Mucoadhesive Electrospun Patch Delivery of Lidocaine to the Oral Mucosa and Investigation of Spatial Distribution in a Tissue Using MALDI-Mass Spectrometry Imaging

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Supporting Information

ABSTRACT: Many oral mucosal conditions cause considerable and prolonged pain that to date has been difficult to alleviate via topical delivery, and the use of injection causes many patients dental anxiety and needle-prick pain. Therefore, developing a noninjectable drug delivery system as an alternative administration procedure may vastly improve the health and wellbeing of these patients. Recent advances in the development of mucoadhesive electrospun patches for the direct delivery of therapeutics to the oral mucosa offer a potential solution, but as yet, the release of local anesthetics from this system and their uptake by oral tissue have not been demonstrated. Here, we demonstrate the fabrication of lidocaine-loaded electrospun fiber patches, drug release, and subsequent uptake and permeation through the porcine buccal mucosa. Lidocaine HCl and lidocaine base were incorporated into the electrospun patches to evaluate the difference in drug permeation for the two drug compositions. Lidocaine released from the lidocaine HCl-containing electrospun patches was significantly quicker than from the lidocaine base patches, with double the amount of drug released from the lidocaine HCl patches in the first 15 min (0.16 ± 0.04 mg) compared to that from the lidocaine base patches (0.07 ± 0.01 mg). The permeation of lidocaine from the lidocaine HCl electrospun patches through ex vivo porcine buccal mucosa was also detected in 15 min, whereas permeation of lidocaine from the lidocaine base patch was not detected. Matrix-assisted laser desorption ionization-mass spectrometry imaging was used to investigate localization of lidocaine within the oral tissue. Lidocaine in the solution as well as from the mucoadhesive patch penetrated into the buccal mucosal tissue in a time-dependent manner and was detectable in the lamina propria after only 15 min. Moreover, the lidocaine released from lidocaine HCl electrospun patches retained biological activity, inhibiting veratridine-mediated opening of voltage-gated sodium channels in SH-SYSY neuroblastoma cells. These data suggest that a mucoadhesive electrospun patch may be used as a vehicle for rapid uptake and sustained anesthetic drug delivery to treat or prevent oral pain.

KEYWORDS: electrospinning, lidocaine, oral mucosa, ex vivo tissue permeation, MALDI-mass spectrometry imaging, drug distribution in tissue

INTRODUCTION

Control of pain is a major unmet clinical need in a range of oral conditions including mucositis, ulceration, and trauma. The effects of pain may be so severe as to affect eating, drinking, speech, or sleep. Pain is also associated with a number of dental procedures and is most commonly alleviated by the injection of a local anesthetic. Unfortunately, the use of injection is a major factor in dental anxiety and nonattendance at the clinic. A wide range of technologies to alleviate localized pain in the oral cavity have been investigated, including tablets, films, gels, and mouth washes. These are chiefly based on topical delivery of drugs, but all current approaches have limitations related to the very challenging conditions found within the oral cavity, where salivary flow and tissue movement contribute to a hostile mechanical and chemical environment.

We have recently fabricated an electrospun patch with mucoadhesive properties with the potential to deliver drugs to oral lesions for a sustained period. The patch displays good adhesion to oral tissues and is well tolerated with no significant cytotoxicity or irritation in both a porcine model and healthy conditions.
volunteers. The delivery of clobetasol-17-propionate was demonstrated in a tissue-engineered oral mucosal model and in ex vivo porcine mucosa. Electrospinning can create relatively thin-fiber (typically 100 nm to 50 μm fiber diameter) patches, where the polymer and solvent selection, as well as the electrospinning conditions, may alter the mechanical properties of the mat and its behavior in the oral cavity. These electrospin patches have high mucoadhesive properties due to the high surface area to volume ratio and to the incorporation of a hygroscopic polymer that rapidly absorbs saliva at the site of application leading to rapid and prolonged drug release. However, electrospun patches have not been subject to a detailed investigation of transbuccal delivery of lidocaine. Porcine mucosa is frequently used in transbuccal permeation experiments since it is physiologically similar to that of humans, and is, therefore, considered the gold standard for modeling oral drug delivery. Ex vivo permeation studies provide quantitative data; however, to visualize the spatial distribution of drugs in the tissue, mass spectrometry imaging (MSI) may be used. MSI allows for a multitude of compounds to be detected including the drug, metabolites, excipients, and endogenous compounds, such as lipids, which may serve as tissue biomarkers. There are a number of different MSI techniques, with matrix-assisted laser desorption ionization (MALDI) MSI being one of the most commonly used. The method requires deposition of a matrix on the tissue sample to aid the ionization of analytes when targeted by the laser. The process is relatively nondestructive, and, therefore, the tissue sample can be stained using hematoxylin and eosin (H&E) for histological evaluation after the process. We have previously used MALDI-MSI to examine permeation of nicotine across porcine buccal mucosa, and a similar experimental setup was used in this study to determine the time-dependent permeation of lidocaine.

