant population expressed TRPV4 and TRPV1/TRPA1 in the trigeminal nerve system.
Discussion

The present study is the first report to demonstrate prostaglandin- and PAR2-dependent TRP channel-mediated pain following mechanical trauma in the oral mucosa (Figure 6). The pathophysiological status of the oral traumatic mucositis showed tissue injury-induced inflammation rather than infectious inflammation, together with no efficacy of the antibiotic pretreatment. During dental treatment, iatrogenic oral traumatic mucositis caused by intraoral appliances has received little attention because patient complaints are transient after removal of the injurious parts. However, many patients suffer such trauma-induced pain, leading to poor quality of life and decreased cooperation in therapy. The results in this study provide an understanding of the pain mechanism associated with oral traumatic mucositis and effective analgesic approaches for many dental clinicians.

Pathophysiological properties of traumatic oral mucositis

During the one-day installation, repetitive wire-tip attachment to the oral mucosa during jaw movements when eating, drinking, and grooming would induce mechanical inflammation with tissue swelling and generate a traumatic ulcer on the mucosal surface following puncture of the wire-tip. The cellular damage induced leukocyte infiltration and the formation of an abscess...
in the submucosal layer until day 1 (Figure 6). The level of phagocytosis by the abundant neutrophils in the injured mucosa was also effective against bacterial infection caused by leukocyte infiltration due to cell damage. After removing the appliance on day 1, the accumulation of leukocytes forms an abscess below the traumatic ulcer. The ulceration was cured by day 2, but the abscess persisted for a few days. On day 1, proinflammatory cytokines promote prostanoid production in a cyclooxygenase-dependent manner on day 1, and the abscess releases elastase over a longer duration up to days 4 to 5. Arrows with double lines represent chemical release. Prostaglandin E₂ (PGE₂) stimulates EP₁, followed by thermal sensitization of TRPV₁ on peripheral sensory fibers, resulting in activation at body temperature (B.T.). The cyclopentane prostanoid 15-deoxy-Δ¹²,¹⁴-prostaglandin J₂ (15d-PGJ₂), which originates from prostaglandin D₂ (PGD₂), directly activates TRPA₁ on peripheral sensory fibers, in addition to TRPV₁ activation-induced sensitization. Elastase stimulates PAR₂, followed by mechanical sensitization of TRPV₄ on peripheral sensory fibers, which is known to be activated by stretch of the cell membrane. Black arrows with solid lines represent agonistic effects of the chemicals on the receptors. Thick blue arrows with twisted lines represent intracellular pathways of TRP channel sensitization. Dashed arrows represent cation influx through channels, resulting in the generation of an action potential following depolarization. QX-314 passes into the intracellular space via the channel pores of TRPV₁ and TRPA₁, as indicated by a clear blue arrow, and then blocks voltage-dependent sodium channels, providing an anesthetic for both spontaneous pain and mechanical allodynia.

Figure 6. Schematic of the mechanisms underlying oral mucositis-induced pain following mechanical trauma. Following mechanical injury caused by the intraoral appliance, the oral mucosa demonstrates a traumatic ulcer with inflammation that is characterized by leukocyte infiltration due to cell damage. After removing the appliance on day 1, the accumulation of leukocytes forms an abscess below the traumatic ulcer. The ulceration was cured by day 2, but the abscess persisted for a few days. On day 1, proinflammatory cytokines promote prostanoid production in a cyclooxygenase-dependent manner on day 1, and the abscess releases elastase over a longer duration up to days 4 to 5. Arrows with double lines represent chemical release. Prostaglandin E₂ (PGE₂) stimulates EP₁, followed by thermal sensitization of TRPV₁ on peripheral sensory fibers, resulting in activation at body temperature (B.T.). The cyclopentane prostanoid 15-deoxy-Δ¹²,¹⁴-prostaglandin J₂ (15d-PGJ₂), which originates from prostaglandin D₂ (PGD₂), directly activates TRPA₁ on peripheral sensory fibers, in addition to TRPV₁ activation-induced sensitization. Elastase stimulates PAR₂, followed by mechanical sensitization of TRPV₄ on peripheral sensory fibers, which is known to be activated by stretch of the cell membrane. Black arrows with solid lines represent agonistic effects of the chemicals on the receptors. Thick blue arrows with twisted lines represent intracellular pathways of TRP channel sensitization. Dashed arrows represent cation influx through channels, resulting in the generation of an action potential following depolarization. QX-314 passes into the intracellular space via the channel pores of TRPV₁ and TRPA₁, as indicated by a clear blue arrow, and then blocks voltage-dependent sodium channels, providing an anesthetic for both spontaneous pain and mechanical allodynia.

