Omega-3 Fatty Acids and other FFA4 Agonists Inhibit Growth Factor Signaling in Human Prostate Cancer Cells

Ze Liu, Mandi M. Hopkins, Zhihong Zhang, Chrystal B. Quisenberry, Louise C. Fix, Brianna M. Galvan, and Kathryn E. Meier

Department of Pharmaceutical Sciences
College of Pharmacy
Washington State University
Spokane, Washington
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b) Corresponding author:

Kathryn E. Meier, Ph.D.
Department of Pharmaceutical Sciences
College of Pharmacy
Washington State University
Spokane, Washington 99210-1495

Telephone: 509-358-7631
Fax: 509-358-6673
E-mail: kmeier@wsu.edu

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Abstract

Omega-3 fatty acids (n-3 FAs) are proposed to have many beneficial effects on human health. However, the mechanisms underlying their potential cancer preventative effects are unclear. G protein-coupled receptors (GPCRs) of the free fatty acid receptor (FFAR) family, FFA1/GPR40 and FFA4/GPR120, specifically bind n-3 FAs as agonist ligands. In this study, we examined effects of n-3 FAs in human prostate cancer cell lines. Initial studies established that the long-chain n-3 FAs, eicosapentaenoic acid (EPA) and docosahexaenoic acid (DHA), inhibit proliferation of DU145 cells in response to lysophosphatidic acid (LPA), a mitogenic lipid mediator. When added alone to serum-starved DU145 cells, EPA transiently activates signaling events including p70S6K phosphorylation. However, when added 15 minutes prior to LPA, EPA suppresses LPA-induced activating phosphorylations of Erk, FAK, and p70S6K and expression of the matricellular protein CCN1. The rapid onset of the inhibitory action of EPA suggested involvement of a GPCR. Further studies showed that DU145 and PC-3 cells express mRNA and protein for both FFA4 and FFA1. TUG-891, a selective agonist for FFA4, exerts inhibitory effects on LPA- and EGF-induced proliferation and migration, similar to EPA, in DU145 and PC-3 cells. The effects of TUG-891 and EPA are readily reversible. The FFA1/FFA4 agonist GW9508 likewise inhibits proliferation at doses that block FFA4. Knockdown of FFA4 expression prevents EPA- and TUG-891-induced inhibition of growth and migration. Together, these results indicate that activation of FFA4 initiates signaling events that can inhibit growth factor-induced signaling, providing a novel mechanism for suppression of cancer cell proliferation.
Introduction

Consumption of omega-3 fatty acids, as dietary constituents or supplements, has been linked to various positive health outcomes, particularly with respect to cardiovascular health [Calder and Yaqoob, 2009]. With respect to cancer prevention or therapy, EPA and DHA have been shown to suppress the growth of carcinoma cells in culture, including prostate cancer cells [Chung et al., 2001; Wang et al., 2012]. Over-expression of FAT1, which converts n-6 to n-3 FAs, inhibits proliferation of DU145 and PC-3 prostate cancer cells [Lu et al., 2008]. Inhibitory effects of n-3 FAs on prostate tumor growth in animal models have also been reported [Brown et al., 2006; Akinsete et al., 2012]. Whether omega-3 fatty acid supplements are of benefit for prostate cancer prevention, based on human epidemiologic data, is quite controversial [Maclean et al., 2006; Brasky et al., 2013]. Nonetheless, the inhibitory effects of n-3 FAs on prostate cancer cells in culture suggest interference with growth factor signaling, which is worthy of further investigation.

