Molecular Cloning and Characterization of a Lysophosphatidic Acid Receptor, Edg-7, Expressed in Prostate

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Received November 22, 1999; accepted December 29, 1999

ABSTRACT

Two G protein-coupled receptors (Edg-2) and (Edg-4) for the lysolipid phosphoric acid mediator lysosphatidic acid have been described by molecular cloning. However, the calcium-mobilizing receptor Edg-4 is not expressed in some cell lines that exhibit robust calcium responses to this ligand, thus predicting the existence of additional receptor subtypes. We report here on the characterization of a third human lysosphatidic acid receptor subtype, Edg-7, which mediates lysosphatidic acid-evoked calcium mobilization. In a rat hepatoma Rh7777 cell line that lacks endogenous responses to lysosphatidic acid, this lipid mediator, but not others, evokes calcium transients when the cells have been transfected with Edg-7 or Edg-4 DNAs. Furthermore, frog oocytes exhibit a calcium-mediated chloride conductance in response to mammalian-selective lysosphatidic acid mimetics after injection of Edg-7 mRNA. Edg-7-expressing Rh7777 cells do not show inhibition of forskolin-driven rises in cAMP in response to lysosphatidic acid. However, membranes from HEK293T cells cotransfected with Edg-7 and Gαi1 protein DNAs show lysosphatidic acid dose-dependent increases in [γ-35S]GTP binding with an EC50 value of 195 nM. When we used this assay to compare various synthetic LPA analogs at Edg-2, Edg-4, and Edg-7 receptors, we found that ethanolamine-based compounds, which are full LPA mimetics at Edg-2 and Edg-4, exhibit little activity at the Edg-7 receptor. Edg-7 RNA was detected in extracts of several rat and human tissues including prostate. Together, our data indicate that Edg-7 is a third lysosphatidic acid receptor that couples predominantly to Gαi1,α proteins.

Lyosphosphatidic acid (LPA) is a potent extracellular lipid mediator that is released, for example, during platelet activation or by stimulation of adipocyte α2 adrenoceptors (Gerard and Robinson, 1989; Eichholtz et al., 1993; Valet et al., 1998). LPA elicits a wide variety of responses by cells; prominent among these are cell proliferation (van Corven et al., 1998; Howe and Marshall, 1993; Tokumura et al., 1994) and antiapoptosis (Levine et al., 1997; Weiner and Chun, 1999). LPA and the structurally similar lipid mediator sphingosine-1-phosphate (S1P) are recognized now to signal cells through a set of G protein-coupled receptors (GPCRs) known colloquially as the Edg receptors. Discovered initially as “orphan” receptors (Hla and Maciag, 1990), two members of the group, Edg-2 and Edg-4, have been shown to be LPA receptors. Edg-2 was shown first by Chun’s laboratory to mediate LPA activation of MAP kinase and inhibition of adenylyl cyclase in a pertussis toxin (PTX)-dependent manner and to induce cell shape changes in a Rho-dependent manner (Hecht et al., 1996). An and colleagues (1998a,b) showed that LPA binding to Edg-4 results in activation of PLCβ with subsequent calcium mobilization in a PTX-independent manner. However, expression of Edg-4 (as judged by RNA accumulation) is restricted primarily to leukocytes, suggesting the presence of another LPA receptor that couples to the Ca2+ mobilization elicited by LPA treatment in a wide variety of other cell types (An et al., 1998a). In this study, we report cloning and characterization of a novel human LPA receptor cDNA, Edg-7, from HEK 293 and PC-3 cells and show functional coupling of Edg-7 to calcium mobilization. While this manuscript was in initial review, another group published a paper characterizing this same LPA receptor (Bandoh et al., 1999). Although much of our data agrees with the findings of Bandoh and colleagues, our results contradict their observation that

This work was supported by National Institutes of Health Research Grants R01 GM52722 and R21 CA69848. C.E.H. is supported by a National Research Service Award predoctoral traineeship (T32 GM07065).