Herein, we demonstrate for the first time the targeted permeation of lidocaine into ex vivo porcine buccal mucosa released from mucoadhesive electrospin patches to be used as a local oromucosal drug delivery vehicle. Both lidocaine HCl and lidocaine base were successfully incorporated into the electrospin patches, for which the release, ex vivo permeation, and bioactivity were measured. Our findings show a greater release of lidocaine HCl from the electrospin patches compared to lidocaine base, and only lidocaine permeation from the lidocaine HCl patches was detected ex vivo. Additionally, using MALDI-MSI, the localization and time-dependent penetration of the drug within the tissue were established. This study shows that lidocaine-loaded electrospin patches deliver functionally active drug directly to the buccal mucosa in a therapeutically useful timeframe. This drug delivery device has the potential of improving the control of localized oral pain that is currently a major unmet clinical problem.

### METHODS

**Materials.** Poly(vinylpyrrolidone) (PVP) (M<sub>W</sub> 2000 kDa) and Eudragit RS100 (M<sub>W</sub> 30 kDa) were kindly donated by BASF (U.K.) and Evonik Industries AG (Germany), respectively. Poly(caprolactone) (PCL) (M<sub>W</sub> 80 kDa), poly(ethylene oxide) (PEO) (M<sub>W</sub> 400 kDa), trifluoroacetic acid (TFA, 99%), 2,5-dihydroxybenzoic acid (DHB, 98%), carboxymethylcellulose (CMC), potassium dihydrogen phosphate, disodium hydrogen phosphate, sodium hydroxide, alcoholic eosin Y solution, hematoxylin solution Mayer’s, fetal calf serum, nonessential amino acids, penicillin, streptomycin, ethylenediaminetetraacetic acid (EDTA), dimethylsulfoxide (DMSO) were purchased from Sigma-Aldrich (U.K. & DK). Lidocaine HCl (M<sub>W</sub> 270.8 Da), lidocaine (M<sub>W</sub> 234.3 Da), and veratridine were purchased from Abcam (U.K.). Solvents such as ethanol (≥99.8%), chloromethane (DCM, ≥99%), dimethylformamide (DMF, 99.98%), and acetonitrile (ACN, ≥99.9%), media such as DMEM/Ham’s F12 with L-glutamine and Fluo-4 Direct assay kit (Invitrogen) were purchased from Fisher Scientific (U.K.). Human neuroblastoma cell line SH-SY5Y was purchased from LGC Standards (U.K.). Methanol (≥99.8%) and sodium chloride were obtained from Th. Geyer GmbH and Co. KG (Germany). Ultrapure water was collected from the water system SG Ultra Clear 2002 from Evoqua Water Technologies LLC. Isotonic phosphate-buffered saline (PBS) was prepared by dissolving 2.38 g of disodium hydrogen phosphate with 0.19 g of potassium dihydrogen phosphate and 8 g of sodium chloride in 1 L of ultrapure water and adjusted with hydrochloric acid and sodium chloride to achieve a pH of 7.4 and an osmotic pressure of 290 mOsm kg<sup>−1</sup>.