The mechanisms underlying the potential cancer preventative effects of omega-3 fatty acids at the molecular level have not been firmly established [Gu et al., 2013b]. Although numerous mechanisms have been considered, the prevailing hypothesis is that metabolites of n-3 FAs are responsible for the anti-cancer effects. It is known that uptake of omega-3 fatty acids modify membrane phospholipid composition in a manner that decreases production of arachidonic acid (n-6)-derived eicosanoids that are pro-inflammatory [Norris and Dennis, 2012], and increases n-3 metabolites (e.g., “resolvins”) that are anti-inflammatory [Zhang and Spite, 2012]. For example, PGE₂ levels are decreased in prostate tumors following dietary supplementation of mice with
n-3 FAs [Kobayashi et al., 2006]. Thus, effects of n-3 fatty acids are usually tested after long-term exposure (e.g., overnight treatment in cell culture, or dietary interventions). More rapid effects have also been reported. For example, fatty acid derivatives can alter the organization of membrane microdomains [Ibarguren et al., 2014]. One study suggests that modification of the FA content of PIP3 is responsible for some of the inhibitory effects on prostate cancer cells [Gu et al., 2013a]. Other reported molecular targets for n-3 FAs include the nuclear receptor PPARγ [Edwards and O’Flaherty, 2008].

Recently, omega-3 fatty acids were shown to act as agonist ligands for FFA1/GPR40 and FFA4/GPR120, G protein-coupled receptors (GPCRs) in the free fatty acid receptor (FFAR) family [Oh et al., 2010]. This finding has been of particular interest with respect to the beneficial effects of omega-3 fatty acids on metabolism [Hara et al., 2011; Talukdar et al., 2011], but has not yet been fully explored with respect to cancer prevention or treatment. The roles of GPCRs in regulating growth and invasion of cancer cells are of increasing interest from a therapeutic standpoint [O’Hayre et al., 2014]. GPCR ligands initiate cellular responses within minutes, in contrast to the longer term effects of n-3 FAs studied previously with respect to lipid composition. However, responses to GPCR agonists can also be sustained over time. Both FFA1 and FFA4 are coupled to G<sub>q/11</sub> as well as arrestin-3, but many of their actions appear to be arrestin-mediated [Mancini and Poitout, 2013; Butcher et al., 2014]. Selective agonists for FFA1 and FFA4 are rapidly being developed [Hudson et al., 2013b; Du et al., 2014], making it possible to test whether such agents elicit therapeutic effects of interest.

In the current study, we tested whether omega-3 fatty acids exert short-term inhibitory effects on growth factor signaling in human prostate cancer cell lines, and
found that this was the case. We then used FFAR agonists to investigate whether GPCRs are responsible for the inhibitory effects observed. Lysophosphatidic acid (LPA) was used as the primary growth factor, based on the experience of our research group in studying LPA response [Xie et al., 2002; Gibbs et al., 2009] and on the importance of this lipid mediator in prostate cancer cells [Wang et al., 2004; Bektas et al., 2005; Guo et al., 2006; Choi et al., 2009]. Our study also characterizes effects of LPA and n-3 FAs on induction of CCN1, a matricellular protein that facilitates proliferation and migration [Jun and Lau, 2011]. Together, the results raise new possibilities for mechanisms by which n-3 FAs can inhibit cancer cell growth and survival.
Materials and Methods