ABBREVIATIONS: LPA, 1-oleoyl lysosphosphatidic acid; S1P, sphingosine 1-phosphate; PTX, pertussis toxin; RT-PCR, reverse transcription-polymerase chain reaction; RACE, rapid amplification of cDNA ends; ORF, open reading frame; EST, expressed sequence tag; GPCR, G protein-coupled receptor; NOEPA, N-oleoyl ethanolamide phosphoric acid; NPSPA, N-palmitoyl serine phosphoric acid; NAEPA, N-acyl ethanolamide phosphoric acid.
Edg-7 does not respond to LPAs with saturated acyl functionalities.

Materials and Methods

Cloning of Human Edg cDNAs—Edg-7. A partial nucleotide sequence similar to Edg-2 and Edg-4 was found in a published patent application by Ellis et al., wherein the cDNA clone was named HOFNH30 (patent EP0878479A). Using that nucleotide sequence as a guide, we amplified via reverse transcription-polymerase chain reaction (RT-PCR) a 1022 base pair fragment from HEK293 cell RNA (oligonucleotide primers: forward 5′-caactaatgagttgcct, reverse 5′-tttataaggctggcct). Although stated in the patent application to be the full translational open reading frame (ORF), our nucleotide sequence analyses of multiple cDNAs from independent RT-PCR reactions differed consistently from the patent application sequence, most problematically in that all our cDNAs lacked an in-frame translational termination codon. Subsequent 3′ rapid amplification of cDNA ends (RACE), nested oligonucleotide primers 5′-ttgacagcttcaga and 5′-acatctctactctcagag, and subcloning allowed us to identify an in-frame termination codon and, ultimately, to use RT-PCR to isolate full-length DNAs from a mixture of HEK293 and PC-3 cell RNAs (primers: forward 5′-caactaatgagttgcct, reverse 5′-aaatcagagttgattggag). The resultant product (1191 base pair) of one of these cDNAs, which contained the full coding sequence, was subcloned into the plasmid expression vector pcRI. Edg-4, the full translational ORF of human Edg-4, as reported by An et al. (1998a) is contained within an expressed sequence tag (EST) cDNA (accession no. aa419064). We obtained this cDNA from the I.M.A.G.E. Consortium via Research Genetics (Birmingham, AL) and subcloned the ORF into the expression plasmid pcDNA3.1neo. However, this cDNA contains a translational frame shift compared with the sequence of two Edg-4 genomic clones (accession nos. ac002306 and ac011458) as well as another EST sequence resulting in a different C sequence of two Edg-4 genomic clones (accession no. aa419064). We obtained this cDNA from the I.M.A.G.E. Consortium via Research Genetics (Birmingham, AL) and subcloned this cDNA into the expression plasmid pcDNA3.1. Edg-4, the full translational ORF encoding mouse Edg-2, was subcloned into the expression plasmid pcDNA3 as reported previously by us (Hooks et al., 1998).

Oocyte Expression. Using the T7 RNA polymerase and the Edg-7 pcRI3.1 DNA as a template, we transcribed Edg-7 mRNA in vitro in the presence of a capping analog. This mRNA was injected into defolliculated stage V–VI Xenopus laevis oocytes. After ~60 h, responses to applied compounds were recorded from individual oocytes held under a two-electrode voltage clamp. The preparation of the oocytes and the conditions for our recordings were as described previously by us (Lynch et al., 1997).

Transient Expression in HEK293T Cells. The appropriate Edg plasmid DNA was mixed with an equal amount of an expression plasmid (pcDNA3) encoding a mutated (C351F) rat Gi2 protein, and these DNAs were used to transfect monolayers of HEK293T cells (in which T indicates expression of the simian virus 40 large T antigen) using the calcium phosphate precipitate method (Wigler et al., 1977). After ~60 h, cells were harvested and membranes were prepared, aliquoted, and stored at ~70°C until use.

Stable Expression in Rh7777 Cells. Rh7777 cell monolayers were transfected with the indicated Edg plasmid DNAs using the calcium phosphate precipitate method, and clonal populations expressing the neomycin phosphotransferase gene were selected by addition of geneticin (G418) to the culture medium. The RH7777 cells were grown in monolayers at 37°C in a 5% CO2/95% air atmosphere in growth medium consisting of 90% MEM, 10% fetal bovine serum, 2 mM glutamine, and 1 mM sodium pyruvate.

