Discovery and Development of a Novel mPGES-1/5-LOX Dual Inhibitor LFA-9 for Prevention and Treatment of Chronic Inflammatory Diseases

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Background: Non-steroidal anti-inflammatory drugs, cyclooxygenase (COX)-2 selective inhibitors, have been explored for prevention and treatment of several inflammatory chronic conditions including arthritis, and cancer. However, the long-term use of these drugs is associated with gastrointestinal, renal, and cardiovascular side effects. Later, COX-5-lipoxygenase (5-LOX) dual inhibitors (eg, licofelone) have been developed but did not enter into the market from the clinical trials due to COX-1/2 inhibition-associated side effects. Hence, targeting microsomal prostaglandin E synthase-1 (mPGES-1) and 5-LOX can be an ideal approach while sparing COX-1/2 activities for development of the next generation of anti-inflammatory drugs with better efficacy and safety.

Materials and Methods: In silico (molecular modelling) studies were used to design a mPGES-1/5-LOX dual inhibitory and COX-1/2 sparing lead molecule licofelone analogue-9 (LFA-9) by modifying the pharmacophore of licofelone. In vitro cell-free enzymatic (mPGES-1, 5-LOX, COX-1/2) assays using fluorometric/colorimetric methods and cell-based assays (LPS-induced PGE2, LTB4, and PGL2 productions from macrophages) using ELISA technique, isothermal calorimetry, and circular dichroism techniques were performed to determine the mPGES-1/5-LOX inhibitory efficacy and selectivity. Anti-inflammatory efficacy of LFA-9 was evaluated using a carrageenan (inflammogen)-induced rat paw edema model. Infiltration/ expression of CD68 immune cells and TNF-α in paw tissues were evaluated using confocal microscope and immunoblot analysis. Anti-cancer effect of LFA-9 was evaluated using colon spheroids in vitro.

Results: LFA-9 inhibited mPGES-1/5-LOX and their products PGE2 and LTB4, spared COX-1/2 and its product PGL2. LFA-9 bound strongly with human mPGES-1/5-LOX enzymes and induced changes in their secondary structure, thereby inhibited their enzymatic activities. LFA-9 inhibited carrageenan-induced inflammation (70.4%) in rats and suppressed CD68 immune cell infiltration (P ≤ 0.0001) and TNF-α expression. LFA-9 suppressed colon tumor stemness (60.2%) in vitro through inhibition of PGE2 (82%) levels.

Conclusion: Overall study results suggest that LFA-9 is a mPGES-1/5-LOX dual inhibitor and showed anti-inflammatory and colorectal cancer preventive activities, and warranted detailed studies.

Keywords: mPGES-1/5-LOX dual inhibitor, LFA-9, drug design, anti-inflammatory agent, cancer chemoprevention

Introduction
Non-steroidal anti-inflammatory drugs (NSAIDs) such as aspirin, naproxen, and sulindac, cyclooxygenase (COX)-2 selective inhibitors (eg, celecoxib), have been explored for prevention and treatment of several chronic inflammatory conditions including arthritis, and cancer. However, the application of these drugs, particularly
long-term use, is associated with gastrointestinal, renal, and cardiovascular side effects and metabolic pathway diversion of substrate arachidonic acid into pro-inflammatory leukotrienes by activation of 5-LOX enzyme.\(^1\)\(^2\) Later, COX and 5-lipoxygenase (CLOXIB) dual inhibitors (eg, licofelone) were developed but most of them did not enter into the market from clinical trials due to safety concerns.\(^3\) These side effects are due to COX-1/2 inhibition and their downstream product prostacyclin I\(_2\) (PGI\(_2\)) and renal function disruption.\(^4\)\(^5\) Hence, mechanistic studies suggest that sparing COX-1/2 and selectively targeting microsomal prostaglandin E synthase-1 (mPGES-1) and 5-LOX would be a safer and effective strategy to inhibit PGE\(_2\) and leukotrienes (LTs), respectively, and to spare PGI\(_2\). In this scenario, several mPGES-1/5-LOX dual inhibitors have been developed for the next generation of anti-inflammatory drugs (Figure 1A).\(^6\)\(^7\)

Licofelone (formerly known as ML-3000), a potent inhibitor of COX-1/2, 5-LOX, and weak inhibitor of mPGES-1 inhibitor, has been evaluated in clinical trials for several inflammatory and oncolgic diseases\(^8\)\(^9\) (Figure 1). Previously, we have shown that licofelone had highly preventive efficacious effects in several preclinical models of colon, lung, pancreatic and urinary bladder cancer.\(^10\)\(^11\)\(^12\)\(^13\) However, its COX-inhibition-associated side effects have somewhat limited its clinical applications.\(^14\)\(^15\) Licofelone is a potential pharmacophore with which to develop mPGES-1/5-LOX dual inhibitory, and COX-1/2 activity sparing analogs.\(^8\) Previously, several groups made efforts to develop licofelone analogs by modifying its pharmacophore at the carboxylic end to yield mPGES-1/5-LOX inhibitor with COX-1/2 sparing property, but were not fully successful in developing dual mPGES-1/5-LOX inhibitors for clinical translation.\(^8\) In addition, the structural variability and the absence of crystal structures of the target enzymes, among experimental animals (rodents) and humans are limiting the lead optimization and pre-clinical development and clinical translation of mPGES-1/5-LOX inhibitory agents,\(^16\)\(^17\) (Supplemental Information Figures 1–4; Supplemental Information Table 1). So far, none of the reported potent inhibitors of human/mouse/rat specific mPGES-1 and 5-LOX inhibitors for pre-clinical evaluation and clinical translation.\(^18\)