Materials. EPA and DHA (prepared in ethanol) were from Cayman Chemical (Ann Arbor, MI). The FFAR agonists TUG-891 (4-[(4-fluoro-4'-methyl[1,1'-biphenyl]-2-yl)methoxy]-benzenepropanoic acid; prepared in DMSO) and GW9508 (4-[3-phenoxyphenyl)methyl] amino]-benzenepropranoic acid; prepared in ethanol) were from Millipore (Billerica, MA) and Cayman Chemical, respectively. Vehicle controls for FFAR agonists had a final concentration of 0.03% (v/v) ethanol. LPA (18:1; oleoyl) was obtained from Avanti Polar Lipids (Birmingham, AL), and was delivered to cells as a 1000X stock solution that had been prepared in 4 mg/ml fatty acid-free bovine serum albumin (BSA). Vehicle controls for LPA had a final concentration of 4 µg/ml BSA. EGF was from Sigma (St. Louis, MO). Anti-CCN1 (lot # F0509; 1:1000 dilution) was from Santa Cruz Biotechnology (Santa Cruz, CA). Antibodies recognizing the phosphorylated forms of p70S6 kinase (Thr389) (reference 01/2014, lot 15; 1:1000 dilution), ERK (T202/Y204) (reference 05/2007, lot 21; 1:1000 dilution), and Akt (Ser 473) (reference 12/2006, lot 10; 1:1000 dilution) were obtained from Cell Signaling Technologies (Beverly, MA). Antibody recognizing human focal adhesion kinase (FAK) phosphorylated at pY397 (lot #33526; 1:1000 dilution) was obtained from BD Transduction Laboratories (Lexington, KY), as was anti-actin (lot # 51711), which was used at a 1:5000 dilution. Anti-FFA4 (lot #GR26308-20) was purchased from Abcam (Cambridge, MA) and used at a 1:1000 dilution. Goat anti-rabbit secondary antibody (lot #083M4752) was purchased from Sigma (St Louis, MO), and used at 1:20,000 dilution, while goat anti-mouse secondary antibody (lot #1124907A) was purchased from Invitrogen/Life Technologies (Grand Island, NY), and used at a 1:5000 dilution.
Cell culture. DU145, PC-3, LNCaP, and Caco-2 cells were obtained from the American Type Culture Collection (Manassas, VA). The prostate cancer cells were grown in RPMI 1640 medium with 10% FBS (Hyclone/Thermo-Fisher Scientific, Waltham, MA). Caco-2 cells were maintained in DMEM medium supplemented with 10% fetal bovine serum (FBS) (Invitrogen/Life Technologies, Grand Island, NY). All cell lines were grown in an incubator at 37°C in 5% CO2 on tissue culture plastic.

Cell proliferation assays. DU145 cells were seeded in 6-well plates at 3x10^5 cells/well in serum-containing medium. After 1 day, the medium was generally changed to RPMI 1640 without serum. For some experiments testing serum response, cells were maintained continuously in serum. On the next day, the medium was changed to RPMI 1640 with 10% FBS, 10 µM LPA, or 10 nM EGF, in the absence or presence of 20 µM DHA, 20 µM EPA, or various concentrations of TUG-891 or GW9508. Control cells were incubated with the appropriate vehicle (0.03% ethanol, v/v). Duplicate wells were prepared for each experimental condition. Cell numbers were evaluated after 24, 48, and 72 hours by removing medium, incubating cells with trypsin/EDTA for 5 minutes, adding trypan blue, and counting the suspended live cells (excluding trypan blue) using a hemacytometer.

Cell migration assays: Cell migration was assessed using a modified Boyden chamber method, as previously described [Zhang et al., 2011]. Cells were serum starved for 24 hours and then seeded in serum-free medium at 2.5 X 10^4 cells per insert in the upper chambers of 8-µm transwell inserts (BD Biosciences, San Jose, CA). Cells were then treated with 10% FBS, 20 µM EPA, 1 µM TUG-891, 10 µM LPA, or 10 nM EGF, either alone or in combination. Serum-free medium was added to the lower wells. Following a
6 hour migration period, the insert membranes were fixed and stained using methanol and crystal violet. Cells that invaded the lower chambers were counted by microscopy.

**RT-PCR:** RNA from cells was isolated using an RNeasy Mini kit (Qiagen, Valencia, CA). First-strand complementary DNA (cDNA) was synthesized with SuperScript II Reverse Transcriptase (Invitrogen) following the manufacturer’s instructions using 20 μl of reaction mixture containing 2 μg RNA. PCR was carried out using Platinum Pfx DNA Polymerase (Invitrogen) and Integrated DNA Technology (San Diego, CA) probes: FFA4 (F: 5’- CCTGGAGAGATCTCGTGGGA-3’; and R: 5’- AGGAGGTGTCCGAGGTCTG-3’); FFA1 (F: 5’-CTCCTTCGGCCTCTATGTGG-3’; and R: 5’-AGACCAGGCTAGGGGTGAGA-3’); RPLP0 (F: 5’-CGCTATCCGCGGTTTCTGAT-3’; and R: 5’-AGACGATGTCACTTCCACGA-3’).

Following the manufacturer’s instructions, 5 μg cDNA template was used for each reaction. In brief, three-step cycling was used for 35 cycles, with denaturing at 94°C for 15 seconds, annealing at 55°C for 30 seconds, and extending at 68°C for 60 seconds. Products were separated by ethidium bromide agarose gel electrophoresis, and were then imaged using a ChemiDoc with Image Lab software (Bio-Rad, Hercules, CA).