Measurement of Calcium Transients and cAMP Accumulation. Assays of calcium mobilization and adenyl cyclase activity were performed as described previously by us (Lynch et al., 1997). Briefly, intracellular calcium fluxes were measured on cell populations (2–4 × 10⁶ cells) that had been loaded with the calcium sensitive fluorophore INDO-1 in the presence of 2 mM probenecid. Responses were measured in a temperature-controlled fluorimeter (Aminco SLM 8000C, SLM Instruments, Urbana, IL). Lipids were delivered as aqueous solutions containing 0.1% (w/v) fatty acid-free BSA; this vehicle was determined to elicit no response. Assays of adenyl cyclase activity were conducted on populations of 5 × 10⁵ cells stimulated with 1 μM forskolin in the presence of the phosphodiesterase inhibitor isomethylbutykanthine. cAMP was measured by automated radioimmunoassay.

[γ-35S]GTP Binding. Briefly, 25 μg of membranes from Edg-7 (or Edg-2 or Edg-4) and GαsαβγCS151F DNA transiently transfected HEK293T cells were incubated in 1.0 ml of GTP-binding buffer (in mM: HEPES 50, NaCl 100, MgCl₂ 10, pH 7.5) containing 25 μg saponin, 10 μM GDP, 0.1 nM [γ-35S]GTP (1200 Ci/mmol), and indicated lipid for 30 min at 30°C. Samples were analyzed for membrane-bound radiolucine using a Brandel Cell Harvester (Gaithersburg, MD). For this assay, receptor was coexpressed with rat Gαsα in which amino acid 351 (normally cysteine) had been changed by mutagenesis to phenylalanine. The C351F mutation renders the protein resistant to inactivation by PTX or the allosteric agent N-ethylmaleimide.

RNA Analyses. RT-PCR analysis was performed using the Titanone One Tube kit. Oligonucleotide primers used to amplify human Edg-7 were forward 5′-ttgacagcttcagaca and reverse 5′-caacc. RNA extraction, Northern blotting, and hybridization of radiolabeled Edg-7 DNA were as described previously by us (O’Dowd et al., 1996).

Sources of Materials. Rh7777 cells (CRL 1601) were from the American Type Culture Collection (Manassas, VA). HEK293T cells were a gift from Dr. Judy White (University of Virginia), PC-3 cells were a gift from Dr. Charles Myres (University of Virginia), human Edg-1 cDNA was a gift from Dr. Tim Hla (University of Connecticut), human Edg-3 cDNA was a gift from Dr. Songzhu An (University of California at San Francisco), N-acylglucosamide phosphates and N-palmitoyl-1-serine phosphate were gifts from Dr. Timothy L. MacDonald (University of Virginia), 1-octyl LPA, 1-palmitoyl LPA and 1-myristoyl LPA, and other lysophospholipids were obtained from Avanti Polar Lipids (Alabaster, AL). S1P was obtained from Biomol (Plymouth Meeting, PA). [γ-35S]GTP from New England Nuclear (Boston, MA), 3′ RACE kit, geneticin, cell culture media, and sera from Life Technologies (Bethesda, MD), human northern blot membrane from Clontech (San Diego, CA), oligonucleotides from Operon Technologies (Alameda, CA), and the RT-PCR kit (Titanon One Tube) from Boehringer Mannheim (Indianapolis, IN). Expression plasmids were from Invitrogen (La Jolla, CA), and other chemicals were from Sigma (St. Louis, MO).

Results

Figure 1A shows the DNA sequence and deduced amino acid sequence of human Edg-7. This sequence is from a single cDNA, but it is the same sequence we found in four independent cDNAs obtained by RT-PCR and three partial cDNAs obtained by 3′ RACE of HEK293 cell RNA. Thus we are confident that the sequence we report represents the coding region of the Edg-7 mRNA population in HEK293 cells. However, our sequence differs substantially from the patent application sequence (EP0878479A) most prominently regarding a translational reading frame shift in the C-terminal region. We do not know why these sequences differ; perhaps they represent two alleles. Using the BLAST (Atschul et al., 1990) and FASTA (Pearson and Lipman, 1988) search tools, we found no record of a human Edg-7 DNA sequence in any division of the Genbank database. However, recently a mouse kidney cDNA sequence that is >90% identical (amino acids) to human Edg-7 appeared in the EST division. Our Edg-7...
sequence has been deposited with the GenBank (accession no. AF186380).

