Specialized Pro-resolving Mediators Reduce Pro-nociceptive Inflammatory Mediator Production in Models of Localized Provoked Vulvodynia


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Abstract: Localized provoked vulvodynia (LPV) is the most common cause of chronic dyspareunia in premenopausal women, characterized by pain with light touch to the vulvar vestibule surrounding the vaginal opening. The devastating impact of LPV includes sexual dysfunction, infertility, depression, and even suicide. Yet, its etiology is unclear. No effective medical therapy exists; surgical removal of the painful vestibule is the last resort. In LPV, the vestibule expresses a unique inflammatory profile with elevated levels of pro-nociceptive proinflammatory mediators prostaglandin E2 (PGE2) and interleukin-6 (IL-6), which are linked to lower mechanical sensitivity thresholds. Specialized pro-resolving mediators (SPMs), lipids produced endogenously within the body, hold promise as an LPV treatment by resolving inflammation without impairing host defense. Ten of 13 commercially available SPMs reduced IL-6 and PGE2 production by vulvar fibroblasts, administered either before or after inflammatory stimulation. Using a murine vulvar pain model, coupling proinflammatory mediator quantification with mechanical sensitivity threshold determination, topical treatment with the SPM, maresin 1, decreased sensitivity and suppressed PGE2 levels. Docosahexaenoic acid, a precursor of maresin 1, was also effective in reducing PGE2 in vulvar fibroblasts and rapidly restored mouse sensitivity thresholds. Overall, SPMs and their precursors may be a safe and efficacious for LPV.

Perspective: Vulvodynia, like many pain conditions, is difficult to treat because disease origins are incompletely understood. Here, we applied our knowledge of more recently discovered vulvodynia disease mechanisms to screen novel therapeutics. We identified several specialized pro-resolving mediators as likely potent and safe for treating LPV with potential for broader application.

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Key Words: Specialized pro-resolving mediators, vulvodynia, pain, inflammation, polyunsaturated fatty acids.
tampoon, or even walk,
resulting in a substantial decline in quality of life.

There are no prescribed medical therapies for vulvodynia that outperform placebo. The only reliable therapy to eliminate pain is surgical removal of the vestibule following unsuccessful trials of less invasive therapies. A clinical trial has demonstrated that this perineoplasty procedure is more effective than behavioral therapy. However, it is considered by many to be disfiguring and comes with significant risks.

A key reason current vulvodynia therapies fall short is that the origins of the disease are poorly understood; vulvodynia is defined as unexplained vulvar pain lasting for ≥3 months. To address this unmet need, our research team has embarked upon mechanism-based drug development for LPV. We have implemented 2 models to study the underlying mechanisms of LPV: 1) an in vitro vulvodynia model utilizing fibroblasts isolated from the anatomic site of LPV pain and 2) a new in vivo mouse vulvodynia model described herein. Compelling evidence, including our work and others, establishes a link between inflammation and vulvodynia.

Fibroblasts from painful areas are inherently sensitive to inflammatory stimuli and will respond to stimuli that normally do not elicit a response, such as the resident microbial flora. In LPV patients, vestibular fibroblasts produce more pro-nociceptive interleukin-6 (IL-6) and prostaglandin E2 (PGE2) than fibroblasts from non-painful vulvar or regionally sampled fibroblasts from women without disease, which is linked to lower pain thresholds. Fibroblasts play an important role in the immune response, maintain their phenotypes in culture, and produce proinflammatory mediators, which makes them a useful model for investigating what is unique about the vestibule (site of pain) in LPV patients.

A mechanistic approach to LPV therapeutic development might therefore target inflammation and could be extended to other painful chronic diseases, including fibromyalgia. In vestibular fibroblasts, inhibiting nuclear kappa factor B (NFkB), a master regulator of inflammation, ablates proinflammatory signaling. However, even local inhibition of NFkB risks impairing inflammatory defenses at a site that must respond to infectious challenge. Topical steroids have been used to treat other vulvar conditions, such as lichen sclerosus, but they pose the same risks as NFkB inhibitors and have not shown efficacy for vulvodynia.

Inflammation is a dynamic process modulated by naturally derived products, referred to as specialized pro-resolving mediators (SPMs). All SPM classes (lipoxins, resolvins, protectins, and maresins) have been shown to impart analgesic effects in rodent pain models, including models of acute and chronic inflammatory, neuropathic, postoperative, and cancer pain. Furthermore, SPMs have virtually no toxicity, and several are in clinical trials for other indications. SPMs are not traditional anti-inflammatory agents and are not immunosuppressive; they are a critical component of the resolution machinery, deficits of which are associated with chronic pain.

Therefore, we investigated whether SPMs have efficacy for vulvodynia treatment. In addition to testing SPMs in our human fibroblast in vitro model, we optimized a mouse model that mimics key features of vulvodynia. We found that SPMs, and even the polyunsaturated fatty acids from which they are derived, are effective in reducing pro-nociceptive IL-6 and PGE2 levels and restoring pain thresholds to baseline levels in mice.