Thus, the main aim of the present work was to modify the licofelone pharmacophore at its carboxyl end using a hit library (active fragments over 1000) through fragment-based drug design strategy to target mPGES-1/5-LOX while sparing COX-1/2 activities of human, mouse, and rat to facilitate its preclinical development and further to clinical translation. In this study, we modeled the three-dimensional structures of some of the target proteins as the crystal structures of some target proteins are not available. Among over 1,000 designed molecules, an optimized lead molecule (licofelone analogue (LFA-9) (Figure 1B) was selected and identified as a potential candidate for development of mPGES-1/5-LOX inhibitor based on molecular docking studies and druggable properties. Further, LFA-9 was synthesized and in vitro cell-free and cell-based assays were performed to determine its selective inhibitory effect on human/mouse/rat mPGES-1/5-LOX and their respective products PGE\(_2\) and LT\(_B\), and sparing effect on COX-1/2 activity and PGI\(_2\) production. Isothermal calorimetric (ITC) and circular dichroism (CD)-based studies were performed to determine the binding affinity and structural changes induced by LFA-9 on human mPGES-1/5-LOX enzymes. Further, in vivo anti-inflammatory activity of LFA-9 was evaluated using carrageenan (inflammogen)-induced rat paw edema model. In addition, the anti-cancer effect of LFA-9 was evaluated by in vitro assay using colon spheroids, since inflammation causes the tumorigenesis of colorectal cancer.\(^19\)\(^20\)

### Materials and Methods

#### General Information

Human mPGES-1 was expressed and purified from *Escherichia coli* Rosetta\(^21\) and also purchased from Cloud-Clone Corp. (Item No. RPA167Hu01) (Houston, TX, USA). ELISA kits for PGE\(_2\) (Item No. 500141), LT\(_B\)\(_4\) (Item No. 520111), PGI\(_2\) (\(\alpha\)-keto prostaglandin) (Item No. 515211), and TX\(_B_2\) (Item No. 501020) quantification, recombinant human COX-1 (Item No. 17616) and PGG\(_2\) (Item No. 17020) were purchased from Cayman Chemical (Ann Arbor, MI, USA). Glutathione (GSH) (Cat No. GI317), lamda-carrageenan (Cat. No. sc-216084) was purchased from Santa Cruz Biotechnology (Dallas, TX, USA). Fluorometric assay kits for 5-LOX (cat no. K980), COX-2 (Cat No. K547), and COX-1 (Cat No. K548) activity assays were purchased from Biovision, Inc. (Milpitas, CA, USA) and Cloud-Clone Corp. (Houston, TX, USA). CD68 antibody with Alexa Fluor\(^6\) 700 (Novus Biologicals, CA, USA; 600985), CD68 antibody (catalog sc-9139; Santa Cruz, Dallas, TX, USA), TNF-\(\alpha\) antibody (Bios, Woburn, MA, USA), (BS-2018R), beta-actin (abcam, ab8227) are purchased from respective vendors and used in the study. Mouse- and rat-specific COX, mPGES-1 enzymes were isolated from their vas deferens and seminal vesicles.\(^22\)\(^23\)\(^24\) Licofelone and LFA-9 were
Synthesis and Characterization of (2-[[2-[2-(4-Chlorophenyl)-6,6-Dimethyl-1-Phenyl-5,7-Dihydropyrrolizin-3-Yl]acetyl] amino]acetic Acid) (LFA-9)

The synthesis of LFA-9 was achieved in a two-step synthetic strategy (Supplement Figure 5A) as per previously patented procedure. Briefly, to a solution of licofelone (1) (28.45 g, 75 mmol) in dichloromethane (300 mL) was added N-hydroxysuccinimide (9.50g, 82.5 mmol) in dichloromethane (300 mL) and N,N'-dicyclohexylcarbodimide (17.03 g, 82.5 mmol) in dichloromethane (100 mL). The reaction mixture was stirred at room temperature overnight. The progress of the reaction was monitored by thin layer chromatography (TLC). After completion of the reaction, N,N'-dicyclohexylurea (byproduct) was filtered.

The Health Sciences Center (OUHSC) Molecular Biology-Proteomics Facility using a Waters Synapt G2-s mass spectrometer.

A

Membrane Phospholipids

Aspirin
Nimesulide
Ibuprofen
Indomethacin
Celecoxib
Acetaminophen
Licofelone

Phospholipase A₂

Arachidonic acid

Prostaglandin H₂

Leukotrienes

Prostacyclin I₂

Normal physiological functions of cardiovascular, renal and gastrointestinal tract

Inflammation and Carcinogenesis

B

HOOC

Cl

N

COOH

(i)

(ii)

Figure 1 (A) Arachidonic acid pathway and its inhibitors in clinical use for inflammatory and oncological diseases. (B) Chemical structures of licofelone (i) and LFA-9 (ii).
on a sintered funnel, and filtrate was evaporated on a rotary evaporator under reduced pressure to obtain licofelone-NHS in a quantitative yield (2). This compound was used immediately for the next step without further purification.

To a solution of licofelone-NHS (35.79 g, 75 mmol) in N,N-dimethylformamide (650 mL) was added glycine (5.63 g, 75 mmol) in 0.2 M Na₂HPO₄ (100 mL, pH 8.0). The reaction mixture was stirred at room temperature overnight. The progress of the reaction was monitored by TLC. After completion of the reaction, solvents were evaporated on a rotary evaporator under reduced pressure, and the product was extracted into ethyl acetate (2 x 300 mL). The combined organic layer was washed with water (2 x 100 mL) and with brine (2 x 100 mL). The organic layer was dried over sodium sulfate, and solvents were evaporated on a rotary evaporator under reduced pressure to obtain the crude LFA-9 (3). The crude compound was purified on a silica gel column by eluting with ethyl acetate/hexane solvent mixture (26.4 g, 80.5%). Purity of LFA-9 as determined by high performance liquid chromatography (HPLC). Melting point, H and ¹³C, NMR and high-resolution mass spectroscopic (HRMS) techniques were used to determine the chemical structure of LFA-9.

**Cell-Free Enzyme Assays**

**5-LOX Assay**

5-LOX (human/mouse/rat) assays were performed using fluorometric-based 5-LOX inhibitor screening kits according to the manufacturer’s instructions (Biovision, Inc., Milpitas, CA, USA). 5-LOX converts arachidonic acid into 5-hydroperoxyeicosatetraenoic acid (5-HPETE). This intermediate interacts with the specific probe and leads to fluorescence emission, which can be measured at Eₓ/Eₘ 500/536 nm.