The conceptualized human Edg-7 protein (353 amino acids, 40,081 Da) is 52% identical with human Edg-2, 48% identical with human Edg-4, 34 to 37% identical with the human S1P receptors Edg-1, Edg-3, Edg-5, and Edg-8 (Im et al., 2000), and 36% identical with the human orphan receptor Edg-6. All other known rhodopsin family GPCRs share <28% identical amino acids. Hydropathy analysis of Edg-7 (data not shown) suggests that the heptahelix structure assumed to be common to GPCRs and the protein has the conserved amino acid motifs expected of a rhodopsin-like (family A) GPCR. The high similarity between Edg-7 and the known LPA receptors Edg-2 and Edg-4 (Fig. 1B) prompted us to test the former as a potential LPA receptor.

In testing for LPA receptor activity, we compared Edg-7 with Edg-2 and Edg-4. We introduced these DNAs individually into Rh7777 rat hepatoma cells by transfection and selected for geneticin-resistant clonal populations. Rh7777 cells were chosen because they were reported (Fukushima et al., 1998) to exhibit minimal responses to high concentrations (10 μM) of LPA but respond to LPA after transfection with Edg-2. In Edg-7 transfected cell populations, calcium transients were evoked by LPA (Fig. 2A) with an EC₅₀ value estimated to be ~100 nM. As predicted from previous reports (An et al., 1999a,b), we also detected calcium mobilization in Edg-4 DNA transfected, but not Edg-2 DNA transfected, cell populations (Fig. 2B). This calcium mobilization was not blocked by previous treatment with PTX (data not shown), suggesting the involvement of PTX-insensitive G proteins, most probably Gq/11α. It is noteworthy that other phospholipids, including lysophosphatidyl-choline, -serine, -inositol, -ethanolamine, -glycerol, and sphingosinephosphorylcholine did not evoke calcium transients in these cells at concentrations up to 10 μM (data not shown).

As a second test of calcium mobilization mediated by Edg-7, we injected the cognate mRNA into frog oocytes and measured changes in the calcium-gated chloride conductance in response to applied lipids. In this assay system, a strong endogenous response to LPA (Durieux et al., 1992) necessitates the use of mammalian receptor-selective LPA mimetics in lieu of LPA. Two such compounds, N-oleoyl ethanolamide phosphoric acid (NOEPA) and N-palmitoyl serine phosphoric acid (L-NPSPA), in which ethanolamine or L-serine replaces the glycerol of LPA (Suguira et al., 1994), meet this requirement (Hooks et al., 1998). As is shown in the recordings (Fig.

