INTRODUCTION

High-dense corneal innervation is essential to ocular surface integrity, and nerve damage impairs tear production, blinking reflex, and epithelial wound healing that results in the loss of transparency and vision. Anatomically, the axons from sensory nerves derived from the ophthalmic branch of the trigeminal ganglion (TG) neurons penetrate the corneal stroma from the limbal area and branch out as the subepithelial nerve network before terminating as free nerve endings in the superficial corneal epithelia.

Factors that can alter corneal innervation include aging, diabetes, and viral and bacterial infections. Moreover, nerve damage also occurs after refractive surgery, such as...
laser-assisted in-situ keratomileusis (LASIK) and photorefractive keratectomy,7,4 and it takes between 3 and 15 years to recover the integrity of corneal nerves.7,10 Consequently, corneal sensitivity decreases and dry-eye disease can develop, causing neuropathic pain, corneal ulcers, and in severe cases, corneal transplant11-13 new therapeutic approaches are needed for treating these complications.

Previous studies from our laboratory have demonstrated that pigment epithelium-derived factor (PEDF), a glycosylated neurotrophic protein, in combination with docosahexaenoic acid (DHA), a member of the ω-3 fatty acid family, play an important role in regenerating the corneal nerves after damage and in maintaining a healthy cornea.14,15 We have successfully tested the action of PEDF + DHA in corneas after experimental surgery and pathologies such as diabetes and herpes simplex virus (HSV1) infection.16-20 Recently, a novel Resolvin D6-stereospecific isomer (named RvD6si) was discovered as the main pro-resolving lipid mediator underlying the mechanism of PEDF + DHA.21 The new lipid stimulates a specific transcriptome in the TG, increasing the induction of genes related to axon growth and modulating genes involved in neuropathic pain.21 Even though RvD6si (4,17-di-hydroxy-DHA) from tears was characterized by LC-MS/MS and its biological activity evaluated by in vivo studies, the lack of a defined stereo structure is a big factor to establish the possible therapeutic use to recover ocular surface health and integrity. This study focused on investigating the configuration of the two hydroxy groups of the RvD6si as well as the biological activities on corneal nerve regeneration after an experimental injury that mimics the nerve damage after refractive surgery. The action of the chemically synthesized pure isomers on the TG transcriptome was also investigated.

2 | MATERIALS AND METHODS

2.1 | Animals

Nine-week-old male CD1 mice were purchased from Charles River (Wilmington, MA, USA) and maintained in a 12-hours dark/light cycle at the Neuroscience Center of Excellence animal care facility, Louisiana State University Health New Orleans, LA. The animals were handled following the guidelines of the Association for Research in Vision and Ophthalmology Statement for the Use of Animals in Ophthalmic and Vision Research, and the experimental protocols were approved by the Institutional Animal Care and Use Committee at Louisiana State University Health New Orleans.

2.2 | Corneal injury and collection of tears

Mice were anesthetized using intraperitoneal injection of ketamine (200 mg/kg) and xylazine (10 mg/kg) and topically anesthetized using one drop of proparacaine hydrochloride solution (0.5%). The center of the cornea of the right eye was demarcated with a 2 mm trephine, and the epithelium and 1/3 of the anterior stroma were gently removed under a surgical microscope with a corneal rust ring remover (Algerbrush II; Alger Equipment Co., Lago Vista, TX, USA), as described previously.20,21 To prevent postoperative infection, 0.3% of tobramycin ophthalmic solution (Henry Schein, Melville, NY, USA) was topically applied to the eye. For consistency, only one investigator (JH) performed all surgeries. For the tears collection, injured mice were treated topically with 10 µL of a mixture of PEDF (0.5 µg/mL) and DHA (50 nM) every 30 minutes for 4 hours after injury, and tears were collected in 1 mL of MeOH as described21 and stored at −80°C until used.