Methods

Patient/Sample Selection

LPV-afflicted cases (fulfilling Friedrich’s Criteria) and age/race-matched pain-free controls were recruited from the Division of General Obstetrics and Gynecology clinical practice at the University of Rochester between December 2012 and November 2019. All subjects provided informed consent, and the research was approved by the University of Rochester Institutional Review Board (RSRB #42136). Expanded details on our selection criteria and sampling procedures have been previously published.

In brief, all subjects denied the use of corticosteroids and nonsteroidal anti-inflammatory medications and had no chronic inflammatory illnesses other than LPV. Prior to biopsy of the vestibular and external vulvar sites, sampling sites underwent Wagner mechanical algometry. We used a Method of Limits technique for vulvodynia mechanical pain threshold initially described by Zolnoun et al and replicated in our earlier publication. Using the Wagner algometer, an increasing 0.5 N per second force (range 0–5 N) was applied perpendicular to the mucocutaneous surface by a moistened dacron tipped swab affixed to the Wagner algometer. Force was terminated at the point of pain development (subject signaled by hand-held clicker) or when the mucocutaneous force reached 5 N. Algometer-site tissue was sampled and used to create fibroblast strains as previously described. A total of 2 existing paired (vulvar vestibule and external vulva) case and 2 paired control fibroblast strains (8 total) were used for this study. Key patient characteristics are listed in Table 1. These strains are representative examples of a much larger panel of cases and controls; they respond consistently and predictably from experiment to experiment.

Fibroblasts are a useful model for vulvodynia, not traditional anti-inflammatory agents and are not immunosuppressive; they are a critical component of the resolution machinery, deficits of which are associated with chronic pain.

Table 1. Patient Characteristics for Fibroblast Strains

<table>
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<th>STRAIN</th>
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<th>VESTIBULE THRESHOLD</th>
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<td>Control B</td>
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because they 1) participate in the immune response and secrete proinflammatory mediators into the media that can be readily quantified, 2) maintain their phenotype in culture, and 3) are viable over a number of passages without immortalization.18-21,26-28,58

**Fibroblast Culture**

Previously established primary fibroblast strains (each obtained from a different patient or healthy control) were cultured in Minimum Essential Medium (MEM) supplemented with 10% FBS, GlutaMAX, gentamicin, and antibiotic/antimycotic solution (Gibco/Invitrogen/Thermo Fisher Scientific, Grand Island, NY). Early passage (4-10) external vulvar and vestibular fibroblast strains were seeded at 2.5 x 10^4 cells/cm². After cultures reached full confluence, fibroblasts were typically serum-starved for 48 hours in MEM lacking FBS prior to stimulation with inflammatory agonists (e.g., IL-1β). Fibroblast cellular identity was previously confirmed by microscopic inspection and with fibroblast-specific markers (e.g., vimentin, collagen). At the same time, the cells were confirmed to be negative for epithelial cell markers (e.g., cytokeratin), smooth muscle and myofibroblast markers (e.g., α-smooth muscle actin), endothelial cell markers (e.g., CD34), and bone marrow derived cell markers (e.g., CD45).59 These markers remain constant across passages, and the cells consistently respond to stimuli throughout culture.

**Impact of SPM Pre- and Post-treatment on Proinflammatory Mediator Production in Vulvar Fibroblasts**

Cells were grown to confluence in 24-well plates then serum started for 48 hours prior to any subsequent treatments. Two representative paired case strains were used. All treatments were administered to quadruplicate wells. For SPM pretreatment (prior to the initiation of proinflammatory signaling), cells were treated with 5 nM RvD1, AT-RvD1, 17S-HDHA, RvD2, RvD3, RvD4, RvD5, RvE1, LXA4, LXB4, maresin 1, 7(S) epi-Maresin 1, protectin D1, or protectin DX for 18 hours (Cayman Chemical Company, Ann Arbor, MI). Cells received a final SPM dose 18 h later and supernatants were collected 48 hours after stimulation. Previous dose ranging experiments were conducted to determine 5 nM was the lowest effective dose of SPMs that significantly reduced IL-6 and PGE₂ production. For the few SPMs that did not reduce IL-6 and PGE₂ levels, 10 or even 100-fold higher concentrations were ineffective. Standard sandwich ELISAs were performed to measure production of IL-6 (BD Biosciences, Franklin Lakes, NJ) and competitive EIA assays were performed to measure PGE₂ production (Cayman Chemical Company, Ann Arbor, MI).

**3D Tissue Culture of Mouse Vulvar Biopsies**

Six mm punch biopsies were collected, encompassing the entire vulvar area immediately posterior to the vaginal opening, and each biopsy tissue was oriented to permit non-tangential cross-sectioning and was equally bisected to create 2 samples. Each biopsy half was washed several times in sterile saline prior to being placed in a 24-well plate. Twelve mice donated a total of 24 biopsy pieces, which were divided among treatments; 3 pieces from different individuals (n = 3) were used for each treatment. Each well, containing a single biopsy piece, was flooded with 0.5 mL serum-free MEM, supplemented with GlutaMAX, gentamicin, and antibiotic/antimycotic (Gibco). 3D cultures were pretreated with lipoxin A₄, resolin D₂, or maresin 1 at a concentration of 1, 10, or 100 nM for 24 hours. Each treatment was applied to three pieces of biopsy tissue. After 24 hours, cells were treated a second time with lipoxin A₄, resolin D₂, or maresin 1 30 minutes prior to treatment with 10 pg/mL murine IL-1β (R&D Systems) for another 24 hours before harvesting supernatants and assaying for PGE₂, as described earlier.