5-LOX + (Arachidonic acid) Substrate → 5-HPETE (Intermediate) + Probe → Fluorescence (Eₓ/Eₘ: 500/536 nm)

Different concentrations of LFA-9 or licofelone (0.1–100 µM) were incubated with 5-LOX (2 µU (units)) in enzymatic assay buffer mixture and the addition of substrate (20 µL of 100 µM arachidonic acid). Then, we read fluorescence (Eₓ/Eₘ 500/536 nm) kinetically from 0–10 minutes. The untreated 5-LOX enzymatic wells were considered a control (100% activity of the enzyme). We calculated relative fluorescence units (ΔRFU) for all samples by subtracting RFU at time t₁ (initial) from RFU at time t₂ (end) (10 minutes). Slopes for all samples were calculated by dividing ARFU by time Δ (t₁-t₂). 0.01% DMSO or methanol (solvents used to dissolve the compound) was used for solvent control. Non-enzymatic ΔRFU were measured and subtracted from the control and test values. Experiments were conducted in triplicate. Percent inhibition of 5-LOX activity was calculated based on the following equation:

% inhibition = (slope of the enzyme control – slope of test compound/slope of enzyme control) x 100% relative activity = [slope of test/slope of enzyme control] x 100

**mPGES-1 Assay**

An mPGES-1 assay was performed according to the method described previously by Kim et al.¹¹ The appropriate amount of mPGES-1 in 100 µL of assay buffer (15 mM Tris-HCl, pH 7.4, and 1 mM GSH) was added to each well of a 96-well plate. The plate was incubated for 15 minutes at 20°C and an enzymatic reaction was initiated by addition of 20 µL of cold PGH₂ (final conc. 2.8 µM). The plate was further incubated for 45 seconds at room temperature. Incubation was immediately terminated by adding 20 µL of SnCl₂ solution in 1N HCl. PGE₂ levels were estimated using PGE₂-monoclonal antibody-based ELISA kits (Cayman Chemical, Ann Arbor, MI, USA). Enzyme inhibitory activity of control (without compound) and/or test chemical compounds was measured by the incubation of 2 µL of a compound (0.1–100 µM) in buffer with mPGES-1 for 20 minutes prior to the addition of substrate PGH₂. Percent inhibition (triplicate) was calculated from PGE₂ reduction in test wells (compound-treated) compared with control (untreated).²¹ The results were expressed in IC₅₀.

**COX-1/2 Assay**

COX-1/2 (human/mouse/rat) assays were performed using a fluorometric-based COX inhibitor screening kit according to the manufacturer’s instructions (Biovision, Inc., Milpitas, CA, USA). Rat and mouse COX-1/2 were isolated from seminal vesicles. The assays were performed using kits, which are provided with isoform-specific inhibitors. COX converts arachidonic acid into PGG₂. This intermediate interacts with a specific probe that leads to fluorescence emission, which can measure fluorescence at Eₓ/Eₘ 535/587 nm.
Different concentrations of LFA-9 or licofelone (0.1–100 μM) were incubated with COX (2 μU) in enzymatic assay buffer mixture with addition of substrate (20 μL of 100 μM arachidonic acid). Then, we read fluorescence (Ex/Em 535/587 nm) kinetically from 0–10 minutes. Untreated COX enzymatic mixture was considered a control (100% activity of the enzyme). We calculated relative fluorescence units (ΔRFU) for all samples by subtracting RFU at time t1 from RFU at time t2 (10 minutes). Slopes for all samples were calculated by dividing ΔRFU by time Δt (t1-t2). 0.01% DMSO or methanol (solvents used to dissolve the compound) was used for solvent control. Non-enzymatic ΔRFU were measured and subtracted from the control and test values. Experiments were conducted in triplicate. Percent inhibition of COX-1/2 activity was calculated.

Cell-Based Assay Using Human/Mouse/Rat Macrophages

Human macrophages were obtained from commercially derived THP1-human monocytes (ATCC; batch number:70005912) using phorbol 12-myristate 13-acetate. Rat/mouse peritoneal macrophages were isolated as per the protocol described previously. Briefly, after euthanasia to the animals, the superficial skin was disinfected with 70% ethanol, and then the skin on the abdomen was removed while the peritoneum was left intact. Animals were then injected with 5 mL ice-cold Dulbecco’s Modified Eagle Medium (DMEM) media, followed by gentle shaking to allow for the distribution of the lavage fluid throughout the peritoneal cavity. The DMEM lavage fluid was then collected back using a 5-mL syringe attached to a 25-gauge needle. Peritoneal macrophages were determined by adherence to glass slides (1 hour, 37°C). Unattached cells were washed off and attached cells were studied for morphology under light microscopy for the experiment. Cell number and viability was confirmed based on the Trypan-blue exclusion method. Then, cells were washed three times with DMEM + 10% fetal bovine serum and seeded in 96- or 24-well plates with an equal number (density of 10,000 cells/well for 96-well; 40,000 cells/well for 24-well) treated with non-toxic doses of LFA-9 and licofelone, and then stimulated with LPS (10 μg/mL) for 24 hours. The culture supernatant was collected for estimation of exogenous PGE2, PGI2, and LTB4 levels. Percent inhibition and IC50 were calculated.

Molecular Modelling and Docking Studies

Human mPGES-1 (trimeric form) (PDB ID: 4YL3), human 5-LOX (PDB ID: 3O8Y), human COX-2 (PDB ID: 5KIR), and mouse COX-2 (PDB ID: 4COX) three-dimensional structures were obtained from Protein Data Bank and used in docking studies. mPGES-1 (mouse and rat), 5-LOX (mouse and rat), and COX-1 (human, mouse and rat), COX-2 (mouse) crystal structures were not available. Thus, three-dimensional structures of these proteins were modelled for molecular docking studies. Crystal structures of the aforementioned proteins were modelled based on their protein sequences (source: NCBI protein) using Swiss-Model via the ExPaSy web server (https://swissmodel.expasy.org). We validated the structure through Ramachandran plots using Rampage (http://mordred.bioc.cam.ac.uk/~rapper/rampage.php). Sequence similarity between different species of mPGES-1 was determined using the NCBI blast program (https://blast.ncbi.nlm.nih.gov/Blast.cgi). Co-crystallized ligands and water molecules were removed from target protein using ArgusLab. Ligands were prepared using ChemOffice (Cambridge, UK). Energy minimization was done using molecular mechanics, which was executed until the root mean square value reached below 0.001 Kcal/mol. Such energy-minimized ligands and receptors were used for docking studies using GEMDOCK (Generic Evolutionary Method for molecular DOCKing), which is a generic evolutionary method with an empirical scoring function for molecular docking (protein–ligand docking). A population size of 300 with 70 generations and three solutions were used in docking accuracy setting. PyMol was used for better visualization of interactions.