**Electrospin Patch Manufacture.** Electrospin mucoadhesive patches were produced as described by Santocildes et al., with modifications. Briefly, the electrospin dope consisted of 10% (w/w) PVP, 12.5% (w/w) Eudragit RS100, and 10% (w/w) PEO (400 kDa) and amounts of lidocaine base or lidocaine HCl theoretically resulting in a 3% (w/w) drug loading in the electrospin patch (i.e., 0.1 g of either lidocaine base or lidocaine HCl added to 0.326 g of polymers in the dope solution). In the case of lidocaine HCl, the 3% (w/w) loading includes the salt content. Polymers and drug were dissolved in 97% (w/w) ethanol prepared in deionized water, and solutions mixed at room temperature using a magnetic stirrer for 24 h or until they had dissolved. The electrospinning rig was configured with the following parameters: 19 kV voltage, 14 cm tip-to-collector distance, and 1.5 mL h<sup>−1</sup> flow rate. Additionally, 10% (w/w) PCL dissolved in DCM/DMF in a ratio of 93:7 (w/w) was electrospun using the same conditions and used as a backing layer. The dual-layer patch was formed by melting the PCL backing layer onto the drug-containing mucoadhesive layer by placing the layers between two glass slides followed by incubation at 70 °C for 5 min.

**Characterization of Electrospin Patches.** The morphology of the different electrospin patches was analyzed using scanning electron microscopy (SEM). The samples were gold-sputtered and imaged using Tescan Vega3 LMU SEM for 5 min. The fiber diameters were subsequently measured from the SEM images for the non-drug-loaded and drug-loaded patches (n = 3 patches for each condition, ~35 fiber measurements per image) using the Image<sup>+</sup> software (National Institute of Health). Differential scanning calorimetry (DSC) was applied using a Pyris 1 calorimeter (PerkinElmer) to determine the temperature transitions of the drug- and non-drug-containing electrospin patches. Electrospin patches (5--9 mg), lidocaine HCl or lidocaine base powders (~3 mg) were
weighed and placed in 50 μL aluminum pans, and aluminum lids crimped on top. Patches or powders were heated from 25 to 200 °C at 10 °C min⁻¹. The pH was measured after fully dissolving the different electrospun patches (n = 3 patches for each condition, 110 ± 4 mg; mean ± standard deviation (SD)) in 10 mL of deionized water using a FiveGo pH meter (Mettler Toledo, Switzerland) at 20.5 °C.

**Quantification of Lidocaine.** Reverse phase high performance liquid chromatography (RP-HPLC) with UV detection was performed to detect the presence of lidocaine using a XBridge BEH-C18 column (4.6 mm × 250 mm; 130 Å pore size) (Waters, U.K.) and a mobile phase composed of ACN/water (1:1, v/v) containing 0.1% (v/v) 7.5 M ammonia, with a flow rate of 1 mL min⁻¹, wavelength 262 nm, a 20 μL injection volume, and at room temperature. The retention time of lidocaine was 10.1 min, and the method was validated in terms of linearity and precision using lidocaine HCl in PBS standards of known concentrations. Good linearity (R² = 1.00) was determined with repeated injections of lidocaine HCl in PBS ranging from 39 to 1250 μM, where previous lower-range concentration injections determined 1.8 μM as the limit of quantitation (10 times the background reading created by PBS in the lidocaine peak region).

**Dosage Uniformity of Lidocaine in the Electrospun Patches.** Electrospun patches containing lidocaine HCl or lidocaine base were weighed, dissolved in 1 mL of 99.8% ethanol with stirring, and analyzed by UV RP-HPLC. Electrospun samples (n = 3) were taken from three separately spun mats to determine batch variation, where lidocaine HCl patches weighed 6.2 ± 1.2 mg and lidocaine base patches weighed 9.3 ± 3.3 mg.

**Release of Lidocaine from the Electrospun Patches in the Buffer.** Electrospun patches (Ø 10 mm) containing lidocaine HCl or lidocaine base were weighed and placed in a 12-well plate containing 2 mL of PBS. Samples of 200 μL were withdrawn every 5 min for 1 h, with the solution being replaced with a fresh buffer each time. The amount of lidocaine released was quantified using the RP-HPLC method previously described.

**Tissue Preparation.** Ex vivo porcine buccal mucosa was obtained from healthy experimental control pigs (approximately 30 kg of Danish Landrace/Yorkshire × Durox (D-LY)) (Department of Experimental Medicine, University of Copenhagen) and stored at −80 °C in cryomedia (40% (w/v) glycerol, 20% (w/v) sucrose in PBS at pH 7.4 and 290 mOsm kg⁻¹). The mucosa was thawed and washed by removal from the cryomedia into 4 °C PBS positioned on a 4 °C rotating plate and transferred to fresh PBS four times with 10 min incubations. The submucosa was trimmed to a thickness of 1011 ± 133 μm (mean ± SD, n = 10) and mounted onto an Ussing slider (P2413, Physiologic Instruments Inc.) with an exposed area of 0.71 cm².