2.3 | Corneal organ culture and inhibitors treatment

Twenty-four hours after injury, 50 mice were euthanized, and the eyes were enucleated. The corneas were excised and kept in ice-cold PBS. Each cornea was placed in a single well of a 48-well plate. A volume of 100 µL of DMEM medium containing inhibitors described in Table 1 was added to cover the entire cornea and incubated for 1 hour. Then, 100 µL of medium with the inhibitors and containing PEDF (50 ng/mL) + DHA (300 nM) was added, and the corneas were incubated for an additional 4 hours. Afterward, 2 mL of media (from 10 corneas) was collected and centrifuged at 19 000 g for 15 minutes at 4°C to remove cellular debris. The supernatant was collected and subjected to lipid extraction.

| Target name   | Inhibitor/abs name | Company—Cat#            | Conc.   | Effect |
|---------------|--------------------|-------------------------|---------|--------|
| P450 (CYP1A2) | Fluvoxamine        | Cayman chemical—15617   | 100 nM  | Good   |
| 15-LOX        | ML351              | Cayman chemical—16119   | 200 nM  | None   |
| 5-LOX         | Zileuton           | Cayman chemical—10006967| 60 µM   | None   |
| 5-LOX Abs     | Mono-Rb            | Abcam—ab169755          | 10 µg/mL| None   |

TABLE 1 | Characteristics of the enzymes inhibitors
2.4 Lipid extraction and LC-MS/MS-based lipidomic analysis

Lipids from organ culture experiments were extracted by the Blight and Dyer method. Briefly, 3.75 mL of a mixture of CHCl₃:MeOH (1:2) was added to 2 mL of sample and 5 μL of the deuterium-labeled internal standard mixture of lipids [AA-d₈ (5 ng/μL), PGD-d₄ (1 ng/μL), EPA-d₅ (1 ng/μL), and LTB₄-d₄ (1 μg/μL)]. The samples were vortexed and stored at −80°C overnight. The next day, to make two phases, 2.5 mL of CHCl₃ was added, vortexed, followed by the addition of 2.5 mL of water (pH 3.5), vortexed, and the pH of the upper phase was adjusted to 3.5-4.0 with 1 N HCl. The lower phase was collected, dried under N₂, resuspended in 1 mL of MeOH, and stored at −80°C until used. Lipids from the injured mice were collected and extracted as described previously. Briefly, 5 μL of sterile PBS was applied in the inferior cul-de-sac of the mouse eye for 30 seconds before the tears were collected with a 5 μL of sterile PBS pipette in 1 mL of ice-cold MeOH, followed by the addition of 2 mL of CHCl₃ and 5 μL of an internal standard mixture of deuterium-labeled lipids. The samples were stored at −80°C overnight, and the lipids were extracted the following day.

For LC-MS/MS analysis, a Xevo TQ equipped with Acquity UPLC with a flow-through needle (Waters Corporation, Milford, MA, USA) was employed as described previously. Briefly, samples were dried under N₂, resuspended in 20 μL of MeOH/H₂O (2:1), and injected into a CORTECS C18 2.7 μm 4.6 × 100 mm column (Water, MA, USA). The column temperature was set at 45°C with a flow of 0.6 mL/min. The initial mobile phase consisted of 45% solvent A (H₂O + 0.01% acetic acid) and 55% solvent B (MeOH + 0.01% acetic acid) and then a gradient to 15% solvent A for the first 10 minutes followed by a gradient to 2% solvent A for 18 minutes, 2% solvent A run isocratically until 25 minutes, and then a gradient back to 45% solvent A for re-equilibration until 30 minutes.

For the spike-in experiments, isolated RvD6si from tears was mixed with chemically synthetized RR-RvD6 or SR-RvD6 at the same molarity. The samples were dried under N₂, resuspended in 20 μL of MeOH/H₂O (2:1), and injected into a CHIRALPAK AD-RH column (Chiral Technologies, Inc, PA, USA). The column temperature was set at 45°C with a flow of 0.6 mL/min. The initial mobile phase consisted of 60% solvent A (H₂O + 0.01% acetic acid) and 40% solvent B (acetonitrile +0.01% acetic acid) and then a gradient to 10% solvent A for the first 15 minutes, followed by 10 minutes of 100% solvent B and then a gradient back to 60% solvent A for re-equilibration until 30 minutes.

Lipid standards (Cayman, Ann Arbor, MI, USA) were used for tuning and optimization. SS-RvD6 standard, [4S,17S-dihydroxy-5E,7Z,10Z,13Z,15E,19Z-docosahexaenoic acid] was a generous gift of Dr Charles Serhan (Harvard University, Boston, MA, USA).