Binding Assay Using Isothermal Titration Calorimetry

Isothermal titration calorimetry (ITC) is a well-established technique that can determine all the thermodynamic parameters, including affinity, of a binding interaction between an enzyme and a small molecule. ITC (VP-ITC manufactured by MicroCal [now part of GE Healthcare]) was used to evaluate the binding efficacy of LFA-9 on the target enzymes. Compounds (up to 1 mM) and proteins (1 mM) are dissolved in phosphate buffered saline and degassed before use in the experiment. A series of injections of the compound were administered to an enzyme-containing well. The heat signal will approach zero as the limiting reactant becomes saturated. Accurate fitting of the
isotherm was performed using Origin software to calculate the binding affinity dissociation constant (KD).  

**Determination of Conformational Changes in the Secondary Structure of mPGES-1/5-LOX/COXs Using the Circular Dichroism (CD) Technique**

Native and LFA-9 (0.1–10 µM)-treated mPGES-1/5-LOX/COXs were dissolved in phosphate-buffered saline (pH 7.4) with a final protein concentration of 200 µM. The samples were transferred to a 1-mm path length quartz cuvette. Circular dichroism spectra in the wavelength range 185–300 nm was acquired in-house with a Jasco 715 Circular Dichroism (CD) Spectropolarimeter (Jasco Corp., Japan) using a bandwidth of 1 nm and a response time of 1 second. Multiple scans were accumulated for each sample. All spectra were corrected by subtraction of buffer blanks. The percent change in secondary structure of proteins was calculated using a CAPITO (CD analysis and plotting tool) server.

**Comparative Analyses of the Drug-Like Properties of LFA-9 and Licofelone**

The Lipinski rule of five helps in distinguishing between drug-like and non-drug-like molecules. Lipinski’s rule of five states that most drug-like molecules have: - Log P ≤ 5; molecular weight ≤ 500; number of hydrogen bond acceptors ≤ 10; and number of hydrogen bond donors ≤ 5. In addition, the number of rotatable bonds are also calculated to determine their bioavailability. Molecules violating more than one of these rules may have problems with bioavailability. The drug-like properties of LFA-9 and licofelone were calculated using a molinspiration tool.

**In vitro Metabolic Stability of LFA-9 Was Assessed on S9 Fraction of Mouse/Rat Liver**

In vitro metabolic stability of LFA-9 was assessed on S9 fraction of mouse/rat liver activated with glutathione, UDP-glucuronic acid, NADPH, PAPS (sulfonation). S9 fraction of mouse/rat liver was isolated and further activated with NADPH (40 mM), UDPGA (20 mM), and GSH (2 mM), respectively while that of PAPS (2 mg/mL) before treatment with LFA-9 and licofelone (500 µM) at different time intervals (0, 15, 30, 45, and 60 minutes). Further, the compound was extracted using acetonitrile and quantified using HPLC. Percent of remaining compound was calculated by using the following formula: (Peak area at 0 min - Peak area at 60 min/Peak area at 0 min) x 100.

**Colon Tumor Spheroid Culture Using Apc<sup>Pirc/+</sup> (Pirc) Rats**

Isolated crypts or single cells were isolated from colorectum of Apc<sup>Pirc/+</sup> rats and cultured as previously described. Briefly, crypts or single cells were plated in ultra-low attachment wells of 24-well plates. After cell seeding, 500 µL of culture medium containing 1% B27/N2, 1% FBS, EGF (50 ng/mL), Noggin (100 ng/mL), R-spondin 1 (500 ng/mL), and Y-27632 (10 µM) was added. Cell culture medium was changed every other day. Spheroids were counted after 14 days. The results were expressed as average number of spheroids. The PGE<sub>2</sub> levels in culture supernatant in treated and untreated wells were estimated using PGE<sub>2</sub> ELISA kits (Cayman Chemical). The results were presented in pg/mL.

**MTT Assay**

MTT assay was performed to assess the cytotoxic effect of compounds (LFA-9 and licofelone) on normal colonocytes of rats. Cells were treated with compounds at various concentrations (0.1–100 µM) for 24 hours. Then, 20 µL of freshly prepared MTT reagent was added to cells and incubated at 37°C for 2 hours. The supernatant growth medium was removed and replaced with DMSO (100 µL) to dissolve the formazan crystals. The optical density (absorbance) was read using a microplate reader at 570 nm with a reference wavelength of 620 nm. Three individual experiments were performed and results were expressed as percent viability at various doses of each compound.

**In vivo Anti-Inflammatory Activity by Carrageenan-Induced Rat Paw Edema Model (Animal Studies)**

The carrageenan-induced paw edema model is the most widely used for the evaluation of anti-inflammatory activity. Male Fischer (F344) rats weighing 350–400 g were housed under standard conditions (temperature of 21°C with an alternating 12-h light-dark cycle and humidity of 50% and 20 air changes per hour, before and also during the experiment. The animals were fed with American Institute of Nutrition (AIN) modified-76A standard laboratory diet. During the experiment, the rats were allowed water and food ad libitum. Animal experimental protocol was approved by Institutional Animal Care and
Use Committee at the University of Oklahoma Health Sciences Centre (Protocol # 18–104-FH). Animal experiments were conducted according to the Guide for the Care and Use of Laboratory Animals (National Research Council of the National Academies) and as per the Animal Welfare Act39 (https://www.nal.usda.gov/awic/animal-welfare-act).