**Buccal Mucosa Permeability Study.** Permeability studies were performed using modular EM-CSY-8 modified Ussing chambers (Physiologic Instruments Inc.). Receptor chambers were filled with 2 mL of PBS (pH 7.4) for all experiments. Donor chambers either contained 2 mL PBS with 3% (w/v) lidocaine HCl (pH 6.8) or were left empty for lidocaine-containing electrospun patch experiments. Patches were punched from the dual-layer electrospun mats (Ø 10 mm), and the drug-containing side of the patch was placed on the buccal epithelium and held in place with Parafilm M to assure direct contact of the patch with the tissue for the duration of the experiment (Figure 1). The dual-layer patches containing lidocaine HCl weighed 15.6 ± 2.3 mg (mean ± SD, n = 10), where the backing membrane PCL layer weighed 3.2 ± 0.14 mg (mean ± SD, n = 3). Sink conditions were maintained in the receptor chambers. A slight temperature increase over the 5 h period was observed (~37.0 to ~38.4 °C).

Aliquots of 200 μL were taken from the receptor chamber every 15−30 min over a 5 h period and were replaced with fresh PBS at each time point. The lidocaine concentration of each sample was determined using UV-HPLC. The accumulated permeated amount (JₜSS, μg min⁻¹ cm⁻²) was plotted against time (t, min), and the steady-state flux (dQ/dt, μg min⁻¹), where Q (μg) is the drug concentration, was determined by taking the linear part of the slope between 180 and 300 min using the eq 1 for steady-state flux (JₜSS), where A is the diffusion area (0.71 cm²)

\[
J_{SS} = \frac{dQ}{dt} \times \frac{1}{A}
\]

(1)

For lidocaine HCl in PBS, the apparent permeability (Pₜapp, cm min⁻¹) was calculated using eq 2

\[
P_{app} = \frac{J_{SS}}{C_{donor}}
\]

(2)

where the donor concentration was C_{donor} (μg L⁻¹).

**Spatial Location of Lidocaine in Buccal Mucosal Using MALDI-MSI.** Buccal mucosa was thawed, cut to a thickness of approximately 5 mm and mounted onto Ussing sliders. Three different experimental conditions were imposed on mounted tissues: (1) 70 μL of 0.3% (w/v) lidocaine HCl in PBS directly pipetted onto the epithelium; (2) an electrospun patch containing lidocaine HCl applied to the epithelium and held in place using Parafilm M; (3) an electrospun patch containing lidocaine base applied to the epithelium and held in place using Parafilm M. Three tissues for each condition were prepared, and the sliders were placed in a high humidity chamber filled at the bottom with PBS to cover the connective tissue side and incubated at 37.4 °C. Tissues were removed from the chamber after 15 min, 1 and 3 h. Samples were immediately embedded in 5% (w/v) CMC and stored at −80 °C. Frozen tissue samples were mounted in a Leica CM 3050 S Cryostat (Leica Microsystems A/S, Germany), cut vertically at −20 °C into 30 μm thick cross-sections, thaw-mounted onto glass microscopic slides that were stored at −80 °C. Sections
were removed from −80 °C and thawed using a vacuum desiccator for 5–10 min, then 300 μL of the matrix solution (30 mg mL⁻¹ of 2,5-dihydroxybenzoic acid (DHB) in methanol/water (1:1, v/v) with 1% (v/v) TFA) was sprayed onto the sample at a flow rate of 30 μL min⁻¹ with a nebulized gas pressure of 2 bar while the sample was rotating at 550 rpm, approximately 10 cm from the syringe tip. An AP-SMALDI10 Ion source (TransMIT Gesellschaft für Technologietransfer GmbH, Germany) mounted on a QExactive Orbitrap mass spectrometer (Thermo Scientific GmbH, Germany) was used for MALDI imaging. Imaging was performed in the positive ion mode using a scan range of m/z 200–800 and mass resolving power of 140 000@m/z 200. A peak from the DHB matrix at m/z-value of 295.02131 was used as a lock mass, thereby ensuring a mass accuracy of at least 2 ppm throughout the entire image. The images were acquired with a pixel size of 20 μm, and no oversampling took place (no overlapping between adjacent ablation craters was confirmed by reflected light microscopy of the samples subsequently to the MALDI imaging experiment). The raw data were converted to imzML files, and the MALDI images were generated in a MSiReader v. 0.0926 using a bin width of 0.002 Th.

