Identification of Farnesyl Pyrophosphate and N-Arachidonylglycine as Endogenous Ligands for GPR92*

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A series of small compounds acting at the orphan G protein-coupled receptor GPR92 were screened using a signaling pathway-specific reporter assay system. Lipid-derived molecules including farnesyl pyrophosphate (FPP), N-arachidonylglycine (NAG), and lysophosphatidic acid were found to activate GPR92. FPP and lysophosphatidic acid were able to activate both Gq/11- and Gi-mediated signaling pathways, whereas NAG activated only the Gq/11-mediated signaling pathway. Computational modeling combined with site-directed mutagenesis of GPR92 indicated that Thr97, Gly98, Phe101, and Arg267 of GPR92 are responsible for the interaction of GPP with FPP and NAG. Reverse transcription-PCR analysis revealed that GPR92 mRNA is highly expressed in the dorsal root ganglia (DRG) but faint in other brain regions. Peripheral tissues including spleen, stomach, small intestine, and kidney also expressed GPR92 mRNA. Immunohistochemical analysis revealed that GPR92 is largely co-localized with TRPV1, a nonspecific cation channel that responds to noxious heat, in mouse and human DRG. FPP and NAG increased intracellular Ca2+ levels in cultured DRG neurons. These results suggest that FPP and NAG play a role in the sensory nervous system through activation of GPR92.

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that GPR92 has only a 35% amino acid homology with the LPA4 receptor. Additionally EC_{50} values for various LPA compounds in the production of cAMP and IP in GPR92-expressing cells range between 0.5 and 5 μM. This indicates that LPA is a moderately potent activator of GPR92 (6, 8). More recently, a luminal protein hydrolysat was found to activate GPR92 and possess a higher potency for cAMP production than LPA (9). These findings suggest the possibility that molecules other than LPA can serve as ligands for GPR92 and that perhaps reference to this compound as the LPA_{4} receptor would be premature.

While searching for orphan GPCR ligands using various chemical compounds, we found that many lipid-derived molecules are able to activate GPR92 with different potencies. Of these, farnesyl pyrophosphate (FPP) was more potent than LPA in activating GPR92 as revealed by E_{max} and EC_{50} values. N-Arachidonylglycine (NAG) was as potent as LPA in activation of GPR92. FPP and LPA activated both G_{s} and G_{q/11} linked signaling pathways, whereas NAG was able to activate only the G_{q/11}-linked signaling pathway. Using computer-simulated molecular modeling combined with site-directed mutagenesis, we identified amino acid residues responsible for the interaction of GPR92 with FPP and NAG. Reverse transcription-PCR analysis revealed that GPR92 mRNA was present in high levels in the dorsal root ganglia (DRG) in the nervous system as well as in the spleen, stomach, intestine, and kidney. Further FPP and NAG induced Ca^{2+} mobilization in cultured DRG cells.

**EXPERIMENTAL PROCEDURES**

**Chemicals**—The chemicals 12-O-tetradecanoylphorbol-13-acetate and FPP were purchased from Sigma. U73122 and MDL-12330A were obtained from Calbiochem. We obtained 2-arachidonoylglycerol (2-AG) from Cayman Chemical (Ann Arbor, MI). LPA (14:0) was purchased from Avanti Polar Lipids (Alabaster, AL). All acylamides, including NAG, N-arachidonylglycerol, N-palmitoylsphingosine phosphoric acid, N-arachidonoylanamine, and N-arachidonylether, were synthesized as described previously (10).

**Plasmids**—The cDNA for human GPR92 was constructed at HindIII and XbaI sites of pcDNA3, whereas cDNA for mouse GPR92 was constructed at EcoRI and XhoI of pcDNA3 (Invitrogen). Site-directed mutagenesis of human GPR92 was generated by the PCR overlapping extension method (11). All PCR-derives oligonucleotides were verified by automated sequencing. pCMV β-Gal was purchased from Clontech. SRE-luc, containing a single copy of the serum response element (SRE; CCATATTAGG) followed by a c-fos basic promoter and luciferase, was constructed. Adenoviruses containing human GPR92 or an SRE-luc reporter were obtained from Neugex Co. (Seoul, Korea).

**Adenovirus Infection, Transfection, and Luciferase Assay**—Adenovirus infection and luciferase assays were performed as described previously (4). CV-1 cells were maintained in DMEM in the presence of 10% fetal bovine serum. Cells (5 × 10^{7}) were plated into a 96-well plate. Cells were coinfected with adenoviruses containing human GPR92 and an SRE-luc reporter with 50 multiplicity of infection (m.o.i.) under serum-free conditions for 3 h. Cells were then incubated with 10% fetal bovine serum-containing DMEM.

For SRE-luc analysis, cells were maintained in serum-free DMEM for at least 16 h before treatment. Following treatment of cells with an agonist for 6 h, cells were harvested. Luciferase activity in the cell extracts was determined according to standard methods using a Wallac 1420 VICTOR3 plate reader (PerkinElmer Life Sciences). For transfection of GPR92 mutants, CV-1 cells were plated into 24-well plates and transfected 24 h later with Effectene reagent (Qiagen, Valencia, CA). The total amount of DNA used in each transfection was adjusted to 1 μg by adding appropriate amounts of pcDNA3. Forty-eight hours after transfection, cells were treated with ligands for 6 h.