Fig. 1. A, the nucleotide and deduced amino acid sequence of the human Edg-7 DNA. The thymidylic acid residue at position 1184 was introduced to create a XbaI restriction endonuclease site during the cloning process. This sequence can be retrieved as a GenBank flat file using accession no. af186380. B, alignment of the Edg-2, Edg-4, and Edg-7 proteins created using the PILEUP algorithm of the GCG program. Potential transmembrane regions are overscored; gaps are indicated by dashes.
increased after 2 d, prepared membranes. In these membranes, LPA receptors Edg-1 and Edg-3 as well as the LPA receptors Edg-7, we tested clonal Rh7777 populations expressing the S1P receptors. By way of comparison, we found that Edg-2 but not Edg-4 inhibited cAMP accumulation in response to LPA, whereas S1P treatment (10 μM) resulted in a marked inhibition in both Edg-1- and Edg-3-expressing Rh7777 cells. As expected, pretreatment of cultures with PTX (100 ng/ml, 24 h) blocked the inhibition of forskolin-driven rises in cAMP accumulation in Edg-7-expressing cells, indicating a lack of coupling of to Gi/o proteins in this system (Fig. 4). In keeping with published reports (An et al., 1997; Fukushima et al., 1998; Gonda et al., 1999; Okamoto et al., 1999; Sato et al., 1999), we found that Edg-2 and Edg-4 couple through both Gq/11 and Gi/o proteins when expressed in Rh7777 cells, coexpression of receptor and G protein by cotransfection of their DNAs might result in observable receptor/G protein coupling. If so, we could use a [γ35S]GTP binding assay to detect receptor/G protein pairs. Lacking a fully validated radiolabeled LPA binding assay, such a demonstration is important to demonstrate unequivocally that LPA signals directly through Edg-7. Therefore, we introduced Edg-7 and Gi/oC351F DNAs by transfection into HEK293 T cells and, after 2 d, prepared membranes. In these membranes, LPA increased [γ35S]GTP binding in a dose-dependent manner, with an EC50 value of 195 nM (Fig. 5, A and B). Although HEK293 T cells exhibit an endogenous response to LPA, mock transfected cells showed only 20% of the response to LPA (Fig. 5A). Thus Edg-7 and Gi/oα interactions can occur, albeit when both recombinant proteins are expressed at artificially high levels. L-NPSPA also stimulated [γ35S]GTP binding in this assay (data not shown) as predicted from the oocyte responses shown in Fig. 2. However, a number of other lipids including S1P, dihydro S1P, sphingosylphosphorylcholine, and lysophospholipids with glycerol, choline, serine, inositol, and ethanolamine head groups did not stimulate [γ35S]GTP binding in these membranes (Fig. 5C).

The availability of the [γ35S]GTP binding assay provides an opportunity to measure the relative potencies and efficacies of synthetic LPA analogs at defined receptors without the confounding influence of endogenous LPA receptors. We exploited this opportunity by comparing the activity at Edg-2, Edg-4, and Edg-7 of a set of LPA analogs wherein the glycerol backbone is replaced by ethanolamine. Such com-

![Fig. 2. Calcium mobilization in Rh7777 cells transfected with Edg-2, Edg-4, and Edg-7 DNAs. A clonal population of Rh7777 cells transfected with Edg-7 DNA was loaded with the calcium fluorophore INDO-1 and aliquots challenged with the 1-oleoyl LPA at the indicated concentrations (A). Clonal populations transfected with the indicated Edg DNAs were challenged with 10 μM of 1-oleoyl LPA (B).](image1)

![Fig. 3. Responses of oocytes injected with Edg-77 mRNA to LPA mimetics. Individual frog oocytes were held under two-electrode voltage clamp and challenged with two different LPA mimetics at the concentration shown. L-NPSPA (Sugiura et al., 1994) is a competitive antagonist of the oocyte LPA response (Liliom et al., 1996), but a partial agonist of mammalian LPA responses (Hooks et al., 1998). NOEPA is a fully potent LPA mimetic at mammalian cells (Sugiura et al., 1994; Lynch et al., 1997), but a partial agonist of mammalian LPA responses (Hooks et al., 1998). NOEPA is a fully potent LPA mimetic at mammalian cells (Sugiura et al., 1994; Lynch et al., 1997), but is ~100-fold less potent than LPA on the oocyte (C.E.H., D.-S.-I., and K.R.L., unpublished data).](image2)

![Fig. 4. Inhibition of forskolin-evoked cAMP accumulation in Edg DNA transfected Rh7777 cells. Clonal populations of Rh7777 cells, either mock-transfected (normal) or transfected with the indicated Edg receptor, were treated with forskolin and challenged with S1P or LPA. The absolute values for cAMP accumulation are basal (i.e., IBMX alone) = 8.5 ± 1.4 pmol/well, IBMX + forskolin (100%) = 111.8 ± 13.8 pmol/well.](image3)
pounds are known to be potent LPA mimetics (Sugiura et al., 1994; Lynch et al., 1997), but their activity at individual receptors has not been reported. In response to a very recent report by Bandoh and colleagues (1999) suggesting that LPAs with saturated fatty acids are not active at the human Edg-7 receptor, we also compared 14:0, 16:0, and 18:0 LPAs at Edg-2, Edg-4, and Edg-7 using the GTP[γ-35S] binding assay. The results of these assays are presented in Fig. 6A–C. In agreement with Bandoh et al. (1999), who assayed Edg-4 and Edg-7 expressed in insect Sf9 cells, we found that 18:1 LPA was 1 to 2 log orders less potent at Edg-7 than at Edg-4. Furthermore, we discovered that LPA mimetics in which the glycerol backbone is replaced by N-acyl ethanolamine phosphoric acid (NAEPA) are quite active at Edg-2 (rank order potency, 18:1 > 18:2 > LPA ≈ 20:1 = 14:1) and Edg-4 (rank order, LPA > 18:1 > 18:2 ≈ 20:1 = 14:1) but exhibit strikingly less activity at Edg-7.