The animals were divided into three groups: Group I served as control (carrageenan), group II (saline), group III (LFA-9400 ppm) and group IV (licofelone 400 ppm). The animals were fed with AIN-76A diet containing control or LFA-9/licofelone (400 ppm) and injected (subcutaneous) 0.1 mL of 1% λ-carrageenan solution with saline into sub-plantar region of left hind paw of each rat. The right hind paw of the same rat was treated with 0.1 mL of saline alone in the same manner as the control. Before induction of edema, the dorsiventral thickness (millimeters (mm)) of both the paws of each was measured using digital vernier caliper and also the measurements were taken at 1-, 2-, 3-, 4-, 5- and 6-hours after carrageenan injection. The results are expressed in percent edema inhibition. The stomach and other major organs were examined for ulcers and any other toxicity.40 The paw exudates were collected and used for PGE2, LTB4, TXB2 and PGI2 quantifications using ELISA kits (Cayman Chemical, MI, USA).

Immunofluorescence Technique

CD68-positive inflammatory cell infiltration and TNF-α expression were evaluated using immunofluorescence technique on the paw tissue sections.38 Heat-induced epitope retrieval was performed on 4-μm formalin-fixed, paraffin-embedded sections in citrate buffer (pH 6.0) at 99°C for 30 minutes. Slides were incubated in a normal serum and BSA blocking step at room temperature for 40 minutes. After incubation with primary antibody [Alexa Fluor 700-conjugated CD68 (1:50 dilution) (Novus Biologicals (NB600985)) and TNF-α (1:100 dilution) (Santa Cruz)] overnight at 4°C, slides were labeled with Alexa Fluor 488 dye-conjugated secondary antibody (for TNF-α) and mounted with ProLong Gold (Thermo Fisher Scientific). Further, the slides were examined utilizing SP8 confocal microscopy (Leica). MetaMorph Microscopy Automation and Image Analysis Software was used to quantify the cell number with fluorescent staining.

Immunoblot Analysis

Twenty-micrograms of the total protein was size-separated in a 12% SDS-polyacrylamide gel and transferred electrophoretically onto a PVDF membrane using a semi-dry blot transfer apparatus (Bio-Rad, Hercules, CA, USA). Further, the membrane was blocked and incubated overnight with a primary antibody (CD68 (1: 100 dilution) (Santa Cruz, Dallas, TX, USA) and TNF-α (1:100 dilution)) and was subsequently incubated with horseradish peroxidase-conjugated secondary antibody. The proteins were detected using ECL reagents (Biorad). Beta-actin was used as the loading control.38

Statistical Analysis

Each experiment was conducted in triplicate and expressed as mean ± standard deviation. Inhibitory Concentration (IC50) values were calculated based on the dose-response curve vs log inhibitor concentration through variable slope (four parameters) of non-linear regression analysis using GraphPad prism software. Statistical differences between control and treated groups were evaluated using unpaired t test with Welch’s correction. P-values ≤0.05 for data sets were considered significant. Significance of the results are depicted in all graphs and tables as not significant (NS), (P>0.05); *(P ≤ 0.05); **(P ≤ 0.01); *** (P ≤ 0.001); **** (P ≤ 0.0001).

Results

In silico Screening for Identification of mPGES-1 and 5-LOX Inhibitor LFA-9

A hit library of mPGES-1 inhibitors and its active fragments (~1000) was prepared from BRENDA and NCI databases based on Lipinski’s rule of 3. To remove possible false hits, an online program PAINS Remover (pan-assay interfering compound substructures http://cbligand.org/PAINS) was used to screen all the designed licofelone analogues and found that all these analogues passed this test (Supplemental Information Table 1 and 2).41,42 The licofelone pharmacophore (the lead molecule) was optimized at its carboxylic group using different hit fragments through rational drug design strategy to target mPGES-1/5-LOX while sparing COX-1/2 activities of human, mouse, and rat (Supplemental Information Table 1 and 2). The licofelone derivatives were designed to target mPGES-1 and 5-LOX of human, mouse, and rat for their pre-clinical development and clinical translation. Among all designed licofelone derivatives (Supplemental Information Table 2), the addition of 2-amino acetic acid (glycine) at the carboxyl moiety of licofelone (a lead molecule LFA-9) led to strong binding interactions with
The active site of human, mouse, and rat mPGES-1 and 5-LOX, but exhibited poor interactions with COX-1/2 (Figure 2). Moreover, LFA-9 obeys Lipinski’s rule of 5 for drug-likeness, but licofelone does not (Supplemental Information Table 3). The calculated LogP value of licofelone is 6.15 (more than 5).

LFA-9 strongly bound with human, mouse, and rat-specific mPGES-1 with binding energies ranging between −238.27 and −257.76 Kcal/mol. LFA-9 showed strong interactions with the key amino acid residues of the mPGES-1 active site (Figure 2A; Table 1). mPGES-1 (trimer) forms active site pockets with polar residues like Arg126 for conversion of PGH₂ to PGE₂.⁴³ LFA-9 formed hydrogen bond interactions with Asn74 and Arg126 of the mPGES-1. The extended -COOH group lies within proximity of Asn74 which acts as an h-bond donor and forms conventional hydrogen bond interactions with one of the oxygens (=O) of the carboxylic acid moiety. Arg126 forms the hydrogen bond interactions with oxygen atom of the glycine residue. This interaction is very important as Arg126 is reported to be crucial for the catalysis of mPGES-1, and its mutation caused loss in mPGES-1 enzymatic activity.⁴³⁻⁴⁵

Apart from this a pi-donor hydrogen bond was also observed with Thr131 (similar to licofelone). Another crucial observation in the case of LFA-9 was the hydrophobic interactions formed with Tyr130. The Tyr130 formed strong pi-pi T-shaped interactions with the central 5-membered ring along with pi-alkyl interactions. Other hydrophobic observations included alkyl with Pro124 and Val128. These amino acids are key for mPGES-1 catalytic activity in a previous study by Gupta et al.⁴⁶ An atomic level interaction study demonstrated that LFA-9 formed hydrogen bonds and hydrophobic interactions with active site of amino acids mPGES-1 after inclusion of 2-aminoacetic acid at carboxyl end of the licofelone. The addition of 2-aminoacetic acid (glycine) before the terminal -COOH group showed an augmentation in the nature and strength of the interactions observed at the mPGES-1 active site. LFA-9 showed similar interactions with human, mouse and rat mPGES-1. These interaction analysis results clearly show the reason for improved mPGES-1 inhibitory efficacy of LFA-9 as compared to licofelone (Figure 2A; Table 1).