**GTPγS Binding Assay**—Forty-eight hours after infection of CV-1 cells with adenoviruses containing human GPR92 or vehicle, cells were homogenized in 5 mm Tris-HCl, 2 mm EDTA (pH 7.4) and centrifuged at 48,000 × g for 15 min at 4 °C. The resulting pellets were washed in 50 mm Tris-HCl, 10 mm MgCl_{2} (pH 7.4) and stored at −80 °C until use. Agonist-stimulated [35S]GTPγS binding was determined as described previously (12). Briefly membrane fractions (15 μg) were incubated for 15 min at room temperature in binding buffer (20 mm HEPES (pH 7.4), 100 mm NaCl, 3 mm MgCl_{2}, and 3 μm GDP) in the presence of various concentrations of FPP, LPA, or NAG. [35S]GTPγS (0.2 nM) was added. Then samples were further incubated for 30 min at 30 °C. The incubation was stopped by centrifugation at 1000 × g for 10 min at 4 °C. Bound GTPγS [35S] was counted in a scintillation mixture. Nonspecific binding was determined in the presence of 10 μM GTPγS to be less than 10% of total binding.

**Measurement of Inositol Phosphate Production**—An IP production assay was performed as described previously (13). CV-1 cells were seeded into a 12-well plate and infected with adenoviruses containing human GPR92 with 100 m.o.i. After infection, cells were incubated in M199 medium (Invitrogen) in the presence of 1 μCi/ml [3H]inositol ([3H]inositol (Amersham Biosciences)/well) for 20 h. Medium was removed, and cells were washed with 0.5 ml Buffer A (140 mm NaCl, 20 mm HEPES, 4 mm KCl, 8 mm d-glucose, 1 mm MgCl_{2}, 1 mm CaCl_{2}, and 1 mg/ml fatty acid-free bovine serum albumin). Cells were then preincubated for 30 min with Buffer A containing 10 mm LiCl followed by addition of 1 μM FPP, 10 μM NAG, or 10 μM LPA at 37 °C for 30 min with or without 15-min prior treatment with a phospholipase C inhibitor, U73122. Replacing incubation medium with 0.5 ml of ice-cold 10 mm formic acid terminated the reaction. After 30 min at 4 °C, the formic acid extracts were transferred to columns containing Dowex anion-exchange resin (AG-1-X8 resin, Bio-Rad). Total IPs were then eluted with 1 ml of ammonium formate, 0.1 m formic acid. Radioactivity was determined using a Tri-Carb 3100TR scintillation counter (PerkinElmer Life Sciences).

**cAMP Assay**—Cyclic AMP levels were determined by measuring [3H]cAMP formation from [3H]ATP (13). Twenty-four hours before transfection, HeLa cells were seeded into 12-well plates. Cells were transfected with the pcDNA3-GPR92 plasmid with Effectene reagent. One day later, cells were labeled with 2 μCi/ml [3H]adenine (PerkinElmer Life Sciences) for
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24 h. Cells were first washed in phosphate-buffered saline and then incubated at 37 °C for 20 min in serum-free DMEM containing 1 mM 1-methyl-3-isobutylxanthine. Cells were stimulated with 1 μM FPP, 10 μM NAG, or 10 μM LPA for 30 min with or without 15-min prior treatment with an adenylate cyclase inhibitor, MDL-12330A. The reactions were terminated by replacing the medium with ice-cold 5% trichloroacetic acid containing 1 mM ATP and 1 mM cAMP. [3H]cAMP and [3H]ATP were separated on AG 50W-X4 resin (Bio-Rad) and alumina columns as described previously (13). The cAMP accumulation was expressed as [3H]cAMP/[3H]ATP × 1000.

Ca2+ Mobilization Assay—F11 rat embryonic neuroblastoma × DRG neuron hybrid cells (generous gifts from Dr. Henning Otto, Freie Universitat, Berlin, Germany) were transiently transfected with hGPR92-GFP and grown on poly-D-lysine-coated glass coverslips for 24–48 h. Cells were then incubated in a physiological solution (138 mM NaCl, 6 mM KCl, 1 mM MgSO4, 2 mM CaCl2, 1 mM NaH2PO4, 5 mM NaHCO3, 5 mM glucose, 10 mM HEPES, and 0.1% bovine serum albumin) with 5 μM fura-2/AM (Molecular Probes, Eugene, OR) at room temperature for 45 min. Cells were washed twice with the dye-free physiological solution and mounted in an acrylamide chamber that allows for perfusion of incubation medium. Fura-2/AM fluorescence was measured in GFP-positive cells at excitation wavelengths 340 and 380 nm and at emission wavelength 510 nm using an IX70 fluorescence microscope (Olympus, Tokyo, Japan) coupled to a digital cooled CCD camera (CoolSNAP fx CCD camera, Roper Scientific, Tucson, AZ).