Bandoh and colleagues (1999) reported the unexpected finding that LPAs with saturated acyl groups were entirely inactive at Edg-7 (but active at Edg-4) at concentrations up to 10 μM. Although the low potency of LPA at Edg-7 in our assay system did not allow the determination of full dose-response curves, in contrast to those of Bandoh et al., our results (Fig. 6) indicate that 16:0 LPA and 18:0 LPA are agonists at Edg-7 as well as at Edg-4 and Edg-2. Although all compounds were less potent at Edg-7, the rank order potency of the LPA molecules (18:1 > 16:0 > 18:0 > 14:0) was the same for both Edg-4 and Edg-7. Edg-2, however, did not discriminate between 18:1 and 16:0 LPAs in this assay (Fig. 6C) Thus our data do not permit us to support the contention of Bandoh and colleagues that Edg-7 exhibits a peculiar preference for unsaturated LPAs.

Finally, we investigated the expression pattern of the human Edg-7 gene in human and rat tissues by Northern analysis. As noted previously, we cloned Edg-7 from embryonic kidney 293 and prostate carcinoma PC-3 cells. This expression was reflected in our detection of a signal in RNA extracts of kidney and prostate (Fig. 7, A–D). Other human tissues positive for Edg-7 RNA were heart and several areas of human forebrain in which signals from frontal cortex, hippocampus, and amygdala were particularly strong (Fig. 7C). In rat tissues, we detected a signal in extracts of kidney and

![Fig. 5.](image_url) [γ-35S]GTP binding to HEK293T cell membranes in response to 1-oleoyl LPA and S1P. A, the histogram shows the stimulation of [γ-35S]GTP binding in response to 1 μM LPA or S1P. B, the absolute values for [γ-35S]GTP binding are basal (no drug) 5880 ± 214 dpm and maximum 11789 ± 127 dpm. Each data point represents mean of six determinations ± S.E. C, each lipid was tested at a concentration of 1 μM. LPE indicates lysophosphatidyl ethanolamine; LPI, lysophosphatidyl inositol; LPS, lysophosphatidyl serine; LPC, lysophosphatidyl cholone; LPG, lysophosphatidyl glycerol; H2S1P, dihydrosphingosine 1-phosphate; SPC, sphingosylphosphoryl cholone.

![Fig. 6.](image_url) [γ-35S]GTP binding to HEK293T cell membranes in response to LPA s and NAEPAs. HEK293T cells were transfected with Edg-4 (A), Edg-7 (B), or Edg-2 (C) DNAs as well as with G protein DNAs (see Materials and Methods for details). Lipids were dissolved in chloroform and after determining their purity (by thin-layer chromatography) and concentration (colorimetric phosphate assay), dried and dissolved in buffer containing 0.1% fatty acid-free BSA. Typical values for minimum and maximum stimulation were 3,000 and 11,000 dpm, respectively.
The identification of LPA receptors is made problematic by the difficulty of radioligand binding and the widespread responsiveness of cultured mammalian cells to LPA. We responded to this dilemma in three ways. First, we used a $[\gamma^{35}\text{S}]$GTP binding assay that detects functional receptor/G protein complexes without the possibly confounding influence of downstream effector systems or endogenous GPCR signaling. Second, we used rat hepatoma Rh7777 cells, which are quite unusual in that they do not have detectable LPA responses (calcium mobilization or adenylyl cyclase inhibition). Finally, we used mammalian subtype-selective LPA mimetics that enable the use of the frog oocyte assay. Together, the results of these assays show that the Edg-7 DNA encodes a calcium mobilizing LPA receptor.