2.5 In vivo treatments

After corneal injury, mice were randomly divided into four groups (8-10 animals per group) and treated topically with 10 μL of RR-RvD6 or SR-RvD6 at the concentration of 1 ng/μL three times per day for 12 days. The lipids dissolved in ethanol were aliquoted for each day’s treatment and stored at −80°C. Before treatment, the ethanol was evaporated under N₂, and the lipids were resuspended in PBS. Vehicle (ethanol evaporated and PBS) was used as the negative control, and PEDF + DHA was used as the positive control. The experiments were double-blinded.

2.6 Measurement of corneal sensitivity

For this study, we used the Belmonte non-contact corneal esthesiometer that is a more reliable method than the standard Cochet-Bonnet esthesiometer to determine the corneal sensation threshold, as described previously. Sensitivity was measured in the holding mice with an air output needle at 3 mm from the cornea and a starting airflow rate of 80 mL per minute. When the mouse blinked, the airflow rate was recorded as the final corneal sensitivity index. As a baseline, sensitivity was measured in each mouse before injury.

2.7 Analysis of corneal nerves

Twelve days after injury and treatment, mice were euthanized, corneas obtained, and whole mounts processed, as described previously. The primary antibodies used were: rabbit monoclonal anti-PGP9.5 (1:500, which is equivalent to 0.22 μg/mL) (ab108986; Abcam, Cambridge, MA, USA), and rat monoclonal anti-substance-P (SP; 1:100, which is equivalent to 1 μg/mL) (sc-21715; Santa Cruz Biotechnology, Dallas, TX, USA). The corresponding secondary antibodies were goat anti-rabbit Alexa Fluor 488 (1:1000, which is equivalent to 0.22 μg/mL) (sc-21715; Santa Cruz Biotechnology, Dallas, TX, USA). Corneas were examined with a fluorescent microscope (Deconvolution microscope DP80; Olympus, Tokyo, Japan), and nerve density was measured using the brush tool in Photoshop CC 2014 (Adobe), as described previously.

2.8 Transcriptome of trigeminal ganglion

At day 12 after injury, mice were euthanized, and the ipsilateral TG from the injury eye side of corneas treated with vehicle, RR-, SR-, or SS-RvD6 were collected and homogenized in 1 mL of TRIzol on ice. Total mRNA from a
single TG in each condition was obtained using an RNeasy mini kit (Qiagen, Germantown, MD, USA), as described by the manufacturer. RNA purity and concentration were determined with a NanoDrop ND-1000 spectrophotometer (Thermo Fisher Scientific), and the samples were stored at −80°C until used. RNA sequencing was performed as described previously21 using the adapted Smart-seq2 protocol.29 Briefly, 10 ng of total RNA was reverse transcribed using the Oligo-dT30VN and template-switching oligo (TSO) primers. The total cDNAs were amplified employing ISPCR primers, and the library was constructed with a Nextera XT DNA library preparation kit (Illumina, San Diego, CA, USA). The libraries were pooled together at the same molarity concentration and sequenced using the NextSeq 500/550 High Output Kit v2 (75 cycles, Illumina). After demultiplexing, RNA-seq data were aligned to the GENCODE GRCh38 mouse primary genome assembly (Release M22, gencodegenes.org/mouse/) using the featureCounts function (Subread v1.6.5 in Ubuntu) BAM files for sequencing data alignment were counted to differential gene expression analysis using DESeq2 package (version 1.28.1).30 The out-putted BAM files for sequencing data alignment were counted using the featureCounts function (Subread v1.6.5 in Ubuntu LTS 16.4 operating system).31 The data were then subjected to differential gene expression analysis using DESeq2 package for R.32 The adjusted p values were regarded as false discovery rate (FDR). Significantly changed genes (FDR < .05) between RR-RvD6 vs vehicle, SR-RvD6 vs vehicle, and SS-RvD6 vs vehicle were subjected to enrichment analysis using Enrichr33 and pathway analysis using IPA (QIAGEN Inc, https://www.qiagenbioinformatics.com/products/ingenuity-pathway-analysis).