ERK Activation—CV-1 cells (2 × 106) were plated on a 60-mm dish for 1 day before infection with adenoviruses containing GPR92. Cells were infected with 100 m.o.I. adenoviruses for 3 h in serum-free DMEM. Then the medium was changed to 10% fetal bovine serum-containing DMEM. Before protein preparation for ERK Western blotting, cells were incubated for at least 16 h in serum-free medium prior to agonist stimulation. After agonist stimulation in the presence or absence of 100 ng/ml pertussis toxin, cells were lysed by RIPA buffer (50 mM Tris HCl (pH 8.0), 150 mM NaCl, 1% Nonidet P-40, 0.1% SDS, 1 mM EDTA, and protease inhibitor mixture) followed by boiling at 95 °C for 5 min. Equal amounts of cellular extracts were separated on 10% polyacrylamide gels and transferred to nitrocellulose membranes for immunoblotting. Phosphorylated ERK1/2 and total ERK1/2 were detected by immunoblotting with mouse monoclonal anti-phospho-p44/42 MAPK (Cell Signaling Technology, 1:2000) and anti-p44/42 MAPK (Cell Signaling Technology, 1:2000), respectively. Chemiluminescence detection was performed using the ECL reagent.

Molecular Modeling—A homology model of GPR92 was built on the basis of the 2.2-Å crystal structure (Protein Data Bank code 1U19) of the bovine rhodopsin (14) using the homology modeling program molIDE (15). Hydrogen atoms and Kollman-all charges were added to the homology model of GPR92 using Sybyl v7.0 (Tripos Inc., St. Louis, MO). The three-dimensional structure of FPP and NAG was sketched and refined using the Sybyl program with the Gasteiger-Huckel charge method. The virtual docking of clomipramine was performed using GOLD v3.0, a program that applies stochastic genetic algorithms for conformational searching (16). The number of genetic operations was set to 1 × 105, and the population size was set to 1 × 103. All structural figures were prepared using PyMol v0.98 (DeLano Scientific LLC, San Francisco, CA).

Reverse Transcription-PCR Analysis—Total RNA was isolated from mouse tissues using TRI Reagent (Sigma). Two micrograms of total RNA were reverse transcribed with Molenoy murine leukemia virus reverse transcriptase (Invitrogen) according to the manufacturer’s instructions. The cDNA was amplified by PCR using the following primers: mGPR92-F (5′-TTGCGAGATCTTCTGGACACT-3′; 681–701) and mGPR92-R (5′-GCCAAGGCTTGTTCAGCCG-3′; 989–1010). The PCR cycling parameters were as follows: denaturation at 95 °C for 5 min followed by 35 cycles of denaturation at 95 °C for 30 s, annealing at 57.5 °C for 30 s, and extension at 72 °C for 45 s. PCR products were separated on a 1.5% agarose gel, stained with ethidium bromide, and photographed under a UV light source.

Immunocytochemistry and Histochemistry—The mouse lumbar DRG was isolated from 6-week-old male C57B/6 mice. The human DRG was isolated from fresh cadavers with the informed consent of the relatives of body donors and the approval of the ethics committee of the Korea University College of Medicine, Seoul, Korea. The isolated DRG was fixed for 4 h in 4% paraformaldehyde, washed with 1× phosphate-buffered saline, and cryoprotected overnight in 30% sucrose. The DRG was serially cut (10 μm) on a cryostat and mounted on gelatin-coated slides. Slides were incubated in 1× phosphate-buffered saline containing 5% bovine serum albumin and 0.1% Triton X-100 for 1 h at room temperature. Sections were incubated overnight in primary antibody at 4 °C and washed three times with 1× phosphate-buffered saline. The primary antibodies used in this study are as follows: mouse GPR92 (1:100, MBL International, Woburn, MA), human GPR92 (1:1000, Abcam), mouse and human TRPV1 (1:1000, Abcam), and neurofilament 200 kDa (NF200; 1:1000, Chemicon, Temecula, CA). After several washes, sections were incubated with fluorescence-labeled secondary antibodies (anti-rabbit-Alexa594 antibody for human and mouse GPR92, anti-guinea pig-Alexa488 antibody for TRPV1, and anti-mouse-Alexa488 antibody for NF200, Molecular Probes), and nuclei were counterstained with Hoechst33342 (10 μM) using 5′,6-diamidino-2-phenylindole (DAPI).
μg/ml) for 30 min at room temperature. Fluorescence labeling was observed under an LSM510 confocal laser microscope (Zeiss). Images were optimized for quantification using Adobe Photoshop 7.0 (Adobe Systems, San Jose, CA).