**Histological Staining.** Post-MSI, the DHB matrix was removed by washing with ethanol and tissue sections stained with hematoxylin and eosin as previously described, then imaged using an Olympus BH-2 microscope (Olympus, Japan) equipped with an AxioCam ERC5s camera (Zeiss, Germany).

**Lidocaine Functional Assay.** The human neuroblastoma cell line SH-SY5Y was cultured in 1:1 (v/v) DMEM/Ham’s F12, supplemented with 10% fetal calf serum, 2 mM L-glutamine, 1% nonessential amino acids, 100 IU mL⁻¹ of penicillin, and 100 μg mL⁻¹ of streptomycin and incubated at 37 °C, 5% CO₂ in a humidified environment. Prior to experiments, cells were loaded with the fluorescent calcium indicator, Fluo-4 Direct following the manufacturer’s instructions. Briefly, cells were removed from tissue culture flasks using EDTA, centrifuged, and resuspended at 10⁶ cells/mL in a Fluo-4 Direct calcium buffer for 30 min at 37 °C. To test lidocaine bioactivity, lidocaine HCl-loaded or placebo electrospun patches (containing no lidocaine) were cut to 2 × 2 cm² and fully dissolved in 2 mL of PBS for 30 min, filtered-sterilized, then the solution added to Fluo-4 Direct-loaded cells to give a final concentration of 0.5 mM lidocaine or placebo control, and the cells incubated for a further 30 min at room temperature. To determine cell calcium responses, fluorescence was measured at 488 nm excitation and 530/30 nm emission using a FACSCalibur (BD Biosciences). For each sample, baseline fluorescence was measured for 40 s, then veratridine or vehicle control DMSO was added, and the fluorescence response measured for approximately a further 160 s. Relative fluorescence units (RFU) were calculated by subtracting the baseline median fluorescence intensity from the maximal median fluorescence intensity following stimulation with veratridine.

### RESULTS

**Manufacture of the Electrospun Patches.** Mucoadhesive electrospun patches containing lidocaine or lidocaine HCl were produced by modification of our published protocol. SEM was used to determine if the inclusion of lidocaine affected the structure or morphology of the patches (Figure 2). Patches of three different conditions were manufactured; patches containing lidocaine HCl or lidocaine base along with control patches without the drug. SEM images show that there was no significant difference in the fiber diameter or fiber structure between drug-containing and drug-free patches (Figure 2). All patches showed a similar range of fiber diameters with a random alignment, where the drug-free fiber diameters ranged from 2.28 ± 1.35 μm, the lidocaine HCl-loaded fibers from 1.98 ± 1.50 μm, and the lidocaine base-loaded fibers from 2.42 ± 2.09 μm (~100 fibers in total measured from n = 3 patches; mean ± SD). As two different drug compositions are incorporated into the patches and drug release can be dependent on the pH microenvironment of a system, the pH of the patches dissolved in deionized water was measured. It was found that the pH of the non-drug-containing patches in water was 7.78 ± 0.03, the pH of lidocaine HCl-containing patches in water was 7.25 ± 0.04, and the pH for the lidocaine base-containing patches in water was 8.26 ± 0.13 (n = 3; mean ± SD).

**Lidocaine Content of the Electrospun Patches.** Lidocaine was added to the polymer dope to achieve a drug loading of 3% (w/w). As low-molecular-weight species can be lost during spinning, the content of lidocaine in the electrospun patches in water was 8.26 ± 0.13 μg μm⁻¹ fibers (Figure 3). That is, the data showed the absence of a peak at 81 °C in the lidocaine HCl-containing patch, where lidocaine HCl powder has its melting peak, and the absence of a peak at 70 °C, for the lidocaine base-containing patch, the melting peak of the lidocaine base powder.

**Release of Lidocaine from the Electrospun Patches into the Buffer.** The mean accumulative release of lidocaine HCl and lidocaine base from four electrospun patches for each condition over 1 h is shown in Figure 4, with respective patch
Figure 3. DSC heating profiles of (A) an empty pan, (B) lidocaine HCl powder, (C) lidocaine base powder, (D) PVP/RS100/PEO electrospun fibers, (E) PVP/RS100/PEO electrospun fibers containing lidocaine HCl, and (F) PVP/RS100/PEO electrospun fibers containing lidocaine base. The onset melting temperatures of the upward-facing enthalpy peaks are given to the nearest 0.1 °C. The image is representative of n = 2 independent experiments.