**Effect of Docosahexaenoic Acid Treatment on Vulvar Fibroblasts**

To study the proinflammatory signal modulating effects of DHA, cells were grown to confluence, then treated with 200 nM or 1 μM docosahexaenoic acid (DHA) (Cayman Chemical, Ann Arbor, MI) for 72 hours prior to stimulation with 10 pg/mL IL-1β and a second dose of DHA for another 72 hours. Two paired case and control strains were used. Supernatants were then assayed for PGE₂.

**Maresin and DHA Testing in Mice**

All procedures involving mice were approved by the University of Rochester Committee on Animal Resources (UCAR protocol 2016-006). Zymosan, a proinflammatory yeast cell wall preparation (Millipore Sigma, St. Louis, MO) was used to induce sustained vulvar allodynia, measured by pain threshold testing as described by Farmer et al.22 The Farmer model used live yeast infection or zymosan as the provoking stimuli to elicit vulvar allodynia; we elected to use zymosan because it is more highly reproducible and reduces the technical complexity of the model versus infection with a live pathogen. We initially used a manual von Frey that employs a series of “hairs” of different thicknesses/rigidity that exert differing forces when applied to the injection site.
located at the midline posterior vulva (between the vaginal opening and anus) and then transitioned to an electronic system. Details on model development can be found in the Supplementary Figs 1-4).

During Phase 1 of the experiment, 12 eight-week-old female C57BL/6 mice (Jackson Laboratories, Bar Harbor, ME) were acquired and trained to facilitate electronic von Frey determination of vulvar pain thresholds. During training, the mice were given access to a preferred drinking solution containing 3% dextrose and 0.125% saccharin (in tap water), which was filter sterilized before administration. The mice were given the solution in a standard water bottle (replacing their normal water supply) daily for ~6 h/day in their home cage for ~1 week until they began to steadily drink the solution, at which point they were transferred to the testing apparatus. Once transferred to the testing apparatus (Fig 1), they were given a steady drip (50 μL droplet/s) of solution for ~2 h/day, 2–3 days/week, for a period of ~2 weeks. Once the mice were able to find the solution in the test cage, the flow was tapered to one 50 μL droplet/30 seconds interval, and the mice were again exposed for ~2 h/day, 2–3 days/week, for a period of ~2 weeks to this noncontingent fixed time schedule of delivery. Once the mice held relatively still in the testing apparatus upon introduction to the cage, they began baseline mechanical sensitivity threshold testing. The mice were shaved and tails were inked weekly. Four colors were used to distinguish each mouse in a cage; the same 4 colors were used for each cage, so mice from different cages could not be distinguished. A series of 3 individual threshold tests were conducted to determine the baseline threshold before zymosan injections commenced.

For testing using the electronic von Frey, a cage of 4 mice was tested at once, alternating the mice stimulated, working from left to right and repeating the series until 5 values were collected for each mouse. The electronic von Frey device was gently rotated upwards at approximately 1 g/s, increasing the pressure, until the mouse stepped or jumped off the hair, at which point the peak force was automatically recorded (mechanical sensitivity threshold). The investigator was blinded to cage and mouse identity. Blinding was maintained by reading cage barcodes into custom LabVIEW software for use with the Mousemet electronic von Frey device (TopCat Metrology, Ltd., Cambridge, UK). The software reads the barcode and keeps it blinded from the investigator during testing, after which the data, retaining mouse identities, is exported to an Excel file.

The mice were permitted access to the preferred drinking solution throughout testing at a rate of one 50 μL droplet/30 seconds interval. The intertrial interval is at least 30 seconds to prevent acute desensitization from repetitive stimulation. The tester assays the validity of each test and records test values to each mouse based on test cage position using the button on the electronic von Frey device or the computer keyboard. To be a valid test, all the following must be true: 1) the force increased ~1 g/s across within an acceptable range of 0.5–4 g/s, 2) the hair contacted the injection site, 3) the hair did not leave the injection site prior to mouse response, and 4) the hair and device arm were not depressed by anything other than contact with the

Figure 1. Treatment scheme and testing environment. Panel A: Image of mouse vulva. Arrow indicates injection site. Below image, there is a schematic of in vivo mouse model to establish vulvar allodynia. Panel B: Images of allodynia testing environment. Four mice are placed in separate compartments that permit access to the vulva from below. The front side is masked to reduce visual cues during testing, while permitting imaging from the rear.
Based on threshold; injections ceased after a ≥33% drop in threshold (vs baseline threshold) for any 2 weeks. Four mice received saline and 8 mice received zymosan injection. The person conducting the pain threshold testing was blinded to treatment assignment. Mechanically records the peak value when pressure on the hair is relieved (eg, the mouse steps off the hair), eliminating the need for the tester to make a determination of if/when the mouse reacts to the stimulus.