**DRG Cultures and Intracellular Ca^{2+} Imaging**—Cultured DRG neurons were prepared using a technique modified from Rhim et al. (17). Briefly, DRG were isolated from all levels of the lumbar and sacral spinal cord from Sprague-Dawley rats (100–150 g) and incubated with 0.15% collagenase (Sigma) for 20 min and then with 0.125% trypsin (Sigma) for 10 min in Ca^{2+}- and Mg^{2+}-free HEPES buffer solution at 37 °C. DRG neurons were then mechanically dissociated with Pasteur pipettes by trituration and plated on poly-l-lysine-coated coverslips. Cells were maintained in DMEM supplemented with 10% fetal bovine serum, 1 mM sodium pyruvate, 100 units/ml penicillin, and 100 μg/ml streptomycin under a humidified atmosphere of 95% air, 5% CO_{2} at 37 °C. For intracellular Ca^{2+} imaging recordings, cells were used within 1–2 days after plating. The diameter of each DRG neuron was defined as the average of the distance along the longest and shortest axis of the cell body. All chemicals used for cell preparation were purchased from Invitrogen. For intracellular Ca^{2+} imaging, the acetoxyethylene ester form of fura-2 (fura-2/AM, Molecular Probes) was used as the fluorescent Ca^{2+} indicator. DRG neurons were incubated for 60 min at room temperature with 5 μM fura-2/AM and 0.001% Pluronic F-127 in a HEPES-buffered solution composed of 150 mM NaCl, 5 mM KCl, 1 mM MgCl_{2}, 2.5 mM CaCl_{2}, 10 mM HEPES, and 10 mM glucose (pH adjusted to 7.4 with NaOH). Cells were then washed with HEPES-buffered solution and placed on an inverted microscope (Olympus). Cells were illuminated using a xenon arc lamp. The required excitation wavelengths (340 and 380 nm) were selected with a computer-controlled filter wheel (Sutter Instruments, Novato, CA). Emitted fluorescence light was reflected through a 515-nm long pass filter to a frame transfer cooled CCD camera. The ratios of emitted fluorescence were calculated using a digital fluorescence analyzer. All imaging data were collected and analyzed using Meta Imaging Software (Molecular Devices Corp., Downingtown, PA).

**Cell Transfection**—DRG cells were transfected after 1 day in culture using Lipofectamine 2000. Briefly, GPR92 siRNA (5’-CGUUUGCAUAUGGUGUdTdT-3’) or a non-targeting scrambled siRNA (negative) and the Lipofectamine 2000 diluted in serum-free DMEM/F-12 were combined and incubated at room temperature for 20 min. After the culture media were removed and saved, cells were incubated with the siRNA/Lipofectamine 2000 mixture at 37 °C for 4 h. Cells were then incubated with the saved culture media for 48 h. To monitor transfection efficiency, the GFP expression vector pEGFP-N1 (Clontech) was co-transfected with each siRNA. For each Ca^{2+} imaging experiment, the transfected cells without the GFP vector were loaded with fura-2/AM after separately confirming transfection efficiency (>80%).

**Data Analysis**—All assays were performed in triplicate and repeated three times. The data are presented as mean ± S.E. of at least three independent experiments. Data analysis was performed using nonlinear regression. Data are expressed using sigmoidal dose-response curves. Agonist concentrations inducing half-maximal stimulation (EC_{50}) and maximal -fold increase (E_{max}) were calculated using the GraphPad PRISM4 software (GraphPad, San Diego, CA). One-way analysis of variance followed by Newman-Keuls post-test was used for data analyses. p < 0.05 was considered statistically significant.

**RESULTS**

**Ligand Screening for GPR92**—We previously established a widely applicable screening system to identify ligands for orphan GPCRs (4). In this system, cDNAs for orphan GPCR and GPCR signaling-specific reporter genes are introduced to adenovirus. These reporter genes include a CAMP-responsive element-driven luciferase gene (CRE-luc) for G_{s}- and G_{i/o}-linked pathways and SRE-luc for G_{q/11} or G_{12/13}-linked pathways (4, 13). We screened ligands for GPR92 using CV-1 cells that were doubly infected by adenoviruses containing human

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**TABLE 2**

| Ligands | IP production | cAMP production |
|---------|---------------|-----------------|
|         | E_{50} (μM)  | E_{max} (μM)    | E_{50} (μM) | E_{max} (μM) |
| FPP     | 0.38 ± 0.15  | 4.41 ± 0.56     | 1.46 ± 0.52 | 3.15 ± 0.22  |
| NAG     | >50          | 2.01 ± 0.25     | >50         | 2.44 ± 0.30  |
| LPA     | 1.03 ± 0.42  | 2.98 ± 0.19     | >50         | 2.44 ± 0.30  |