### 2.9 Statistical analysis

To calculate the subbasal nerve density, the fluorescent images were changed to grayscale mode and placed against a white background using imaging software (Photoshop, Adobe System, Inc, Mountain View, CA, USA). Each image was carefully drawn with a 4-pixel line following the course of each fiber using the brush tool in the imaging software. The nerve area and the total area of the image were obtained using the histogram tool. The percentage of total nerve area was quantified for each image, as described previously.26,25 Data expressed as mean ± SD of ≥3 independent experiments were analyzed by one-way ANOVA followed by Tukey's honest significant difference post hoc test at 95% confidence level to compare the different groups and considered significant when P < .05. All statistical and graphs were done using Prism 7 software (GraphPad Software, La Jolla, CA, USA) and BioVinci (BioTuring, La Jolla, CA, USA). For the sequencing data, the normalized counts from DE-Seq2 were used as the input of the ANOVA test.

### 3 RESULTS

#### 3.1 Cytochrome P450 is involved in the synthesis of RvD6 isomers

Our recent study has shown that RvD6si release from tears of injured mice treated with PEDF + DHA when separated by LC-MS/MS eluted earlier than the LTB4-d4 and RvD6 standard (Figure 1A). Differences in the biological activities of these two molecules were also demonstrated.21 Therefore, it was important to elucidate the structure of this new isomer derived from the treatment with PEDF + DHA. It has been shown that the standard RvD6, [4S-17S-dihydroxy-DHA], is generated from DHA by the action of a 15-lipoxygenase (15-LOX) followed by a 5-LOX which forms the 17S- and 4S-OH configurations, respectively (Figure 1B). However, DHA could also be catalyzed by the P450 cytochrome system, which forms the 17R-monohydroxy-DHA.34,35 As a result, there are four possible stereo configurations for the hydroxy groups in the same 4,17-dihydro-DHA backbone (Figure 1B). In injured corneas previously incubated with inhibitors of the enzymes and then stimulated with PEDF + DHA, we found that fluvoxamine, an inhibitor of the cytochrome P450, consistently decreases the synthesis of RvD6si. Although the inhibition is 30%, the reduction is significant at a 95% confidence level (Figure 1C). Since inhibitors and antibodies of 15-LOX and 5-LOX had no effect on RvD6si synthesis, we hypothesized that the RvD6si could be 4R-17R-dihydroxy-DHA or 4S-17R-dihydroxy-DHA. To further investigate the structure of this lipid, the total synthesis of these two lipids, RR-RvD6 and SR-RvD6, was conducted.

#### 3.2 RvD6 stereoisomer structures were confirmed by total chemical synthesis

The scheme of 4R-17R-dihydroxy-DHA synthesis is shown in Figure 2A. Briefly, the synthesis began with small and defined starting materials of known configurations, with the carbon atoms of a C1-C13 fragment. The (R) alcohol chirality at C-4 position was unambiguously confirmed and was retained from enantiomerically pure (R)-γ-carboxy-γ-butylactone, a commercially available building block. The C14-C22 fragment was built from cis-4-Heptenal, via an organocatalytic α-oxidation to establish the C-17 R-configuration with excellent enantiomeric excess. With both the C1-C13 and C14-C22 main intermediates at hand, a Z-stereoselective Wittig condensation between the aldehyde and ylide derivative of C1-C13 yielded the docosahexaenoate carbon skeleton of the desired compound in the methyl ester form. Further manipulations and purification were done by reversed-phase HPLC on a Phenomenex Luna® column 5 μm C18 100 Å 250 x 10 mm, 240 nm, 70% MeOH (B)/
H₂O (A) to 100% (B) for 20 minutes afforded stereochemically pure RR-RvD6i, namely 4(R),17(R)-dihydroxydocosa-5E,7Z,10Z,13Z,15E,19Z-hexaenoic acid. The LC/MS showed a great purity (Figure 2B, left). The synthesized product also gave an ultraviolet (UV) chromophore profile with a broadband of absorption, λ max MeOH, at ~240 nm, characteristic of conjugated 1,3-butadiene moieties present in the molecule at carbon positions C5-C7 and C13-C15.