Figure 4. Accumulative release of lidocaine from the lidocaine HCl or lidocaine base-containing electrospun patches in PBS over 1 h, shown as the drug released in mg per 10 mg of the patch. Data presented is the mean ± SD (n = 4).

weights of 10.8 ± 3.8 and 11.20 ± 2.7 mg (mean ± SD). As there was variation in the patch weight and variation in drug loading, the drug release per 10 mg of the patch was calculated. Lidocaine release from the lidocaine HCl patches increased rapidly to 0.16 ± 0.04 mg (mean ± SD) over the first 15 min, then gradually at 0.21 ± 0.02 mg up to 60 min. In contrast, the release of lidocaine base was slower, 0.06 ± 0.01 mg within 15 min, gradually increasing to 0.11 ± 0.02 mg after 1 h (Figure 4). Overall, the release of lidocaine from the lidocaine HCl patches was significantly quicker than that of lidocaine base from electrospun patches (p < 0.0001; Mann-Whitney U test).

Permeation of Lidocaine HCl through ex vivo Porcine Buccal Mucosa. Dual-layer electrospun patches were used for buccal mucosal permeation studies ex vivo to ensure the unidirectional release of lidocaine directly into the mucosal tissue, thereby preventing release into the donor chamber. Accumulative permeation of lidocaine from patches containing between 0.20 and 0.35 mg of lidocaine HCl per patch was linear over a period of 5 h (Figure 5), resulting in a flux of 0.268 ± 0.009 μg cm⁻² min⁻¹ (mean ± SD; n = 6). In comparison, the application of 3% (w/v) lidocaine HCl in the solution resulted in a flux of 4.092 ± 0.062 μg cm⁻² min⁻¹ and P_app value of 136.4 ± 0.002 (×10⁻³) cm² min⁻¹ (mean ± SD; n = 4) (Figure S1). However, no lidocaine could be detected in the receptor buffer upon application of electrospun patches containing 0.25–0.39 mg of lidocaine base per patch.

Spatial Distribution of Lidocaine HCl and Lidocaine Base Released from Electrospun Patches Applied to ex vivo Porcine Buccal Mucosa using MALDI-MSI. Time-dependent drug penetration and localization using MALDI-MSI was first investigated by exposure to a solution of lidocaine HCl (3 mg mL⁻¹, approximately 0.21 mg applied) for up to 3 h (Figure 6). The control image with no lidocaine applied to the tissue shows that the lidocaine m/z signal is not present in the tissue. After 15 min, lidocaine was located exclusively in the stratified epithelium, where it was evenly distributed and extending into the rete ridges. Drug penetration did not progress beyond the basement membrane. In contrast, after 1 h, lidocaine was concentrated throughout the entire epithelium and was observed progressing into the lamina propria (Figure 6), and by 3 h, lidocaine was found evenly distributed throughout the epithelium, lamina propria, and submucosa (Figure 6). The mass spectrum averaged over 100 pixels from this 3 h time point (Figure 6) is given in Figure S2, to show that the lidocaine signal was best detected in the protonated form at m/z 235.1807 with a mass accuracy of 1.0 ppm.

Lidocaine released from the lidocaine HCl patches was visible in the stratified epithelium and extending into the lamina propria after just 15 min (Figure 7). After 1 and 3 h, lidocaine was distributed throughout the epithelium and lamina propria (Figure 7). The mucosal distribution of lidocaine was more widespread and homogeneous when lidocaine HCl was applied as a solution compared to the electrospun patches (Figure 6). However, unlike administration of a solution in the oral cavity, the electrospun patches maintained local contact to the tissue where lidocaine was not depleted from the patch over a period of 3 h (Figure 7). Similar spatial distribution of lidocaine in tissue was observed from lidocaine base-containing patches applied to the ex vivo buccal tissue (Figure S3).