During the induction phase, the mice received up to 6 weekly injections to the midline posterior vulvar under isoflurane anesthesia and underwent weekly threshold testing. The number of injections administered was based on threshold; injections ceased after a ≥33% drop in threshold (vs baseline threshold) for any 2 weeks. Four mice received saline and 8 mice received zymosan injection. The person conducting the pain threshold testing was blinded to treatment assignment. Mechanically records the peak value when pressure on the hair is relieved (eg, the mouse steps off the hair), eliminating the need for the tester to make a determination of if/when the mouse reacts to the stimulus.

For DHA testing, 36 eight-week-old female C57BL/6 mice (Jackson Laboratories, Bar Harbor, ME) were divided into 3 treatment groups. A topical formulation (cream) of DHA was prepared containing the following: 3% glyceryl monostearate, 3% cetyl alcohol, 2.5% polyoxy-40, 3.5% isopropyl myristate, 4% white petrolatum, 1% benzyl alcohol, 0.5% vitamin E acetate, 15% neosorb 70/20 B, 65.5% purified water, and 2% highly purified fish oil enriched for DHA (Omegatex, Solutex GC, SL, Madrid, Spain; ~70% DHA by volume). To prepare the cream, the oil and water phases were heated to 70°C and combined by high speed homogenization for 5 minutes, and then mixed while cooling. Active phase ingredients were pre-mixed with antioxidant vitamin E and preservative benzyl alcohol and added to the emulsion at 45°C. A vehicle cream containing all the elements, save DHA, was also prepared. Creams were prepared in a small batch under non-GMP conditions by the Ferndale Pharma Group (Detroit, MI). Mice were randomized into 3 groups: vehicle, DHA, and mock. Mice were held by the tail, and a pea-sized amount of cream was applied to the entire shaved area by rolling sterile cotton tipped swab over the area. For mock treatment, a saline-moistened swab was used. Threshold testing was performed blinded each week as described.

**Statistical Methods**

The following statistical programs were used: SAS 9.4 (SAS, Cary, NC), STATA (Stata Corp, LLC, College Station, TX), and Graph Pad Prism 7 (Graph Pad Software, San Diego, CA). Differences in IL-6 and PGE2 production in human vulvar fibroblasts in response to treatment with SPMs were found to be non-normally distributed by Shapiro-Wilk testing. We, therefore, utilized the non-parametric Wilcoxon Rank sum test. Based upon the testing of 10 SPM candidates, significance was Bonferroni corrected to a P < .006 of a two-tailed distribution. In cultured mouse tissue biopsies, differences in PGE2 were determined via one-way ANOVA. In live mice, differences in pain thresholds were analyzed with one-way ANOVA, while differences in PGE2 levels were analyzed via paired t-test. Spearman Rank correlation was performed to examine the relationship between PGE2 levels and threshold in mice. According to the Shapiro-Wilk testing, pain threshold values were found normally distributed but PGE2 values significantly differed from normality and therefore Spearman Rank was selected. Significance was defined at P ≤ .05 of a two-sided distribution.
**Results**

**SPM Treatment Reduces Human Vulvar Fibroblast Proinflammatory Signaling**

To assess whether SPMs reduce proinflammatory signaling linked to pain in vulvodynia, we evaluated the effects of all commercially available SPMs on IL-6 and PGE2 production in vulvar fibroblasts from 2 representative cases used in previous studies. We employed 2 treatment strategies: 1) a pretreatment strategy where cells were treated with SPMs prior to exposure to the inflammatory activator interleukin-1 beta (IL-1β) and 2) a post-treatment strategy where inflammatory signaling was initiated with IL-1β for 30 minutes prior to SPM treatment. We tested lipoxin A₄, lipoxin B₄, resolvin D₁, aspirin-triggered resolvin D₁, resolvin D₂, resolvin D₃, resolvin D₄, resolvin D₅, resolvin E₁, maresin 1, 7S-epi-maresin 1, protectin D₁, and protectin Dₓ. Ten of the 13 SPMs reduced IL-6 and/or PGE₂ levels, as a pretreatment and/or a post-treatment (Fig. 2). Resolvin D₁, aspirin-triggered D₁, and lipoxin B₄ were the only SPMs that were ineffective in reducing IL-6 or PGE₂ levels. The magnitude of reduction by pretreatment was generally greater than post-treatment, and more SPMs were able to significantly reduce IL-6 versus PGE₂ (Fig. 2). Nonetheless, most SPMs tested were capable of reducing proinflammatory mediator levels associated with vulvodynia pain.