in the submucosal layer until day 1 (Figure 6). The level of phagocytosis by the abundant neutrophils in the injured mucosa was also effective against bacterial infection caused by the traumatic ulcer, maintaining a low infection level. We failed to detect differences in bradykinin, which is the most potent extravasation factor, between healthy oral mucosa and traumatic ulcer on day 1, despite the high level of extravasation in the submucosal layer based on the Evans blue test. The activity of bradykinin was likely completed by day 1 in our model. Using the same analogy, mast cell degranulation in the oral traumatic mucositis seemed to be completed by day 1. Therefore, these pain-producing factors might play an initial role in the inflammatory activity, but not contribute to a large extent after day 1. However, the proinflammatory cytokines, COX-2, PGE₂, and 15d-PGJ₂, were significantly upregulated on day 1, suggesting their involvements in the pain mechanism in this model.

Prostanoid-dependent TRPV₁- and TRPA₁-mediated spontaneous pain

Spontaneous pain was inhibited by indomethacin pretreatment, the PGE₂ receptor EP₁ antagonist and TRPV₁ and TRPA₁-selective antagonists, but not by PAR₂ and TRPV₄-selective antagonists. Together with the upregulation of PGE₂ and 15d-PGJ₂ only on day 1, these results suggest that COX-dependent prostanoid
production elicited spontaneous pain through indirect TRPV1 activation following EP1 activation by PGE2 and direct TRPA1 activation by 15d-PGJ2 (Figure 6). Because EP1 activation has been reported to reduce the thermal threshold of TRPV1 below body temperature through phosphorylation, the sensitized TRPV1 of traumatic mucositis-innervated peripheral fibers would be continuously activated during resting conditions by body temperature. Based on the direct activation of TRPA1 by the cyclopentane prostanoid 15d-PGJ2, TRPA1 would be activated continuously under resting conditions. The significant efficacy of the membrane-impermeable, TRPV1/TRPA1-permeable anesthetic QX-314 in swab applications supports TRPV1 and TRPA1 channel opening.

The suppression of spontaneous pain following swab application of these antagonists suggests that the spontaneous pain mechanism occurred primarily in the traumatic ulcer region. Spontaneous pain induction and prostanoid upregulation occurred only on day 1 and disappeared by day 2. The timecourse change corresponded to the development and disappearance of traumatic ulcer, suggesting that the painful region within the traumatic mucositis was an ulcer rather than an abscess.

PAR2-dependent TRPV4-mediated mechanical allodynia

In contrast to spontaneous pain, mechanical allodynia was not suppressed by indomethacin pretreatment and TRPV1 and TRPA1-selective antagonists, in addition to the antioxidative drug and P2X3 and P2X2/3 antagonist. Finally, we observed significant suppression of the PAR2 receptor antagonist in terms of mechanical allodynia. Based on the complete degranulation of mast cells containing tryptase up to day 1, the serine protease does not seem to contribute to PAR2 activation. Recently, neutrophil elastase has been reported to sensitize TRPV4 following PAR2 activation through a distinct intracellular route due to the activation of the representative PAR2 agonist trypsin. The timecourse of neutrophil elastase upregulation was consistent with those of abscess formation and mechanical allodynia induction. The TRPV4 antagonist significantly suppressed mechanical allodynia on days 1 and 2 by swab application and submucosal injection, respectively. Swab application of the TRPV4 antagonist demonstrated no efficacy on day 2, potentially due to the rapid regeneration of the oral mucosal barrier (Figure 6). Furthermore, in naïve rats, submucosal injections of TRPV4 agonist and elastase induced mechanical allodynia, and the elastase-induced mechanical allodynia was blocked by coinjection of TRPV4 or PAR2 antagonist. These results suggest that elastase released from the abscess elicited mechanical allodynia due to mechanical sensitization of TRPV4 through PAR2 activation (Figure 6).

The resistance of spontaneous pain to TRPV4 antagonist in the model suggests TRPV4 channel closure during resting conditions and activation only following mechanical stimulation of oral mucositis. Nevertheless, swab application of QX-314 significantly suppressed mechanical allodynia on day 1. No reports have demonstrated the passage of QX-314 through TRPV4 channel pores. After systemic administration of TRPV1 and TRPA1 antagonist, the anesthetic effect of QX-314 on mechanical allodynia disappeared, suggesting that TRPV4-mediated mechanical allodynia was suppressed by the passage of QX-314 through TRPV1 and/or TRPA1 channel pores. In rats and humans, TRPV4 is abundantly expressed in peripheral sensory neurons, although its expression in mice is debated. In this study, more than 60% of the GSK1016790A-sensitive trigeminal sensory neurons responded to AITC and/or CPS. Taking into consideration the abundant co-expression of TRPV4 and TRPV1/TRPA1, it is likely that the mechanical allodynia-inducible neuronal population partially consisted of a spontaneous pain-inducible subpopulation (Figure 6). However, additional approaches will be required to clear analgesic mechanism of QX-314 in WiM model, for example, electrophysiological experiments for continuous TRPV1/TRPA1 channel opening with QX-314 passing.