LFA-9 bound with 5-LOX (human/mouse/rat) with binding energies ranging between −199.85 and −215.3 Kcal/mol, which is comparable to that of licofelone. LFA-9 interacted with key active site amino acid Tyr181 of the 5-LOX and interferes with the iron that is coordinated by three conserved histidine residues (His 367, 372, and 550), as well as the main-chain carboxylate of the C terminus (Ile673) as part of the active site.⁴⁷ LFA-9 and licofelone are interacting in the similar vicinity of 5-LOX. Addition of 2-aminoacetic acid before the terminal -COOH group of licofelone (LFA-9) retains its 5-LOX inhibitory potential (Figure 2B). LFA-9 also interacts with His432 that helps to define the active site of 5-LOX⁴⁷ (Figure 2B). However, LFA-9 shows poor binding interactions with COX-1 and COX-2 (Figure 2C and D). Licofelone interacts with key active site amino acids of COX-1 (Arg119) and COX-2 (Ser530)⁴⁸⁻⁴⁹ (Figure 2C and D). The addition of 2-aminoacetic acid before the terminal -COOH group of licofelone (LFA-9) led to loss of these interactions with COX-1/2 (sparring effect).

Chemical Synthesis and Characterization of mPGES-1 and 5-LOX Inhibitor LFA-9

LFA-9 (2-[2-[2-(4-chlorophenyl)-6,6-dimethyl-1-phenyl-5,7-dihydropyrrolizin-3-yl]acetyl] amino)acetic acid) was synthesized in a two-step organic synthesis process and purity was found to be >95% (Supplemental Figure 5A and B). The retention time of LFA-9 is 6.7 minutes in HPLC analysis. Chemical structure of LFA-9 was assigned using advanced spectroscopic techniques, such as ¹H, ¹³C, and HRMS (Supplemental Figure 5C and D).

Yield: 80.5%; Melting Point: 106–108°C.; ¹H NMR (300 MHz, CDCl₃) δ (ppm): 1.29 (s, 6H, 2×CH₃), 2.85 (s, 2H, CH₂), 3.55 (s, 2H, CH₂), 3.72 (s, 2H, CH₂), 4.04 (d, Jgem=5.3 Hz, 2H, NH-CH₂), 7.03–7.29 (m, 9H, Ar-H). ¹³C NMR (75 MHz, CDCl₃) δ (ppm): 28.0, 33.0, 40.6, 41.2, 43.3, 58.0, 115.0, 117.2, 124.3, 124.8, 2×128.0, 128.5, 131.2, 131.9, 134.1, 134.9, 135.5, 171.4, 172.9; HRMS: m/z 437.1680 [M +H⁺]]; 459.1508 [M+Na⁺], calc'd. for 436.1680, found 437.1680.

5-LOX and mPGES-1 Dual Inhibitory and COX-1/2 Sparing Effects of LFA-9 in Cell-Free Systems

In a cell-free system, LFA-9 (Figure 3) substantially inhibited human, mouse, and rat mPGES-1 activities (IC₅₀ values: 0.87 µM [human]; 0.52 µM [mouse]; 1.40 µM [rat]), as compared with its parental compound licofelone (IC₅₀ values: 14.04 µM [human]; >15 µM [mouse]; 12.3 µM [rat]). LFA-9 showed comparable 5-LOX inhibitory activity (IC₅₀ values: 2.75 µM [human];
Licofelone inhibited COX-1 (IC$_{50}$ values: 1.65 µM [human]; 1.74 µM [mouse]; 1.81 µM [rat]) and COX-2 (IC$_{50}$ values: 1.36 µM [human]; 2.01 µM [mouse; 3.26 µM [rat]), but LFA-9 did not show inhibition of human, mouse, or rat COX-(1and2) activity up to the tested range of 1 mM (Figure 3).
LFA-9 Shows Strong Binding Affinity Causing Unfolding of the Secondary Structures of Human mPGES-1/5-LOX

To validate results from molecular docking studies, binding studies of LFA-9 and mPGES-1 and 5-LOX were performed using ITC, and found strong binding affinity based on accurate fitting of the isotherm. The KD (dissociation constant) values of LFA-9 for human and rat mPGES-1 were 2.1 and 3.3 µM, respectively (Table 2, Supplemental Figure 6). Human 5-LOX had two KDs, 18 µM and 210 µM, indicating two binding sites (Table 2, Supplemental Figure 6). This finding provides evidence that LFA-9 interacts with mPGES-1 at one binding site, while LFA-9 interacts with the 5-LOX enzyme at two binding sites. Further, the effect of LFA-9 on the secondary structure of mPGES-1, COX-1 and 2, and 5-LOX was evaluated using the circular dichroism (CD) technique. The interaction of LFA-9 (selected dose at 0.12 µM) with the secondary structure of mPGES-1 was determined based on the shift in bands at 208 nm and 220 nm of the CD spectrum, which is indicative of a 19% loss of α-helix content (Table 3, Supplemental Figure 7). This result suggested that the decrease in α-helix content of mPGES-1 led to its partial unfolding, which resulted in inhibition of its enzyme activity. LFA-9 binds strongly and unfolds the α-helix (8.7%) and β-sheets (1%) of 5-LOX at 0.1 µM, thereby inhibiting 5-LOX activity (Table 3, Supplemental Figure 7).

LFA-9 Inhibits Inflammmogen (Carrageenan)-Induced Hind Paw Edema of Rats

In vivo oral anti-inflammatory efficacy of LFA-9 (400 ppm) was tested on a group of rats (n=3) using carrageenan model and found 70.4% edema inhibition (maximum) at 3 hours after carrageenan injection, whereas licofelone (400 ppm) showed 61.7% edema inhibition (Figure 5A and B). LFA-9 (400 ppm) did not show any gastrointestinal toxicity (Supplemental Figure 8; Supplemental Information Table 4). LFA-9 reduced PGE2 (P ≤ 0.0001), and LTB4 (P ≤ 0.01) levels but spared PGI2 in paw exudates compared to control (vehicle), while, licofelone reduced PGE2, TXB2, LTB4, and PGI2 levels. LFA-9 maintained homeostasis of TXB2 and PGI2 production levels as that of control (saline). This study demonstrates the anti-inflammatory potential of LFA-9 by inhibiting production of PGE2 and LTB4, while sparing PGI2 levels (Figure 5C). Further, immunofluorescence and immunoblot analysis of paw tissue sections demonstrated that CD68-positive inflammatory cell infiltration and expression of TNF-α are significantly decreased in the LFA-9-treated group compared to the untreated control (Figure 5D–F). LFA-9 showed better efficacy in the reduction of CD68-positive inflammatory cell infiltration (P ≤ 0.0001) and suppression of TNF-α expression than licofelone (Figure 5D–F).