To parse out the most significant differences, we plotted a concordance bubble chart (Fig. 3) depicting the concordance of the degree of reduction in both IL-6 and PGE₂ levels across all vulvar samples, including external vulvar and vestibular fibroblasts, both with pre- and post-treatment dosing. This strategy enabled the use of high stringency statistical methods to better rank the ability of each SPM to reduce proinflammatory mediator production to identify the most promising hits for further testing. The highest concordance was evident for lipoxin A₄, 7S-epi-maresin 1, and resolvin D₂, indicating they have the strongest suppressive effects for both IL-6 and PGE₂. The relative bubble size reflects specificity of the suppressive response for the vulvar vestibule (painful area), confirming resolvin D₂ and 7S-epi-maresin 1 are highly effective in reducing proinflammatory mediator release. Overall, exogenous SPM treatment has the capacity to reduce proinflammatory signaling associated with vulvar pain, while the maresins, lipoxin A₄, and resolvin D₂ were especially effective in reducing IL-6 and PGE₂ levels in human vulvar fibroblasts.

**SPM Treatment Reduces Proinflammatory Mediator Levels in Cultured Mouse Biopsy Tissues**

Prior to testing a response to SPM treatment in mice, we tested the effects of SPM treatment on PGE₂ production in a three-dimensional tissue culture model using mouse vulvar biopsies. Pretreatment with either maresin 1, lipoxin A₄, or resolvin D₂, SPMs that profoundly reduced both IL-6 and PGE₂ production in human vulvar fibroblasts (Fig. 2 and 3), also reduced PGE₂ release from mouse vulvar tissue (Fig. 4). All 3 SPMs produced dose-related effects over a 1 to 100 μM concentration. Maresin 1 was then selected for further testing in mice, because it was highly effective in reducing IL-6 and PGE₂ levels in both models, especially in the 3D mouse culture model. In addition, exogenous maresin 1 has been shown to reduce inflammatory signals, including IL-6 and alleviate neuropathic pain and allodynia in other mouse models. Reduced levels of endogenous maresin 1 may be even be linked to the development of chronic pain. These observations, coupled with our in vitro data, led us to further investigate the analgesic properties of maresin 1 in a mouse model of vulvar allodynia.

**Maresin 1 Increases Mechanical Sensitivity Thresholds in Mice**

Using the model of vulvar allodynia described by Farmer et al. as a starting point (Fig 1, Supplementary Fig 1), we developed a robust and reproducible model for testing the efficacy of potential topical SPM-based therapies for vulvodynia. Feasibility trials in mice demonstrated once daily treatment with maresin 1 can raise mechanical sensitivity thresholds and reduce PGE₂ levels (Supplementary Fig 2). Enhanced model development (Supplementary Fig 3) produced reproducible and consistent threshold measurements (Supplementary Fig 4).

Following the model development stage, we tested the efficacy of maresin 1 in increasing mechanical sensitivity thresholds. A reliable progressive reduction in mechanical sensitivity threshold values was produced by repeated zymosan injections (Fig. 5). Furthermore, reduced thresholds persisted through and beyond the induction period. After 4 injections, most mice developed allodynia that was sustained for several weeks, creating a window to evaluate analgesic effects. After 4 weeks of maresin 1 treatment, there was an increase in threshold, near or above baseline. Thresholds were increased above the baseline; threshold values increased over time for both maresin 1 and vehicle treated mice but threshold values were higher for maresin 1 treated mice during treatment (P < .05) (Fig. 5). However, a vehicle effect was evident; DMSO has analgesic properties that may contribute to the observed increase in thresholds. This increase is nonetheless enhanced by maresin 1, demonstrating maresin 1 has analgesic properties beyond the effects of DMSO, contributing to a faster and fuller recovery.

PGE₂ is highly conserved between mice and humans and is a surrogate marker for pain in women with LPV; high levels of PGE₂ in vitro predict reduced pain thresholds. Therefore, we also assayed vulvovaginal levels of PGE₂ in mice weekly. The pattern in PGE₂ levels was consistent with an acute inflammatory response preceding sustained allodynia. PGE₂ levels peaked after the fourth injection and decreased as acute inflammation resolved, at which point maresin 1 treatment did not further...
reduce these levels (Fig 5). Mice receiving zymosan had signs of erythema and edema during the induction phase, which resolved after the cessation of the injection series. Plotting average PGE2 levels against average threshold values showed a concomitant decrease in threshold value. Spearman Rank correlation demonstrated an inverse relationship between threshold values and PGE2 levels (rho = .56; P < .002), recapitulating what occurs in women with vulvodynia where high PGE2 levels (produced by fibroblasts in vitro) are associated with low threshold values.26 In this aspect, the mouse model recapitulates the human condition; repeated vulvar insults (eg, chronic yeast infection) produce acute inflammation that resolves over time but leaves a lasting hypersensitivity to normally non-painful stimuli. However, vulvodynia PGE2 levels were not reliable in distinguishing treatment effects.

Mice Do Not Exhibit Conditioned Behavior in Response to Von Frey Testing

To validate the specificity of threshold testing in C57BL/6 mice, we devised a method to evaluate behavioral conditioning to mechanical sensitivity threshold assessment. Because frequent touching of the vulva with a probe might result in avoidance behavior, a subset of mice received topical lidocaine/prilocaine prior to...
threshold testing. Without evidence of a conditioned response to the von Frey filament, lidocaine/prilocaine should minimize pain responsive behaviors.