Differences in comparison to the infectious oral ulcerative mucositis model

In our previous study using a 5-fluorouracil-administered rat model treated with acetic acid, severe oral ulcerative mucositis with excessive bacterial infection, prostanoid-dependent TRPV1-mediated spontaneous pain, and bacteria toxin-dependent TRPA1-mediated mechanical allodynia were observed, which were largely different from the pain mechanism observed in the WiM model, excluding the efficacy of indomethacin pretreatment for spontaneous pain and mechanical allodynia and the involvement of TRPV1 in spontaneous pain. The administration of 5-fluorouracil induced leukopenia and led to deficient leukocyte infiltration into the injured region, indicating that the activity of neutrophils was largely eliminated. The large difference in the pain mechanism between the models led to differences in bacterial infection levels together with leukocyte functions between the models. In the 5-fluorouracil-administered oral ulcerative mucositis model, lipopolysaccharide and N-formyl peptides were derived from extremely infected oral regions that were sensitized to mechanical activation of TRPA1, directly and indirectly via receptor activation. However, in the WiM model, abundant 15 d-PGJ2 production in traumatic mucositis resulted in continuous TRPA1 channel opening, resulting in spontaneous pain but not mechanical allodynia. Furthermore, the release
of elastase from neutrophils in the abscess elicited TRPV4-mediated mechanical allodynia via PAR2 activation without activation/sensitization of TRPV1 and TRPA1. Importantly, PAR2 activation by elastase has been reported to induce mechanical pain but not heat hypersensitivity, which is caused by TRPV1 sensitization.

**Clinical significance**

The results of this study supply basic scientific evidence for abscess formation in the submucosal layer following the installation of an intraoral appliance in dental patients, as suggested by a clinical review,6 and indicates that the pathophysiological status of traumatic mucositis is not equivalent to other types of ulcerative mucositis. During dental orthodontic and prosthodontic treatments, the installation of intraoral appliances is needed to improve oral functions during meals and esthetic dentition during conversation. Unfortunately, many dental patients suffer from severe pain in the mechanical lesion caused by the intraoral appliance.1–4 The pain mechanism underlying traumatic mucositis-induced pain elucidated in this study (Figure 6) suggests that some effective pain treatments will solve pain-related poor quality of life and low cooperation in therapy in dental patients.7,8 According to the pain relief observed in the WiM model on day 1 following swab applications of various drugs, topical treatments of drug-containing ointment and mouthwash may be effective approaches for patients with mechanically induced ulcerative mucositis. Alternately, systemic oral application of non-steroidal anti-inflammatory agents is also effective for the suppression of spontaneous pain based on the prostanoid dependency of the pain modality in this study. However, the ulcer is quickly healed, but submucosal inflammation, including abscess, persists over subsequent days, suggesting continuation of mechanical allodynia induction in patients following the removal and correction of the injurious parts of orthodontic and prosthodontic appliances. To resolve the continuous pain complaints after being subjected to the intraoral appliance,1,7,8 systemic administration of PAR2 antagonistic or neutrophil elastase inhibitory drugs seems to be effective.34,45

In conclusion, prostanoids and PAR2 activation elicit spontaneous pain through TRPV1 and TRPA1 and mechanical alldynia through TRPV4, respectively, independently of bacterial infection, in oral mucositis following mechanical trauma. The pathophysiological pain mechanism suggests that analgesic approaches are effective for dental patients suffering from mucosal trauma-induced pain. To more clarify involvements of TRPV1, TRPA1, and TRPV4 in mucosal trauma-induced pain, it is necessary to perform further genetic approaches (gene knockdown by in vivo siRNA and CRISPR/Cas9 system).

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**Author contributions**

MI, KO, SH, TN, TS, NH, and KY performed animal experiments, biochemical, and molecular biological testing, and Ca2+-imaging and analyzed the data; KO, KG, RH, TK, and KI designed experiments, supervised research, and wrote the manuscript. All authors read and approved the final manuscript.

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