### Table 1 Binding Energies of LFA-9 on mPGES-1, COX-1/2, and 5-LOX

| Enzyme | Species | Binding Energy (Kcal/mol) |
|--------|---------|--------------------------|
|        |         | Licoferone | LFA-9   |
| 5-LOX  | Human   | −208.58     | −199.85 |
|        | Rat     | −212.36     | −209.8  |
|        | Mouse   | −216.74     | −215.3  |
| COX-1  | Human   | −198.37     | −56.91  |
|        | Rat     | −192.52     | −66.28  |
|        | Mouse   | −188.56     | −63.33  |
| COX-2  | Human   | −213.2      | −85.31  |
|        | Rat     | −205.37     | −72.42  |
|        | Mouse   | −211.53     | −66.54  |
| mPGES-1| Human   | −104.35     | −245.34 |
|        | Rat     | −135.25     | −257.76 |
|        | Mouse   | −142.63     | −238.27 |

PGE2 and LTB4 Inhibitory, and PGL2 Sparing Effects of LFA-9 in a Human/Mouse/Rat Cell-Based System

In cell-based assays (Figure 4), the effect of LFA-9 on PGE2, LTB4, and PGL2 production was evaluated in LPS-stimulated macrophages of humans, mice, and rats, and compared with that of licofelone. LFA-9 inhibited PGE2 (IC50 range 0.47–0.78 µM) and LTB4 (IC50 range 0.63–1.24 µM) production and spared PGL2 (IC50 >100) in LPS-stimulated macrophages of humans, mice, and rats, whereas licofelone inhibited production of PGE2 (IC50 range 0.56–1.31 µM), LTB4 (IC50 range 0.53–1.6 µM), and PGL2 (IC50 range 0.85–0.91 µM). Unlike licofelone, LFA-9 effectively inhibited PGE2 and LTB4 production in LPS-stimulated macrophages of humans, mice, and rats, with no effect on PGL2 production (Figure 4).
Figure 3 mPGES-1 and 5-LOX dual inhibitory activity of LFA-9 in a cell-free system. (A). Effect of LFA-9 and licofelone on human (i), rat (ii), mouse (iii) specific mPGES-1 (A), 5-LOX (B), COX-1 (C), COX-2 (D) and on an in vitro cell-free system. IC\textsubscript{50} values were calculated based on the dose-response curve vs log inhibitor concentration though variable slope (four parameters) of non-linear regression analysis.
Table 2 Binding Efficacy Representing KD Values of LFA-9 on Human mPGES-1, COX-1/2, and 5-LOX

| Type of Enzyme                  | KD          |
|--------------------------------|-------------|
| Human 5-LOX (sequential binding sites) | 18 µM (KD1) |
|                                | 210 µM (KD2)|
| Human COX-2                    | >1 mM       |
| Human COX-1                    | >1 mM       |
| Human mPGES-1                  | 2.1 µM      |
| Rat mPGES-1                    | 3.3 µM      |

LFA-9 Prevents Colorectal Tumor Stemness and Spheroid Formation in vitro

LFA-9 significantly inhibited 60.2% of colorectal tumor spheroids at 25 µM as compared with the untreated control, whereas licofelone inhibited only 22.3% of colonic spheroids at 25 µM (Figure 6A). LFA-9 inhibited 82% of PGE2 production as compared with the untreated control, whereas licofelone inhibited 63% of PGE2 production (Figure 6B). Moreover, LFA-9 showed no cytotoxicity to normal rat colonic epithelial cells (Supplemental Information Figure 9). LFA-9 exhibited better efficacy in prevention of colon tumor stemness than licofelone.

LFA-9 Metabolically Stable in Activated-S9 Fraction of Liver

LFA-9 is metabolically stable in activated-S9 fraction of mouse/rat liver for the tested time (1 hour). HPLC analysis demonstrated that 98.3% and 96.1% of LFA-9 remains after 60-minutes incubation in rat and mouse liver S9 fraction, respectively (Supplemental Information Figure 10).

Discussion

Pharmacological inhibitors of mPGES-1/5-LOX are proposed as alternative and safer anti-inflammatory agents to NSAIDs and COXibs.5-7 Licofelone is a potent inhibitor of COX-1/2, 5-LOX, and weak inhibitor of mPGES-1 with anti-inflammatory efficacy, but did not enter into the market due to COX-associated toxicities.14,15 Licofelone is an excellent scaffold for mPGES-1/5-LOX dual inhibitors development with COX-1/2 sparing effects.8 In this study, pharmacophore of licofelone was modified at its carboxylic end by addition of various active fragments of mPGES-1 inhibitors to enhance mPGES-1 inhibitory potential and spares COX-1/2 activity. Interestingly, the addition of 2-amino acetic acid (glycine) at carboxyl end of licofelone led to enhanced interactions with active site of mPGES-1 and loss of interactions with active sites of COX-1/2, but no significant change in its interactions with active site of 5-LOX. In vitro and in silico studies demonstrated that licofelone analog LFA-9 showed substantial 5-LOX and mPGES-1 dual inhibitory activity and spared COX-1/2 activities in human, mouse, and rat-based systems. LFA-9 showed several folds higher mPGES-1 inhibitory activity than did its parental molecule licofelone. In structure activity relationship studies demonstrated that addition of 2-amino acetic acid at the carboxyl end of licofelone improved inhibition of mPGES-1 and 5-LOX, and spared COX-1/2 activity. CD and ITC-based studies demonstrated that LFA-9 binds and induces changes in the secondary structure of human mPGES-1, thereby inhibiting their enzyme activities, based on in vitro studies. Using LPS-stimulated human/mouse/rat macrophages as the cell-based system, LFA-9 inhibited PGE2 and LTB4 due to mPGES-1/5-LOX dual inhibitory efficacy, and spared PGL2 production due to its COX-1/2 sparing effect. Moreover, LFA-9 was more effective in a cell-based system than a cell-free system. Results from in silico, in vitro, and ex vivo studies of LFA-9 that were performed using human/mouse/rat mPGES-1 and 5-LOX are useful to facilitate its pre-clinical development and clinical translation. Moreover, LFA-9 is metabolically stable in activated rat/mice hepatic S9 fraction in vitro for 60 minutes. Lipinski’s rule