C57BL/6 mice showed a significant increase in threshold values following application of the lidocaine/prilocaine solution (Fig 6). Baseline thresholds were $>3$g force and were reduced to $<1$ g after 6 weeks of zymosan injection, but restored to $>3$ g with lidocaine/prilocaine application. Baseline thresholds were indistinguishable from post-lidocaine thresholds. Therefore, avoidance conditioning did not occur or interfere with mechanical sensitivity threshold assessment.

Vulvar fibroblasts can use docosahexaenoic acid (DHA) as a substrate for SPM production, which reduces PGE2 levels in vitro

Natural products rich in omega-3 fatty acids (eg, fish oil) could represent a more efficient option for therapeutic development. Therefore, we sought to determine if human vulvar fibroblasts could utilize polyunsaturated fatty acid (PUFA) precursors to produce SPMs and if PUFA supplementation would be sufficient to reduce proinflammatory mediator levels. Preliminary analysis showed DHA, arachidonic acid (AA), and eicosapentaenoic acid (EPA) are substrates for the production of SPMs while the greatest number of and most abundant SPMs were DHA derived (Supplementary Fig 5). Therefore, we also examined the effects of DHA on proinflammatory mediator production. DHA treatment significantly reduced PGE2 levels dose responsively and was effective even at very low nanomolar concentrations (Fig 7). Taken together, these observations suggest DHA is effective in reducing proinflammatory signaling in vulvar fibroblasts through the production of SPMs.

DHA Increases Sensitivity Thresholds in Mice

We went on to test a topical formulation of DHA in mice using formulations comparable to other topical applications currently approved for human use. Mice were treated twice daily for a total of 7 weeks, at which point the majority of the mice in the vehicle and DHA treated groups had recovered (75%), while only half the mice recovered in the mock treated group (Fig 8A and 8B). Mice in the DHA group recovered fastest; 50% recovered after 2 weeks of treatment, while it took 4 weeks of vehicle treatment and 6 weeks of mock
treatment for 50% to recover. With no recovery in the mock treatment group in the first 4 weeks, these early affects can be attributed to DHA and to some extent the vehicle; a vehicle effect was apparent after 3 weeks. We used the threshold data to calculate the percent improvement score, which represents the percent increase in threshold over the last threshold prior to initiating treatment (Fig 8C). The average percent improvement in mechanical sensitivity threshold was significantly lower in the mock and vehicle treated groups versus the DHA treated group over the first 4 weeks.

**Figure 5.** Increased PGE$_2$ levels are accompanied by a lower mechanical sensitivity threshold; maresin 1 increases mechanical sensitivity thresholds in C57BL/6 mice. This graph depicts the average mechanical sensitivity thresholds for all C57BL/6 mice that developed allodynia (box and whisker plots, showing maximum, minimum, and median values). Baseline thresholds were established prior to allodynia induction by averaging three separate testing sessions during which an average of 5 threshold measures were taken per session (gray bar). Allodynia was then induced through a series of up to 6 weekly zymosan injections (white bars). A final assessment was taken on week 7 (last white bar) before mice were treated with maresin 1 (red bars) or vehicle (yellow bars) for 4 consecutive weeks. The threshold values are plotted as the average of 5 threshold tests for each mouse, with the exception of the baseline (average of 3 tests or 15 values), while the PGE$_2$ levels are the average of the entire group with one weekly sampling per mouse (n = 7 mice). Over the induction period, pain thresholds declined and remained low, while PGE$_2$ levels increased and eventually tapered off as acute inflammation was resolved, consistent with visible inflammation during the induction period (erythema, edema), which resolved prior to initiating treatment. There was an inverse relationship between thresholds and PGE$_2$ levels during the induction phase (rho = 0.56; P < .002), similar to that seen in vulvodynia. During the treatment phase, PGE$_2$ levels remained low and thresholds increased. Thresholds were higher in maresin 1 treated mice (*P < .05 maresin vs vehicle), but thresholds also increased with vehicle. Overall, maresin 1 exhibited effects beyond the vehicle and enhanced recovery in mice.

**Figure 6.** Reduced mechanical sensitivity threshold and lidocaine reversal are not avoidance responses in C57BL/6 mice. Box and whisker plots with maximum, minimum, and median values show baseline touch sensitivity and sensitivity after zymosan treatment before and after lidocaine treatment for C57BL/6 mice. Data are based on 5 determinations in each of 6 mice (one-way ANOVA, *P < .05). Thresholds were reduced with zymosan and increased or were restored to baseline levels with lidocaine. Baseline thresholds were indistinguishable from post-lidocaine thresholds. Therefore, altered mechanical sensitivity threshold was not conditioned avoidance of contact.