Interestingly, Edg-7 couples effectively to $G_{\alpha3}$C351F when both are introduced into HEK293T cells via DNA-mediated transfection, but does not couple to endogenous $G_{\alpha3}$ proteins in Rh7777 cells when the receptor alone is introduced. We chose this mutant $G_{\alpha3}$ protein so that if necessary we could suppress some background $[\gamma^{35}\text{S}]$GTP binding by treatment of cells with PTX or membranes with N-ethylmaleimide. (In practice, these maneuvers proved unnecessary.) Presumably, other $G$ proteins such as $G_{\alpha3}$ would function in this assay also. Our laboratory had shown previously that two compounds structurally similar to LPA, NOEPA, and $\alpha$-NPSPA (Suguiura et al., 1994) are LPA mimetics on human breast cancer MDA MB231 and HEK293 cells (Lynch et al., 1997; Hooks et al., 1998). Our present data using the oocyte indicate the previous observations were from interactions proceeding, at least in part, through Edg-7.

We took advantage of the $[\gamma^{35}\text{S}]$GTP binding assay to measure the relative potencies and efficacies of synthetic compounds at the Edg-2, Edg-4, and Edg-7 receptors. In a previous study (Lynch et al., 1997), we found that structural analogs in which ethanolamine replaced glycerol were full LPA mimetics when measured in calcium mobilization and adenylyl cyclase inhibition assays in MDA MB231 cells (these cells express Edg-2, -4, and -7 RNAs; Lynch, unpublished observations). In this assay, although the most active of these compounds (e.g., 18:1 or 18:2 NAEA) were quite indistinguishable from 18:1 LPA at Edg-4 (Fig. 6A) and more potent than 18:1 LPA at Edg-2 (Fig. 6C), all of the ethanolamine-based compounds exhibited only slight LPA mimic activity at Edg-7 (Fig. 6B). Thus 18:1 NAEA is a prototype for compounds that are Edg-2- and Edg-4- (versus Edg-7-) selective.

While this paper was under initial review, Bandoh et al. (1999) published their independent discovery of Edg-7. Both groups report the same nucleotide and amino acid sequence and agree on Edg-7’s calcium mobilizing properties and tissue localization. However, there are several differences between the two reports. Bandoh and colleagues report that Edg-7 and Edg-4, when expressed in insect Sf9 cells, stimulate cyclic AMP accumulation, whereas Edg-2 does not affect cAMP levels. This result is somewhat surprising, particularly regarding the apparent lack of $G_{\alpha3}$ coupling by Edg-2, which has been demonstrated repeatedly in mammalian systems (e.g., our Fig. 4). Perhaps the cyclic AMP accumulation by Edg-4 and Edg-7 in the Sf9 system is similar to the atypical protein kinase C activation of adenylyl cyclase type II in RAW264.7 macrophages reported recently (Lin et al., 1999). More interesting to us was the demonstration by Bandoh et al. that LPAs containing a saturated acyl group were entirely inactive at Edg-7 at concentrations up to 10 $\mu$M. However, our results using the GTP binding assay contradict those of Bandoh et al. in that we found 16:0 and 18:0 LPAs were active at Edg-7, albeit with lower potency than 18:1 LPA. This difference in results, which we are at a loss to explain, is particularly striking in that the assay system of Bandoh et al. (calcium mobilization in Sf9 cells) has an intrinsic amplification that renders their system more sensitive to LPA. Our results suggest that if one’s goal is to design LPA receptor-selective compounds, there is relatively little to gain in modulating the degree of unsaturation of the acyl chain. Our results with all three LPA receptors suggest the optimal acyl chain length is 16 to 18.

The existence of a third LPA receptor has been suspected because the other calcium-mobilizing LPA receptor, Edg-4, was reported to have limited tissue distribution (An et al.,...
Cloning of the LPA Receptor Edg-7 759


Phospholipase C-Ca 2+ system by sphingo-lysophosphorylcholine with signaling characteristics distinct from EDG1 and AGR16. Biochem Biophys Res Commun 260:203–208.

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