Table 3 Effect of LFA-9 on Secondary Structure of Human mPGES-1 and 5-LOX

| Enzyme            | Concentration of LFA-9 (µM) | Change in Secondary Structure                      |
|-------------------|-----------------------------|---------------------------------------------------|
| Human mPGES-1     | 0.12                        | 19% unfolding in α-helix content                   |
| Human COX-1       | 10                          | No change in secondary structure                   |
| Human COX-2       | 10                          | No change in secondary structure                   |
| Human 5-LOX       | 0.12                        | Unfolding in (8.7%) α-helix and (1%) β-sheets     |
Gupta et al. described the importance of LogP and Nrotb along with dock scores in the discovery and development of mPGES-1 inhibitors. Unlike licofelone, LFA-9 obeyed Lipinski’s rule of 5 and number of rotatable bonds for drug-likeness (Supplemental Information Table 3). Thus, LFA-9 may show better pharmacokinetic and dynamic properties than does licofelone.33,34
In the pre-clinical study using an inflammatory animal model, LFA-9 showed better anti-inflammatory activity than that of licofelone in carrageenan (inflammogen)-induced rat paw edema model. LFA-9 showed maximum edema inhibition at 3 hours after carrageenan injection, which is referred to as the second phase of acute inflammation, in which CD68-positive inflammatory cells are accumulated, pro-inflammatory enzymes of arachidonic acid pathways and TNF-α are over-expressed. Results also demonstrate that LFA-9 inhibited inflammatory response by inhibiting pro-inflammatory PGE₂ and chemotactic LTß₄ levels and thereby, preventing the infiltration of CD68 positive macrophages and granular neutrophils along with TNF-α expression (Figure 5G).

NSAIDs induce gastric damage (ulcers and erosions) due to COX-1/2 inhibition, while mPGES-1 inhibitors show gastric mucosal safety due to COXs sparing effect. Chattopadhyay et al. found that aspirin (180 mg/kg) caused gastric toxicity with potential anti-inflammatory efficacy. Ding et al. found gastric toxicity of celecoxib and developed human/mouse specific mPGES-1 inhibitory benzylidenelbarbituric acid derivatives as anti-inflammatory agents with GI safety. LFA-9 is a human/rat/mouse-specific mPGES-1/5-LOX inhibitor and maintains homeostatic levels of TXB₂ and spared PGI₂ levels as equal to that of the healthy control group (saline treated). But, licofelone reduced the levels of TXB₂ and PGI₂ compared to that of the healthy control group (saline treated) (Figure 5G). LFA-9 (400 ppm) did not show any gastrointestinal and any other toxicities in the rats (Supplemental Figure 8; Supplemental Information Table 4). Therefore, the results of the study demonstrated that LFA-9 can be a safer anti-inflammatory agent and warrants detailed studies for its clinical translation.

A pro-inflammation mediator, PGE₂, promotes tumor stemness and spheroid formation towards colonic tumor initiation and progression. Three-dimensional tumor spheroids in vitro mimic in vivo solid colonic tumors for anticancer drug screening. LFA-9 inhibited colonic tumor spheroid formation of APC(nec) rats (a model mimics colonic familial polyposis formation of humans). It had been well established that PGE₂ production in the microenvironment regulates colon tumor stemness during tumorigenesis, progression, and invasion. Therefore, this study demonstrated the cancer-preventive
effects of LFA-9 against tumorigenesis of CRC via its PGE₂ inhibitory efficacy (Figure 6C). These results support our other study observations of LFA-9 inhibitory effects on APC<sup>min⁻/⁻</sup> mice intestinal polyposis.²⁵

**Conclusions**

In silico and in vitro (cell-free and cell-based) studies demonstrated that LFA-9 is a potent mPGES-1/5-LOX dual inhibitor by inhibiting products PGE₂ and LTB₄, respectively, and spares COXs and its downstream product PGJ₂. LFA-9 showed better druggable properties than did licofelone. In vivo studies demonstrated that LFA-9 showed substantial anti-inflammatory activity by inhibiting inflammogen (carrageenan)-induced paw edema in rats. LFA-9 prevented colon tumor stemness and spheroids formation in vitro, mediated by its PGE₂ inhibitory effects. Therefore, mPGES-1/5-LOX inhibitor LFA-9 may be a new generation anti-inflammatory agent and the results warrant further pre-clinical studies on LFA-9 against several inflammatory diseases, including cancer.

**Abbreviations**

LFA-9, licofelone analogue-9; COX, cyclooxygenase; 5-LOX, 5-lipoxygenase; mPGES-1, microsomal prostaglandin E synthase-1; CD, circular dichroism; IC₅₀, inhibitory concentration 50; ITC, isothermal titration calorimetry; LTs, leukotrienes; PGJ₂, prostacyclin; TXB₂, thromboxane B₂; PGE₂, prostaglandin E₂; CAR, carrageenan.
Data Sharing Statement
The data sets and supporting data are available from the corresponding author upon reasonable request.

Ethics Approval and Consent
Animal experimental protocol (Protocol # 18-104-FH) was approved by Institutional Animal Care and Use Committee at the University of Oklahoma Health Sciences Center.

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Disclosure
Gopal Pathuri reports a patent US10206904B2 issued and a patent WO2017/015013 pending. Hariprasad Gali reports a patent US Patent 10206904 issued. Chinthalapally V. Rao reports a patent US10206904B2 issued and a patent WO2017/015013 A1 pending. The authors report no other potential conflicts of interest for this work.

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