**FIGURE 1. Response of GPR92 to FPP and NAG.** A, CV-1 cells were coinfected with adenoviruses containing human GPR92 and SRE-luc in 96-well plates. Forty-eight hours after infection, cells were treated with the indicated concentration of FPP, NAG, or LPA for 6 h. Cells were harvested, and luciferase activities in the cell extracts were measured. The results are plotted as -fold activity over basal activity. Each bar represents means ± S.E. of three independent experiments performed in triplicate. B, FPP-, NAG-, and LPA-induced [35S]GTPγS binding to membrane fractions of CV-1 cells infected with adenoviruses containing GPR92. [35S]GTPγS binding was determined in a binding buffer containing 100 μM GDP in the presence of various concentrations of FPP, LPA, and NAG. The increase in [35S]GTPγS binding is indicated as the percentage of specific binding to basal binding. Data are expressed as mean ± S.E.
FIGURE 2. GPR92-mediated signaling pathways. A and C, IP production (A) or cAMP accumulation (C) in response to various concentrations of FPP, NAG, or LPA were determined in CV-1 cells infected with adenoviruses containing GPR92. B and D, the specificity of the signaling pathway was verified by pretreating with U73122 (10 μM), a phospholipase C inhibitor (B), and MDL-12330A (10 μM), an adenylate cyclase inhibitor (C), 15 min before treatment with 10 μM FPP, NAG, or LPA, respectively. a, p < 0.05 versus the control group (basal); b, p < 0.05 versus the agonist-stimulated group. E, effects of FPP and NAG on intracellular Ca^{2+} levels in GPR92-expressing cells were examined. Increased intracellular Ca^{2+} was found in F11 cells transiently expressing GPR92-GFP. Cells were treated with various concentrations of FPP and NAG. F, ERK1/2 activation by agonists in GPR92-infected cells. CV-1 cells were infected with 100 m.o.i. adenoviruses with GPR92 and were serum-starved for 16 h prior to agonist stimulation. The mock-infected cells (F, upper panel) were treated with 1 μM FPP, 10 μM of NAG and LPA, and 200 nm 12-O-tetradecanoylphorbol-13-acetate (TPA) as a positive control for 5 min. GPR92-infected CV-1 cells (F, middle panel) were treated with various concentrations of NAG and FPP. Involvement of G, activation in ERK1/2 phosphorylation was determined by pretreatment with 100 ng/ml pertussis toxin (PTX) (F, bottom panel). DP, double phosphorylated. The prime represents minutes. Data are expressed as the mean ± S.E. of three independent experiments.
In an initial screening, FPP, NAG, LPA, 2-AG, N-arachidonyl γ-aminobutyric acid, N-palmitoyleserine phosphoric acid, N-arachidonoylalanine, and N-arachidonoylserylserine were found to activate GPR92. Dose-response studies showed that FPP had the highest potency with an EC50 of 0.26 μM, approximately 10 times lower than that of LPA (Fig. 1A). NAG, 2-AG, N-arachidonyl γ-aminobutyric acid, and LPA showed EC50 values ranging from 2 to 5 μM (Table 1). Further FPP showed 3-fold higher efficacy than other lipid-derived molecules.

To examine whether the agonist-induced SRE-luc activity is mediated by G protein coupling, a [35S]GTPγS binding assay was performed using cells expressing GPR92 in the presence of various concentrations of FPP, LPA, or NAG. FPP dose-dependently increased [35S]GTPγS binding to cell fractions containing GPR92 with a higher potency than NAG and LPA (Fig. 1B). No agonist-induced increases in [35S]GTPγS binding were observed in mock-infected cells.

**Signaling Pathways of GPR92—** To investigate the signaling pathways of GPR92, we determined the levels of various second messengers after treatment with FPP, NAG, or LPA. Treatment of GPR92-expressing cells with these agonists resulted in a concentration-dependent increase in IP production. FPP showed the highest potency for IP production followed by LPA and NAG (Fig. 2A and Table 2). Agonist-induced production of IP was completely blocked by U73122, a phospholipase C inhibitor, indicating that GPR92 is Gq-coupled (Fig. 2C and Table 2). Agonist-induced increases in cAMP levels were suppressed by MDL-12330A, an adenylate cyclase inhibitor (Fig. 2D), indicating that GPR92 is able to couple with Gs. The fold induction of IP and cAMP by FPP was higher than the induction levels produced by LPA and NAG. To further determine the involvement of a cAMP-linked pathway, we performed an additional experiment using a CRE-luc reporter. The cells were coinfected with adenovirus containing GPR92 and CRE-luc and treated with various concentrations of FPP, LPA, and NAG. FPP and LPA increased CRE-luc reporter activity, whereas NAG failed to do so (supplemental Fig. 1A). H89, a cAMP-dependent protein kinase inhibitor, was able to inhibit both FPP- and LPA-induced CRE-luc activity (supplemental Fig. 1B). This study again demonstrated the involvement of the cAMP/cAMP-dependent protein kinase pathway in GPR92-mediated signaling.
We then examined whether FPP and NAG can increase intracellular Ca\(^{2+}\) levels in GPR92-expressing cells. F11 cells transfected with GFP-fused human GPR92 were treated with various concentrations of FPP or NAG. Dose-dependent inductions of intracellular Ca\(^{2+}\) concentration were observed only in GFP-positive cells (Fig. 2E).

Treatment of cells with FPP or NAG for 5 min elevated phospho-ERK levels (Fig. 2F, middle panel), whereas these agonists did not elevate phospho-ERK levels beyond basal amounts in mock-infected cells. A slight increase in phospho-ERK levels by LPA in mock-infected cells is likely due to expression of other types of LPA receptors in these cells (Fig. 2F, upper panel). Increased phospho-ERK levels were maintained for 15 min. Increases were not blocked by pertussis toxin, a specific inhibitor of G\(_i\) (Fig. 2F, lower panel), indicating that the increase in phospho-ERK level is independent of G\(_i\) activation.