FIGURE 1  Effect of inhibitors on RvD6si synthesis. A, Different retention time of standard RvD6 (SS-RvD6) and the RvD6si isomer isolated from tears of PEDF + DHA-treated mice. The blue peak represents the internal standard LTB₄-d₄. The RvD6si eluted earlier than LTB₄-d₄ and RvD6 standard. B, Possible stereoisomers of RvD6 made from DHA. The 15-LOX derivatives of RvD6 (left panel) contain the 17S-OH, while the P450 derivatives (right panel) contain the 17R-OH on the DHA backbone. 5-LOX catalyzes the formation of 4S-OH, while the 4R-OH might be formed by hydrolysis. C, Incubating corneas with inhibitors/antibodies in the presence of PEDF + DHA shows that only the cytochrome P450 inhibitor Fluvoxamine, significantly reduced the synthesis of RvD6si while 15- and 5-LOX inhibitors/antibodies had no effect. Details about inhibitors and antibodies used are described in Table 1. Values are mean ± SD * P < .05 using Student’s t test with PEDF + DHA group as reference. There were 10 corneas/sample per experiment (total 50 corneas for 5 groups of treatment). An average of three independent experiments was calculated.
The structure of RvD6si was confirmed by total synthesis. A. The synthetic scheme of RR-RvD6 involved key synthetic intermediates C1-C13 (black) and C14-C22 (red) were constructed from chirally pure building blocks that were synthesized in conjunction with stereospecific transformations to furnish the target with absolute stereo- and regiochemical control. B. The purity of RR-RvD6 was confirmed by LC-MS chromatogram (left) while its characteristic conjugated 1,3-butadiene moieties are shown in the molecule at carbon positions C5-C7 and C13-C15, confirmed by UV spectrum (right). B, LC-MS/MS analysis of mouse tears-derived RvD6si using a chiral column shows the separation of four peaks. Chemical synethized RR-RvD6 had the same retention time as the first peak of RvD6si, while the SR-RvD6 partially matched the third peak of RvD6 isomer. Only RR-RvD6 has an earlier retention time than the internal standard LTB₄-d₄ (blue), as shown in Figure 1A.
To distinguish the stereoisomer structure of the RvD6si, a chiral column was used that provided enantiomer separation. Surprisingly, we found that our RvD6si isolated from mouse tears is composed of four different stereoisomers (Figure 2C) named RvD6i_{1-4}. Among those, the retention time of RvD6i\textsubscript{1} matched the retention time of the synthesized 4R-17R-dihydroxy-DHA (RR-RvD6) while RvD6i\textsubscript{1} partially matched the 4S-17R-dihydroxy-DHA (SR-RvD6) (Figure 2C). These two chemical synthesized lipids generate the same peaks in a spike-in experiment in which we mixed the standards with RvD6si isolated from mouse tears. Aside from the unidentified RvD6i\textsubscript{2} and RvD6i\textsubscript{4}, our data show that the RvD6si we have recently reported\textsuperscript{21} contains a 4R-17R- and a 4S-17R-dihydroxy-DHA.

3.3 Biological activities of RvD6 stereoisomers on corneal nerve regeneration

For these experiments, we treated the mouse injured corneas with pure RR-RvD6 or SR-RvD6 (10 ng/eye, three times per day) for 12 days. The PEDF + DHA-treated group was used as positive control, while the vehicle-treated group was the negative control (Figure 3A). Corneal sensitivity shows a significant increase at day 3 after injury only with RR-RvD6 treatment (Figure 3B). The corneal sensitivity was 87 ± 10% in the RR-RvD6-treated corneas while the vehicle was 61 ± 8 (P = 0.0015). This early effect is similar to our reported data for the RvD6si isolated from mouse tears.\textsuperscript{21} Interestingly, when we look to the sensation of single mouse in each group, there were 50% (4/total of 8) mice that had more than 90% of sensitivity recovery at day 3, while the percentage of recovery with SR-RvD6, PEDF + DHA, and vehicle treatments were 25% (2/8), 22% (2/9), and 0% (0/9), respectively. On day 6, all mice in the RR-RvD6-treated group recovered their corneal sensation (Figure 3B). This robustness made the RR-RvD6-treated group significantly different to vehicle as well as to PEDF + DHA-treated corneas. SR-RvD6 also showed a higher score of corneal sensitivity recovery than vehicle. By day 9, all treated groups had a significant increase in sensitivity with respect to the vehicle-treated mice.