Lidocaine Released from Electrospun Patches Blocks Voltage-Gated Sodium Channels and Prevents Veratridine-Induced Calcium Responses in SH-SYSY Neuroblastoma Cells. Veratridine, an alkaloid toxin found in Liliaceae plants, causes the persistent opening of voltage-gated sodium channels leading to cell depolarization and downstream intracellular calcium flux. Lidocaine has been shown to exert its biological action by blocking these voltage-gated sodium channels. To test whether the lidocaine released from electrospun patches remained functional, we examined its ability to block endogenously expressed voltage-gated sodium channels by examining intracellular calcium flux following cell stimulation with veratridine. Untreated SH-SYSY cells displayed increased fluorescence associated with intracellular calcium flux when stimulated with veratridine (17.9 ± 2.8 RFU over baseline; Figure 8A). Similar data were observed when SH-SYSY cells were pretreated with the elutant from the placebo electrospun patches that contained no lidocaine (19.4
± 3.0 RFU over baseline, Figure 8B). In contrast, intracellular calcium flux was significantly reduced (11.3 ± 4.8 RFU over baseline; p ≤ 0.05) compared to the placebo patch when cells were preincubated with the eluant from lidocaine HCl electrospun patches (Figure 8C,D), showing that the lidocaine released from the electrospun patches was functional and able to inhibit veratridine-mediated opening of voltage-gated sodium channels. The injection of DMSO alone as control did not induce a calcium flux fluorescence signal, whereas preincubation of cells with a solution of lidocaine HCl (10 mM) inhibited the veratridine-mediated calcium influx by 88% (2.2 ± 1.5 RFU over baseline; p ≤ 0.05).

**DISCUSSION**

Painful oral mucosal conditions and dental pain can greatly affect the quality of life. For dental treatments, injections are commonly used; however, this causes dental anxiety in many patients. A number of topical delivery methods of pain relief are available, although many are inadequate for targeting the affected local tissue and delivering a sustained drug release, where mucoadhesiveness and direct contact of the delivery vehicle to the local tissue are an underlying issue. We have
previously developed a mucoadhesive electrospun oral drug delivery patch to resolve these issues and have used these patches to deliver the corticosteroid clobetasol-17-propionate to the mucosal tissue.\textsuperscript{11,12} The incorporation of lidocaine into electrospun polymers for transbuccal delivery has not previously been performed, although Palo et al. incorporated lidocaine HCl into an ink-jet-printed oral patch in a combined electrospun additive manufacturing process,\textsuperscript{35} thus not directly comparable to the patches used in this study. The data reported here confirms the ability of applying electrospraying to manufacture patches consisting of microscale polymer fibers loaded with lidocaine HCl and lidocaine base during fabrication. The incorporation of lidocaine HCl or base did not appear to influence fiber morphology, consistent with previous reports for unloaded and drug-loaded fibers.\textsuperscript{15,36} DSC analysis showed that both forms of lidocaine were amorphous when incorporated into electrospun fibers. Similar observations have been reported for many other drug compounds where the amorphous form has been shown to aid drug solubility and bioavailability.\textsuperscript{17–39}

Following the manufacture of lidocaine-loaded patches, drug release studies were performed to compare release rates for lidocaine HCl or lidocaine base patches (Figure 4). Significantly greater lidocaine release was observed from lidocaine HCl-containing patches. Around 80\% of lidocaine was released from the electrospun fibers in 1 h, which is similar to the previously reported release of lidocaine from poly-L-lactic acid electrospun fibers containing lidocaine HCl.\textsuperscript{40} Lidocaine base has been previously electrospun into CMC/PEO fibers,\textsuperscript{31} where release in the 1 h was close to 50\%, also agreeing with the findings in this study. One reason for the difference in release between the two drug compositions may be the surface pH microenvironment of the electrospun patches and the difference in acidity between the encapsulated lidocaine HCl and base. The acidic lidocaine HCl protonates and releases more quickly than that of lidocaine base, as shown in this study. The solubility of weakly basic drugs increases in lower pH environments, and hence acidity modifiers within the delivery vehicle may be used to increase their solubility.\textsuperscript{42}