**Identification of Ligand-binding Residues of GPR92**—Although FPP and NAG have different chemical structures, they are able to activate the same receptor, which provided the impetus to investigate the ligand-binding residues of GPR92 used in binding to different agonists. To address this question, we constructed a molecular model of GPR92 based on the rhodopsin structure and performed a virtual docking of the ligands to the receptor to predict the putative binding sites of GPR92 for FPP and NAG. Molecular modeling results revealed that the negatively charged diphosphate moiety of FPP has an electrostatic interaction with Arg276 in the third extracellular loop of GPR92 (Fig. 3A). Gly98 and Phe101 at transmembrane domain 3 have close contacts with the hydrophobic carbon chain of FPP (Fig. 3B). As Gly has the smallest side chain, it may provide a space for docking of the hydrophobic carbon chain of FPP. The glycine moiety of NAG interacts with Thr97 at transmembrane domain 3 (Fig. 3C). Like FPP, the hydrophobic carbon chain of NAG has close contacts with Gly98 and Phe101 (Fig. 3D).

Support for the modeling data was obtained by mutating these potential binding residues. EC\(_{50}\) and E\(_{\text{max}}\) values of FPP and NAG for wild type and mutant GPR92 are summarized in Table 3. Mutation of Thr\(^{97}\) to Ile (T97I), to Leu (T97L), and to Ala (T97A) significantly decreased response to NAG. However, these mutants responded to FPP with potency similar to that of the wild type.
type receptor (Fig. 4, A and B), indicating that Thr97 might be important for NAG interaction. Mutation of Gly98 to Phe (G98F) or to Lys (G98K) completely abolished responsiveness to both FPP and NAG (Fig. 4, A and B). Mutation of Gly98 to Ala (G98A) also reduced responsiveness to both FPP and NAG (supplemental Fig. 2, A and B). This result indicates that the small Gly98 may be important for forming a binding pocket. A tight steric hindrance caused by the bulky side chains of Phe and Lys may hamper binding to both FPP and NAG.

Mutation of Phe101 to Trp (F101W) drastically decreased sensitivity to NAG, whereas it reduced sensitivity to FPP to a level 10-fold lower than wild type (Fig. 4, C and D). Mutation of Phe101 to Ala (F101A), however, did not change sensitivity to either NAG or FPP. Replacement of Arg276 with Ser (R276S) completely abolished the FPP response but resulted in retention of responsiveness to NAG. Lys substitution (R276K) significantly decreased sensitivity to FPP and NAG. This finding suggests that the positively charged side chain of Arg276 may participate in an electrostatic interaction with the negatively charged diphosphate group of FPP.

To examine whether reduced responsiveness of mutant receptors to agonists is due to aberrant surface expression of the receptor, the subcellular localization of GFP-fused mutant receptors was determined. Like the wild type receptor, all mutant receptors were localized to plasma membrane (Fig. 4E). Further we examined protein expression level by Western blot using an anti-GFP antibody, showing no remarkable differences in expression levels except for G98F (24% of wild type expression) (Fig. 4F). This result together with the membrane expression of mutant receptors suggests that reduced or complete loss of responsiveness to agonists is not mainly due to aberrant or reduced expression of the mutant except for G98F.

Expression Pattern of GPR92 in the Mouse Tissues and Human DRG—Regional distribution of GPR92 mRNA in the mouse brain and peripheral tissues was assessed by reverse transcription-PCR analysis (Fig. 5A and supplemental Fig. 3A). GPR92 mRNA was expressed broadly at low levels throughout the brain regions and was expressed at relatively high levels in the DRG, spleen, stomach, small intestine, kidney, and thymus. GPR92 protein expression was further evaluated by immunohistochemical staining. To examine whether antibodies for GPR92 were appropriate for immunohistochemistry, we first performed immunocytochemistry in cells expressing mouse or human GPR92. Antibodies for mouse and human GPR92 specifically recognized mouse and human GPR92, respectively (Fig. 5B).

For DRG immunohistochemistry, GPR92 was doubly stained with the neuronal markers NF200 and TRPV1 to distinguish subgroups of DRG neurons. In both mouse and human DRG, the anti-NF200 antibody recognized large diameter neurons, whereas TRPV1 mainly stained medium and small diameter neurons. In the mouse DRG, GPR92 was faintly labeled in NF200-negative cells, whereas some signals were observed in NF200-positive cells. GPR92 was strongly labeled in TRPV1-positive cells (Fig. 5C). In the human DRG, GPR92 was distributed in small diameter neurons but rarely detected in NF200-positive neurons (Fig. 5D). Double staining of human DRG sections with GPR92 and TRPV1 revealed that the majority of TRPV1-positive neurons also expressed GPR92 (Fig. 5D). No cross-activity between GPR92 and TRPV1 antibodies was observed (supplemental Fig. 3, B and C).