**Figure 7.** DHA reduces proinflammatory mediator levels in human vulvar fibroblasts. Human vestibular and external vulvar fibroblasts from 2 case and 2 control strains were pretreated with DHA for 72 hours, then activated with IL-1$eta$ (10 pg/mL) for another 72 hours. Culture media were collected and analyzed for PGE$_2$. DHA reduced PGE$_2$ levels in the presence of IL-1$eta$, a strong inflammatory stimulus. Mean +/- SEM of n = 4. ANOVA, *P < .05 vs corresponding vehicle treatment.
Figure 8. Topical DHA treatment increases pain threshold and reduces time to recovery in C57BL/6 mice. Panel A depicts the average mechanical sensitivity thresholds for all C57BL/6 mice that developed allodynia (box and whisker plots, showing maximum, minimum, and median values). Mice were treated with DHA, vehicle, or mock treatment for 7 weeks after establishing allodynia (n = 11-12 mice/group). The starting baseline prior to allodynia induction and the last threshold prior to initiating treatment (week 0) are also plotted. At week 0, thresholds were significantly lower than at baseline, but increased over the course of treatment, approaching levels similar to baseline. ANOVA, *P < .05 vs mock treatment. Panel B shows mouse recovery over time with treatment, which takes into account individual improvements for each mouse. Mice were considered to have recovered if they showed at least a 70% improvement in their pain threshold for two consecutive weeks. Mice began to recover as early as week 2 in the DHA group. By week 4, >50% of the mice had recovered in the DHA and vehicle groups, while no mice had recovered in the mock treated group. Panel C shows the mean percent improvement (+/- SEM) over the last pre-treatment threshold throughout seven weeks of treatment, which compares each mouse’s threshold to their pretreatment threshold. The percent improvement in pain threshold was significantly higher in the DHA versus mock treated group at weeks 2, 3, 5, and 6, while vehicle was only significantly higher than mock after 6 weeks of treatment, Two-way ANOVA, *P < .05, n = 12 for vehicle and DHA, n = 11 for mock.
weeks of treatment, again demonstrating that mice treated with DHA recovered more quickly than vehicle or mock treated mice.

Discussion

Developing new therapies for pain is particularly difficult, because there is a paucity of mechanistic pain research in this area, and although there are quantifiable endpoints, the experience can vary from person to person and animal models present complex challenges.\(^\text{16,46,47,75,76}\) New therapies for chronic vulvar pain are direly needed; vulvodynia patients may undergo treatment for years before symptoms resolve.\(^\text{18,56}\) Often, this resolution is achieved by surgical removal of the vestibule, encompassing a substantial portion of the vulvar tissue.\(^\text{6,18,56,70}\)

The etiology of vulvodynia is poorly understood, and as a result the current therapies are not mechanism-based.\(^\text{18}\) Therefore, our work has focused on identifying mechanisms of disease to achieve targeted efficacious therapies. Our findings to date,\(^\text{18,21,26-28}\) along with the findings of other groups,\(^\text{41,42,68,71}\) implicate aberrant proinflammatory signaling in this mechanism. Here, we tested the plausibility of using SPMs as a therapy for vulvodynia; their roles in the resolution of inflammation suggest they might be ideal therapeutics for LPV. In addition, recent publications indicate SPMs may alleviate pain in other inflammatory conditions.\(^\text{2,25,43,60,66,78}\)

Consistent with our hypothesis, we found that most commercially available SPMs can reduce the levels of one or both proinflammatory mediators previously associated with vulvodynia pain,\(^\text{26}\) while a few SPMs have particularly profound effects. We came to focus on the maresins, because both 75-epi-maresin and maresin 1 were among the SPMs that had striking effects in reducing IL-6 and PGE\(_2\) levels in fibroblasts, as well as PGE\(_2\) levels in mouse vulvar tissue. In addition, we found that vulvar fibroblasts are capable of making SPMs, most of which are DHA-derived, among them maresin 1. DHA was also effective in reducing PGE\(_2\) levels in fibroblasts, which could be explained by an indirect effect involving the production of SPMs, namely maresins or D-series resolvins. The D-series resolvins detected via lipidomic analysis have no or weaker effects on PGE\(_2\), suggesting maresin 1 may be responsible for reductions in PGE\(_2\). Our in vitro fibroblast model represents an efficient way to prescreen potential therapeutic agents for vulvodynia, even beyond SPMs. Here, we identified 10 SPMs and one PUFA (DHA) that were highly effective in suppressing pro-nociceptive proinflammatory signaling, which could impart analgesic effects.