Corneal nerve regeneration was evaluated at 12 days after injury. Whole-mount staining of corneal total nerves (PGP 9.5 positive) (Figure 3E,C) and SP-positive nerves (Figure 3F,D) showed that mice treated with RR-RvD6 had the highest corneal nerve density of the two stereoisomers-treated mice compared to vehicle, and higher significance than the mice treated with PEDF + DHA. The percentage of total corneal nerve regeneration in the RR-RvD6-treated corneas was 75.5 ± 11.3, followed by 61.4 ± 6.8, 61 ± 13.4, and 46.9 ± 6.5% in PEDF + DHA, SR-RvD6, and vehicle, respectively. Similar results were found with substance P (SP)-positive nerves.

3.4 Distinctive transcriptome in the TG after RvD6 stereoisomers treatment

Using RNA-seq analysis, we investigate the effect of RR-, SR-, and SS-RvD6 isomers in the transcriptome of the TG at day 12 after corneal injury and treatment. Our previous studies have shown that expression of genes related to neuropathic pain was modulated in the TG of mice treated with the RvD6si isolated from tears.\textsuperscript{21} Analyzing these genes, we found that expression of tachykinin 1 (Tac1) that encodes SP and calcitonin-related peptide alpha (Calca) and calcitonin-related peptide beta (CalcB), both encoding calcitonin gene-related peptide (CGRP), decrease by treatment with the three lipid isomers (Figure 4A). Interestingly, the gene expression of transient receptor potential melastatin 8 (Trpm8), which encodes the TRPM8 ion channel expressed in sensory neurons and contributes to decrease neuropathic pain,\textsuperscript{36,37} was activated by the two new synthetic standards RR-RvD6 and SR-RvD6 but not by SS-RvD6. This result might explain why SS-RvD6 showed a slower corneal nerve regeneration with significant recovery of corneal sensation found at day 9 after injury, as we reported previously.\textsuperscript{21}

Our early studies using the RvD6si isolated from mouse tears had demonstrated that Rictor, a gene that encodes the rapamycin-insensitive complex 2 for mTOR, was involved in corneal nerve regeneration.\textsuperscript{21} Increased gene expression of this upstream regulator was found in all treated mice except for those treated with SS-RvD6 (Figure 4B). We also determined Rictor upregulated and downregulated genes (Figure 4C), and the IPA activation z-score of this transcription factor predicts that RR-RvD6 and SR-RvD6 are modulators of many of these genes (Figure 4C). In the prediction, there are about 5 of the 9 Rictor genes upregulated in the TG (red) that their induction was increased when corneas were treated with the SR-and RR-RvD6 compared to vehicle, and there were 36 of the 38 downregulated genes by Rictor (blue) that were inhibited by RR and SR-RvD6 respect to vehicle. KEGG pathway enrichment analysis of these 38 downregulated genes shows that they were mainly found in degenerative diseases such as Parkinson's disease, Alzheimer's disease, and Huntington's disease (Figure 4D).

Next, we investigated the potential effect of RR-RvD6 on biological systems using its specific changed genes list obtained by RNA-seq. The IPA Bio Functions tool predicts that most of the biological functions upregulated by RR-RvD6 in the TG belong to neuronal and axonal growth with 12 significant increase biological functions (Figure 5A). A graphic summary of the action of RR-RvD6 shows that the core of its action in the TG is hepatocyte growth factor (hgf) that affect functions related to cellular skeleton components and microtubule dynamics, which in turn can increase neuronal branching, quantity, and density of sensory neurons (Figure 5B). We found a significant increase in the
gene expression of *hgf* with both SR- and RR-RvD6 cornea treatments (Figure 5C).

### 4 | DISCUSSION

Previous studies from our laboratory have demonstrated that PEDF + DHA treatment decreases inflammation and increases wound healing and nerve regeneration using different models of corneal injuries and infection.\(^{16-19}\) The mechanism involves the phospholipase A2 activity of the PEDF receptor that releases DHA incorporated in the membrane phospholipids and leads to the synthesis of docosanoids.\(^{15}\) By LC-MS/MS isolation and fragmentation analysis, we identify one of the docosanoids release in tears as an RvD6 isomer with a retention time different than the RvD6 standard with SS-configuration.\(^{21}\) The new isomer enhances corneal wound healing, increased recovery of sensitivity, and activates nerve regeneration.\(^{21}\)

Here, we showed that RvD6si release to the media from injured corneas stimulated by PEDF + DHA were significantly inhibited by Fluvoxamine, an inhibitor of cytochrome P450, but not by 15-LOX or 5-LOX inhibitors. These data suggested that there is a 17R-monohydroxy group in the DHA derivative. Therefore, the chemical synthesis of 4R,17R-dihydroxy-DHA (RR-RvD6) and 4S,17R-dihydroxy-DHA (SR-RvD6) was performed in order to confirm the isomer structure. Using an innovative method, we obtained these two molecules with more than 90% purity (Figure 2B), which by chiral separation, provided two matched peaks to our RvD6si isolated from mouse tears (Figure 2C).