FPP and NAG-induced Ca²⁺ Mobilization in Cultured DRG Neurons—We next examined the function of GPR92 by treating cultured rat DRG neurons with FPP or NAG. Agonist-induced Ca²⁺ responses were mainly observed in small diameter neurons. Some neurons responded to both FPP (1 μM) and NAG (10 μM), whereas others responded to only FPP or NAG (Fig. 6, A–D). Of the small diameter neurons tested (diameter <30 μm, n = 200), 15% responded to FPP, NAG, and capsaicin. Additionally 10% responded to FPP and NAG but not to capsaicin. These FPP- or NAG-mediated Ca²⁺ increases were
FIGURE 6. FPP- and NAG-induced Ca\(^{2+}\) mobilization in cultured small DRG neurons. A time course shows the effects of FPP and NAG on rat cultured DRG neurons using fura-2-based digital imaging techniques. The application of FPP (1 \(\mu\)M) or NAG (10 \(\mu\)M) for 10 s increased intracellular Ca\(^{2+}\) in HEPES-buffered solution (\(n = 5\)). A and B are pooled data obtained from five DRG neurons, whereas C and D are sampled recordings showing differential responses of FPP and NAG on a cultured DRG neuron. The black dot (●) represents acute application of each indicated drug. The response to capsaicin (0.1 \(\mu\)M) was examined in all DRG neurons tested. E and F are pooled data obtained from 12–27 siRNA-transfected DRG neurons. To knock down GPR92, cultured rat DRG neurons were exposed to siRNA against GPR92 (●). As a negative control siRNA (○), a nontargeting scrambled siRNA with no sequence homology to any known gene sequences was used. E and F represent agonist-induced [Ca\(^{2+}\)], signals in capsaicin-insensitive and -sensitive cells, respectively.

repeatedly observed by second application of the agonist. However, treatment of medium (30–40 \(\mu\)M) or large (>40 \(\mu\)M) diameter DRG neurons with FPP or NAG did not produce any changes in Ca\(^{2+}\) responses. To confirm that these agonist-induced Ca\(^{2+}\) increases occur via an activation of GPR92, we used siRNA-mediated silencing of GPR92 in cultured rat DRG neurons. The selective knockdown of GPR92 was confirmed by Western blot analysis (supplemental Fig. 4A). To monitor transfection efficiency in cultured rat DRG neurons, the GFP expression vector pEGFP-N1 was co-transfected with each siRNA (supplemental Fig. 4B). For each Ca\(^{2+}\) imaging experiment, the transfected cells without the GFP vector were loaded with fura-2/AM after separately confirming transfection efficiency (>80%). Agonist-induced [Ca\(^{2+}\)], increases were still observed in the scrambled siRNA-transfected cells. However, GPR92 siRNA-transfected cells showed much smaller [Ca\(^{2+}\)], increases compared with scrambled siRNA-transfected cells. For capsaicin-insensitive cells, GPR92 knockdown resulted in a 68.2 ± 0.8 and 64.3 ± 1.3% inhibition of NAG- and FPP-mediated [Ca\(^{2+}\)], increases, respectively (Fig. 6E). For capsaicin-sensitive cells, the NAG- and FPP-mediated [Ca\(^{2+}\)], increases were completely inhibited in GPR92 knockdown cells (Fig. 6F).

DISCUSSION

Human GPR92 is structurally most closely related to GPR23/LPA\(_4\), a recently identified receptor for LPA (7). Indeed GPR92 binds to LPA. This binding induces \(G_{i11}\), \(G_{12/13}\), and \(G_s\)-mediated signaling pathways (6, 8). Because the EC\(_{50}\) value for LPA in the production of IP and cAMP in cells expressing GPR92 ranges in micromolar levels, there is likely to be an agonist that is more potent than LPA. This possibility was further supported by the observation that a luminal extract is more potent than LPA in inducing cAMP production in GPR92-expressing cells (8).

While searching for potential GPR92 agonists, we observed that GPR92 is activated by various lipid-derived molecules with different affinities. Of these molecules, FPP demonstrated a higher potency and efficacy for activating GPR92 than LPA. FPP showed an approximate 10-fold lower EC\(_{50}\) value and 3-fold higher \(E_{\mathrm{max}}\) value than LPA in agonist-induced SRE-luc activity in cells expressing GPR92. NAG showed a potency similar to that of LPA for inducing SRE-luc activity.

One might suggest that FPP-induced SRE-luc activity is likely not due to G protein coupling because FPP can contribute to transcriptional activation independent of GPCR-mediated G protein activation. FPP is a donor for isoprenylation of many proteins, including the \(\beta\gamma\) subunit of heterotrimeric G proteins and the small GTPases such as Ras and Rho. Isoprenylation is necessary for translocation of these G proteins to the plasma membrane and subsequent activation (18–20). Because SRE can possibly serve as a downstream target of FPP-mediated Ras activation, it can be postulated that FPP-induced SRE-luc activity in this study is due to Ras activation. Further FPP is found to be a novel transcriptional activator for a subset of nuclear hormone receptors (21), suggesting possible G protein-independent

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effects of FPP on SRE-luc activity. However, this FPP-induced SRE-luc activity is only observed in GPR92-expressing cells but not in mock-infected cells. In GPR92-expressing cells, FPP induced immediate responses, such as Ca\(^{2+}\) mobilization, accumulation of IP and cAMP, and elevation of phospho-ERK. Additionally increased \(^{[35]}\text{S}GTP\gamma S\) binding indicates that FPP may have a direct interaction with GPR92, leading to activation of a G protein such as G\(_i\) and G\(_{i1}\).