Proinflammatory cytokines have long been implicated in the elicitation of pain in numerous conditions.\(^\text{63}\) As denoted, IL-6 and PGE\(_2\) are surrogate measures of LPV pain, as measured in our in vitro fibroblast model.\(^\text{26}\) However, this model does not allow evaluation of the potential analgesic effects of SPMs. Studying putative therapeutics in in vitro and in vivo testing systems enhances the likelihood of success in subsequent clinical trials. To this end, we optimized a validated murine model to test new therapeutic agents for vulvodynia.\(^\text{22}\) This model recapitulates important clinical aspects of human vulvodynia, eliminates subjective evaluations of pain, permits randomization and blinding, and assesses mechanical sensitivity in mice as is done in human settings.\(^\text{18,26}\) A lone investigator can conduct blinded testing through the use of appropriate software that exploits barcode scanning for blinding. The software and systems developed here can be readily incorporated in hind paw models commonly used to study pain.\(^\text{15,46,47,54,80}\)

We incorporated both measures of sensitivity and the ability to track inflammatory end points. Although lavage fluid may not completely reflect the profile in the tissue, using vulvovaginal lavage fluid is noninvasive and can be repeated throughout the course of the experiment. We cannot be certain that repeated bouts of inflammation, such as weekly zymosan injection in the mouse model, are clinically linked to vulvodynia. However, several pieces of evidence suggest that inflammatory insults may elicit or at minimum contribute to vulvar allodynia. Greater than 70% of women with vulvodynia report a previous history of chronic or recurrent yeast infection,\(^\text{17}\) and at least some women with vulvodynia exhibit cutaneous hypersensitivity to yeast.\(^\text{51}\) The Farmer model established that repeated injection of zymosan or infection with live Candida albicans resulted in vulvar allodynia that remained after infection and inflammation were resolved, very similar to what is observed in women with LPV disease.\(^\text{22}\) In addition, fibroblasts from the vulvar vestibule are exquisitely sensitive to proinflammatory stimuli,\(^\text{18,21,26,28}\) there is an increased abundance of and altered organization of inflammatory cells in the painful vestibule of LPV patients,\(^\text{41,42,68,71}\) and there is a correlation between mechanical thresholds in patients and the production of proinflammatory mediators by fibroblasts cultured from these painful areas.\(^\text{26}\) Here, we demonstrate that agents that help to resolve inflammation also impart analgesic effects in mice with vulvar allodynia, suggesting inflammation is involved in the vulvar pain mechanism and is a suitable target for analgesic therapy.

Using this mouse model, we tested first the effects of maresin 1 and then DHA on vulvar sensitivity and found that both were effective in raising sensitivity thresholds reflective of increased tolerance of force and presumably reduced sensitivity. For maresin 1, we also assayed PGE\(_2\) levels in vulvovaginal lavage fluid, but did not find a significant treatment effect, although we did find a correlation between threshold and PGE\(_2\) levels, similar to what is observed in women with LPV. The absence of a treatment effect on PGE\(_2\) levels could reflect the differences in anatomical sampling, or the levels detected during the treatment phase may already be too low to further suppress. The primary outcome measure, change in mechanical sensitivity threshold, denotes significant drug effects. For both maresin 1 and DHA treatment, there was a vehicle effect, although mice receiving active treatment recovered more quickly and fully
than mice receiving vehicle. The DMSO in the maresin 1 vehicle and the long chain alcohols in the DHA vehicle have known analgesic effects, which would account for their vehicle effects.\textsuperscript{45,48} This vehicle effect could also explain failures to detect differences in vulvovaginal PGE\textsubscript{2} levels. Nonetheless, the effects of maresin 1 and DHA clearly surpassed their respective vehicles, confirming their potential efficacy for the treatment of vulvodynia. These observations are consistent with at least one other study reporting analgesic effects for maresin 1.\textsuperscript{23}

In the context of translating these findings to patient applications, such vehicle effects could further enhance the therapeutic response in women. However, careful design of the treatment vehicle in future mouse studies would be helpful in eliminating background effects to better evaluate the effects of SPM or PUFA treatment alone. As far as deciding which approach is best, a chemically synthesized SPM or highly purified PUFA-enriched oil, both offer advantages and disadvantages. A pure SPM is more likely to have direct and potent effects, but would require lengthy drug development steps, while fish oil, which is already consumed in diet, could lead to a considerably faster translation. Determining optimal strategies for SPM-based drug development would be a next logical step and could easily be evaluated in our mouse model.

In summary, combined in vitro fibroblast and in vivo mouse modeling represents an effective strategy for testing novel mechanism-based therapies for vulvodynia and possibly other pain conditions. Using this approach, we have identified a class of molecules, the SPMs that may be highly effective in reducing vulvar pain in the clinic. The pathogenesis of vulvodynia involves a hypersensitivity to inflammatory stimuli, which may be overcome by exogenous SPM treatment, which helps to resolve inflammation without inhibiting this vital process. Many of the SPMs tested reduced both IL-6 and PGE\textsubscript{2} levels in vulvar fibroblasts. Topical application of maresin 1 or the PUFA from which it is derived, DHA, are highly effective in reducing measures of pain and inflammation in a mouse model of vulvodynia. Overall, SPMs and their precursors are potentially potent and safe treatments for vulvodynia that address at least part of the disease mechanism.

**Supplementary data**

Supplementary data related to this article can be found at https://doi.org/10.1016/j.jpain.2021.03.144.


35. Hayden MS, Ghosh S: Signaling to NF-kappaB. Genes Dev 18:2195-2224, 2004


47. Mogil JS, Davis KD, Derbyshire SW: The necessity of animal models in pain research. Pain 151:12-17, 2010