The lidocaine-loaded electrospun patches were subsequently investigated for their ability to deliver lidocaine to the porcine buccal mucosa. For this experiment, cryopreserved frozen porcine buccal mucosa was used, as previous studies have shown no change in the permeability barrier functions between fresh and frozen mucosal tissues.\textsuperscript{7,24} The permeation profile of lidocaine across porcine buccal mucosa released from a lidocaine HCl-containing patch had a similar linear profile with a slight time lag as observed for the lidocaine permeating esophageal porcine tissue and rabbit ear skin\textsuperscript{13,48} after release from a lidocaine HCl-containing polyvinyl alcohol film. The lidocaine apparent permeability coefficient in solution to cross porcine buccal epithelium has been previously reported as 17.0 ± 1.8 (×10\(^{-6}\)) cm s\(^{-1}\), which is 7 times greater than shown in this study.\textsuperscript{17} However, here we have used a thicker section of mucosal tissue containing both epithelium and lamina propria compared to that study by Kokate et al. In the present study, lidocaine released from lidocaine base patches was not detected in the receptor chamber, suggesting that lidocaine release from the electrospun fibers was too slow, and/or the amount of permeated lidocaine through the tissue was too low to be detected by RP-HPLC. Lidocaine base has previously been reported to permeate the buccal tissue when released from a polymeric thin film,\textsuperscript{45} likely because the film contained approximately 100 times more lidocaine than in this study. Notably, the permeation of lidocaine observed in the study by Cavallari et al. was surprisingly quick as the authors were able to detect the drug in the receptor chamber after only 5 min, and permeation was shown to be at a similar rate to its release rate in PBS despite the tissue thickness being reported to be 1.3 cm.\textsuperscript{45}

Although ex vivo permeation studies provide valuable information on the rate of permeation, these experiments do not provide information on the spatial localization of drugs within the tissue over time. MSI is a powerful tool for high-resolution detection of substances in tissue sections, and a number of MSI techniques have been used to detect the presence of lidocaine, and metabolites thereof, in the skin.\textsuperscript{46–48} Here, we show for the first time the spatial distribution of lidocaine within porcine buccal mucosa following delivery from a mucoadhesive electrospun patch. Our data shows that lidocaine penetration into the oral tissue is most pronounced in the region of the patch placement, especially in the first 15 min, where lidocaine was mainly present in the stratified epithelium directly adjacent to the patch. For the lidocaine HCl-containing patches, the lidocaine signal was most intense within the electrospun patch compared to the tissue, especially for 15 min and 1 h. When associating this to the lidocaine permeation data where only 20 μg of potentially up to 300 μg of lidocaine contained in the patch had permeated through the mucosa after 1 h, it is understandable that the signal in the patch is much stronger. Once released from the patch, lidocaine, being a lipophilic molecule, permeated through the epithelium and into the lamina propria over time that was consistent with the homogeneous time-dependent tissue distribution displayed by lidocaine HCl in the solution. The tissue penetration depth may also be dose-dependent, as has been shown previously following the application of lidocaine to the skin.\textsuperscript{47} MALDI-MSI also allowed us to review the localization of lidocaine released from the lidocaine base-containing patches (Figure S3). Although lidocaine permeation was below the detection limit by HPLC, we found that lidocaine was localized throughout the epithelium and in the lamina propria after just 15 min exposure to the patch.

Although previous studies have shown the release of lidocaine from a number of fabricated films, none has shown that the drug retains its functional biological activity once released. Lidocaine exerts its anesthetic activity by blocking several voltage-gated sodium channels expressed by neuronal cells.\textsuperscript{32} Here, we show that lidocaine released from electrospun patches was able to interact with voltage-gated sodium channels expressed by SH-SYSY neuroblastoma cells preventing their opening upon stimulation veratridine.\textsuperscript{32} These data give clear evidence that levels of lidocaine released from patches directly into tissue are therapeutically active, enabling blockage of oral mucosal neuronal cell depolarization.

\section*{CONCLUSIONS}

Electrospun patches containing lidocaine HCL and lidocaine base were successfully manufactured by electrospinning, and the release of biologically active lidocaine from both types of the patch was demonstrated. Detailed investigations of diffusion through ex vivo porcine buccal mucosa suggested that lidocaine permeated into tissues following the release from a patch. This was confirmed by MALDI-MSI where, for the first time, the distribution of electrospun-delivered lidocaine within oral soft tissue was visualized. These data provide strong...
evidence that electrospun patches have potential as a delivery vehicle for lidocaine into buccal mucosa. Further clinical investigation is required to determine the feasibility of using this patch system as a topical dental anesthetic or as an analgesic for patients with painful oral mucosal disease conditions.

■ ASSOCIATED CONTENT

5 Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.molpharmaceut.9b00535.

Accumulative permeation of lidocaine HCl through porcine buccal mucosa and additional mass spectrometry data (PDF)

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Notes

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■ REFERENCES