Most GPCRs respond to a single endogenous ligand. Some phylogenetically related receptors share a common ligand. However, some receptors respond to more than two different compounds, each with different affinities (2, 3). GPR92 responds to various lipid-derived molecules including FPP, NAG, and LPA. These compounds possess a long hydrophobic carbon chain with a polar moiety. Our molecular modeling study, combined with site-directed mutagenesis, revealed that hydrophobic moieties of FPP and NAG have common binding sites, Gly\(^{98}\) and Phe\(^{101}\), at transmembrane domain 3. Gly\(^{98}\) is likely important for forming a binding pocket that allows a hydrophobic interaction between Phe\(^{101}\) of GPR92 and the hydrophobic carbon chains of FPP and NAG. In addition to common binding sites, GPR92 has specific binding residues for charged moieties of FPP and NAG. The negatively charged diphasphate moiety of FPP and the glycine moiety of NAG interact with Arg\(^{276}\) in the third extracellular loop and Thr\(^{97}\) at the transmembrane domain 3, respectively. In this regard, molecular interactions between GPR92 and other molecules such as LPA and 2-AG remain to be elucidated.

FPP and NAG are endogenously synthesized. FPP is a key intermediate in the biosynthesis of steroids, carotenoids, and polyisoprenoids. FPP is easily transported into plasma. The steady-state human plasma concentration of FPP is ~7 ng/ml (~15 nm) (22). As the mRNA for GPR92 is expressed in a variety of peripheral tissues, including spleen, thymus, stomach, small intestine, and kidney, it is possible that circulating FPP may exert its effects in these tissues through activation of GPR92.

NAG is naturally produced in a variety of tissues including the spinal cord, small intestine, kidney, glabrous skin, and brain (10). These tissues also contain GPR92 mRNA. Recently NAG was identified as an endogenous ligand for GPR18; it activates the G\(_i\)-mediated signaling pathway (23). This suggests that NAG plays a role in immune regulation because GPR18 is highly expressed in lymphoid tissues such as the peripheral blood leukocytes, spleen, and thymus (24, 25). The biological functions of NAG with regard to GPR92, however, are not yet understood. However, as tissue distributions of GPR92 and NAG are highly correlated (5, 10), it is postulated that NAG has a role in these tissues through activation of GPR92.

In good agreement with a previous observation (6), GPR92 was highly expressed in the DRG where FPP and NAG are able to induce Ca\(^{2+}\) elevation. This indicates possible roles of FPP and NAG in the DRG. FPP and its synthase activities are present in the spinal cord (26), suggesting a role of FPP in sensory neurotransmission through activation of GPR92. NAG is highly concentrated in the spinal cord and glabrous skin (10), which indicates a physical interaction of NAG with GPR92 in sensory neurons. However, it is not known how FPP and NAG are regulated in physiological conditions and whether FPP and NAG have direct actions on DRG neurons.

An interesting finding of this study is that GPR92 was expressed at high levels in small and medium diameter neurons of the DRG, which include primary sensory fibers associated with acute and neuropathic pain. Many GPR92-positive cells in the DRG coexpress TRPV1, which functions as a nociceptor (27). FPP- and NAG-induced Ca\(^{2+}\) elevation was observed only in small diameter DRG neurons. Some FPP- and NAG-responsive cells responded to capsaicin, a TRPV1 agonist. Of the capsaicin-responsive DRG neurons, ~38% of the cells responded to either FPP or NAG. This observation, together with coexpression of GPR92 and TRPV1, indicates a possible role of GPR92 in TRPV1-mediated pain sensing. Recently involvement of GPR92 in neuropathic pain has been suggested because GPR92 knock-out mice display significantly less sensitivity in the spinal cord to noxious mechanical and thermal stimulation than wild type littermates (28).

Several reports suggest involvement of NAG in nociception in the sensory nervous system. No evidence for the role of FPP in nociception has been provided. Peripheral administration of NAG in the hind paw is capable of suppressing the phase 2 response of formalin-induced pain behavior in the rat (10). Further recent studies suggest involvement of NAG in inflammatory and neuropathic pain. Intrathecal administration of NAG reduces the mechanical allodynia and thermal hyperalgesia induced by either intraplantar injection of Freund’s complete adjuvant or partial ligation of the sciatic nerve (29, 30). NAG is a structural analog of the endogenous cannabinoid anandamide. However, the pain suppressive effects of NAG are not likely mediated by activation of the cannabinoid CB1 receptor because NAG has a very poor affinity to the CB1 receptor (31, 32). One possible explanation for the effects of NAG is that NAG increases anandamide concentration by inhibiting the hydrolytic activity of fatty acid amide hydrolase on anandamide (33). Alternatively the coexpression of GPR92 and TRPV1 in the DRG raises the possibility that NAG can exert its pain suppressive effects through GPR92 in the sensory nervous system. As NAG acts as a partial agonist at GPR92, it is possible that it serves as an antagonist to FPP under physiological conditions thereby reducing activation of GPR92, although this possibility should be further investigated.

In summary, we have demonstrated that multiple lipid-derived molecules can activate GPR92 with different potencies. Of these, FPP was by far the most potent ligand and exerted effects with much greater efficacy than any of the other compounds tested, including LPA. However, we do not exclude the possible presence of other, as yet unknown, GPR92 agonists that have even greater efficacy. The presence of multiple ligands for GPR92 portends a wide range of biological and medicinal roles for this receptor. The roles of FPP and NAG through activation of GPR92 in peripheral tissues and the DRG require further investigation.

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