Our experiments demonstrate that 5-LOX inhibitors and antibodies did not show a decrease in the synthesis of RvD6si. Therefore, we do not have a satisfactory explanation for how the synthesis of SR-RvD6i occurs. One possibility is that the inhibitors were not specific enough to block the synthesis of this isomer. Alternatively, the isomer is produced by a pathway that does not include 5-LOX. Further studies will be needed to determine the origin of SR-RvD6 in tears after injury. When the biological activities of RR-RvD6 and SR-RvD6 on corneal sensation and nerve recovery were assessed, we found that RR-RvD6 was the only isomer that increased the sensitivity 3 days after injury, suggesting that this lipid mediator has a higher potency than SR- and SS-RvD6 isomers. Moreover, the percentage of total corneal nerve recovery and the SP-positive nerves in the RR-RvD6 group were even higher than in the PEDF + DHA-treated group (Figure 3). The results demonstrate that the RR-RvD6 isomer has a stronger effect on corneal nerve regeneration than the mixture of isomers obtained after PEDF + DHA treatment. Even though PEDF + DHA was applied in the same manner (three times per day for 12 days), the higher efficiency of RR-RvD6 could be caused by its defined structure and higher molarity since the conversation of PEDF + DHA to RvD6si is low. For this reason, the administration of 1 ng/µL of RR-RvD6 is much higher than the actual amount synthesized by PEDF + DHA in vivo.

One of the advantages of RvD6si treatment was pain management.\(^{21}\) Unlike other ongoing therapies, such as NGF,\(^{38}\) RvD6si provided an alternative way to increase corneal nerve density without stimulating the expression of genes associated with pain in the TG. Using RNA-seq analysis of the homolateral TG, we show that both RR-RvD6- and SR-RvD6-treated mice show a significantly lower gene expression than the vehicle of genes encoding pain-related neuropeptides, such as SP and CGRP while Trpm8, the transient channel that diminished in dry eye-like pain conditions,\(^{36,37,39,40}\) was significantly elevated in SR- and RR-RvD6-treated corneas than vehicle and SS-RvD6-treated groups (Figure 4B). Tac1 encodes SP, one of the most abundant neuropeptides expressed in corneal nerves\(^{25-27}\) that causes pro-inflammatory effects and chronic clinical pain,\(^{41}\) while Calcb encodes CGRP, another abundant neuropeptide\(^{42}\) that plays an essential role in neurogenic inflammation and pain.\(^{43}\) The inhibition of these neuropeptides in the TG suggests successful management of potential neuropathic pain by treatment with RR-RvD6 and SR-RvD6. Future studies to further investigate the action of the new lipids in models of corneal pain will be needed. In addition, the topical application of these lipid mediators increases corneal nerve outgrowth by fostering the Rictor-related signaling as well as other neurotrophic genes. Due to the important role of Rictor in downregulated several genes in the TG, we discovered an important role of Rictor in decreasing the induction of genes involved in degenerative diseases using the KEGG pathway enrichment analysis. It will be important to investigate the role of Rictor in models of brain diseases.

Our RNA-seq data also shows a significant increase in the expression of *Hgf*, which encodes HGF, a paracrine growth factor that is expressed in corneal stroma and stimulates...
epithelial proliferation and suppresses inflammation after injury through its receptor Met tyrosine kinase.44-46 IPA predicts that RR-RvD6 activates a group of axonal outgrowth genes in the TG involved in neuritogenesis and branching of neurites, which are important to regenerate corneal nerves, with HGF being a central factor in these biological functions.
FIGURE 4  Transcriptome signature in the trigeminal ganglion unravels the role of RvD6 isomers. A, RNA-seq normalized counts of genes involved in inflammation and pain, including Tac1, Calca, Calcb, and Trpm8. The p values are derived from ANOVA post hoc Dunnett’s multiple comparisons test with vehicle as a reference. Mean and SD are depicted as the lines. Each data point represents an ipsilateral TG of a mouse. B, The Rictor signaling activated by RvD6 isomers. The blue columns represent the relative normalized fold changes of Rictor gene analyzed by RNA-seq, while the red columns represent the activation z-score of Rictor signaling predicted by IPA using RNA-seq data as input. The SS-RvD6 z-score was insufficient to show the activation of RICTOR pathway in the IPA. C, The RNA-seq normalized counts heatmap of RICTOR regulated genes. The genes activated (red) and inhibited (blue) are grouped and clustered across all treatment groups. There are about 5 upregulated genes and 36 downregulated genes by SR- and RR-RvD6. D, The KEGG-pathway prediction of RICTOR downregulated genes from C. Bars were sorted by p value. The length of the bar represents the significance of the pathway, while the lighter the color, the higher the significance.

FIGURE 5  Predicted action of RR-RvD6 on nerve growth in the trigeminal ganglia. A, The IPA of nervous system development and functions in the TG after RR-RvD6-treated corneas in mice. The bar chart shows the activation z-score (blue) and p value (red) of 12 significant groups of biological functions. B, The graphical summary of RR-RvD6 induced axon regeneration with hgf expression as a central upstream modulator. Dash lines represent that Hgf requires several intermediate signaling pathways to reach the biological function. C, Hgf was found significantly elevated by SR- and RR-RvD6 treatment but not by SS-RvD6.
(Figure 5). Previous studies have shown that HGF signaling enhances axonal growth of periphery sensory neurons from the skin in a mice model\(^7\) and increases peripheral nerve regulation after sciatic nerve injury.\(^8\) It is interesting that cornea nerves can transmit signals to the TG,\(^9\) but further studies will be needed to decode the working mechanism of RR-RvD6 on the cornea-TG axis, especially how topical treatment with RR-RvD6 can affect the cornea-innervating neuron in the TG as well as how RR-RvD6 stimulated corneal epithelial cells can interact with corneal nerve axons to enhance nerve recovery.

Recently, in an alkali burn rat model as well as in human corneal epithelial cells in culture challenged with IFN\(\gamma\), we found that treatment with RR-RvD6 decreases inflammation, cytokine expression, and senescence programming, demonstrating the protective bioactivity of this lipid.\(^50\) Interestingly, RR-RvD6 decreases the expression of ACE2, the receptor of coronavirus SARS-CoV-2, and the binding of the receptor-binding domain (RBD) of the virus spike protein to ACE2. The data suggest that RR-RvD6 may have therapeutic value in decreasing the damage produced by this viral infection.

In conclusion, this study uncovers the stereochemical structure of RvD6si released from tears in a mouse injury model as a mixture of four isomers, with 4R-17R-dihydroxy-DHA being a more biologically potent isomer than the mixture obtained after PEDF + DHA treatment. In addition, RR-RvD6 stimulates a specific transcriptome in the TG with Rictor and hgf as key regulators to modulate axonal growth and decrease pain, which is not reported by other docosanoids.\(^51-55\)

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CONFLICT OF INTEREST

The subject matter of this manuscript is protected by a US provisional patent application assigned to the Board of Supervisors of Louisiana State University and Agricultural and Mechanical College (LSU) and licensed to a company, CurVirBiotech, founded by co-author Nicolas G. Bazan.

AUTHOR CONTRIBUTIONS

H. E. P. Bazan and T. L. Pham designed the experiments. T. L. Pham performed acquisition and analysis of the data, and writing and critical review of the manuscript; A. H. Kakazu coded and prepared the drugs, performed lipid extraction, and helped with RNA extraction and sensitivity measurement. J. He did the injury, treatments, and isolation of corneal tissues. R. Nshimiyimana and N. A. Petasis synthesized RR- and SR-RvD6i. B. Jun performed the LC-MS/MS lipid analysis and acquisition of data. J. He, A. H. Kakazu, and B. Jun provided a critical review of the manuscript. N. G. Bazan contributed to the interpretation of results and reviewed the manuscript. H. E. P. Bazan supervised the study, wrote and reviewed the manuscript, and was responsible for the integrity of this work.

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