**Cell incubations for signal transduction assays:** Cells were grown in F12K or RPMI 1640 medium supplemented with 10% FBS until ~80% confluent. Cells were serum-starved for 24 hours in F12K or RPMI 1640 medium, then incubated with 10 μM LPA, 10 nM EGF, 20 μM DHA, and/or various concentrations of EPA, TUG-891 or GW9508 for various times. Cells were rinsed twice with ice-cold phosphate-buffered saline (PBS), harvested by scraping into 1 ml ice-cold PBS, collected by centrifugation at 10,000xg for 10 min at 4°C, and resuspended in ice-cold lysis buffer (20 mM HEPES [pH=7.4]), 1%
Triton X-100, 50 mM NaCl, 2 mM EGTA, 5 mM β−glycerophosphate, 30 mM sodium pyrophosphate, 100 mM sodium orthovanadate, 1 mM phenylmethylsulfonyl fluoride, 10 µg/ml aprotinin, 10 µg/ml leupeptin). Insoluble debris was removed after centrifugation. Protein concentrations of these “whole-cell extracts” were determined using a Coomassie binding assay (Pierce Chemical, Rockford, IL).

**Immunoblotting:** Whole-cell extracts containing equal amounts of protein (30 µg) were separated by SDS-PAGE on 10% Laemmli gels, transferred to nitrocellulose, and incubated with primary (overnight at 4 °C) and then secondary (one to two hours at room temperature) antibodies. Blots were developed using enhanced chemiluminescence (GE Healthcare), and imaged using a Gel Doc system (BioRad). Protein expression was quantified by densitometry using Quantity One software (Bio-Rad). Results were normalized to the actin loading control, and then to the value obtained for untreated control cells.

**RNA interference:** Pre-designed human Silencer Select siRNA specific to FFA4 (s50346) was purchased from Life Technologies (Grand Island, NY), along with Opti-MEM reduced serum medium and Lipofectamine 3000 reagent. The siRNA was re-suspended in RNase-free water at a final concentration of 60 µM and stored at -20°C. AllStars Negative Control siRNA oligos were purchased from Qiagen and were used for all negative controls. According to the manufacturer’s directions for RNAi transfections, cells were plated in 1.7 ml medium per well in RPMI medium plus 10% FBS without antibiotics in 6-well plates one day before transfection. At the time of transfection, cells were approximately 50% confluent. For each transfection, two mixtures were prepared: 1) duplex siRNA added to 150 µl Opti-MEM, and 2) 4 µl Lipofectamine 3000 added to
150 µl Opti-MEM. The two solutions were gently mixed and then incubated at room temperature for 5 minutes. Final concentrations of siRNA were 180 nM. The mixtures were added to each well and cells were cultured for an additional 24-72 h. For LPA- and EGF-treated cells, the same procedure was followed, except cells were seeded in 6-well plates two days before transfection and serum starved overnight the day before transfection.

*Statistical analysis.* Data were analyzed by two-way ANOVA followed by either Tukey’s or Sidak’s multiple comparisons test. The only exception were assays in which there was only one time point (e.g., migration assays), which were analyzed by one-way ANOVA followed by Tukey’s multiple comparisons test. All analyses were done using Prism software (Graphpad, San Diego, CA). The data presented are representative of at least two experiments, unless otherwise indicated in the figure legends.
Results

Effects of EPA and DHA on DU145 cell proliferation

DU145 cells were serum-starved for 24 hours to remove serum growth factors (including LPA), then treated with 10% serum, 10 µM LPA, and/or 20 µM DHA, or 20 µM EPA for 24, 48, and 72 hours. The doses initially used for EPA and DHA were selected based on a literature survey; detailed dose-response data will be presented below. Notably, the doses used here are considerably lower than those used in previous studies with prostate cancer cells [Chung et al., 2001; Wang et al., 2012]. Ethanol was used as a vehicle control for EPA/DHA, and BSA as a vehicle control for LPA; neither vehicle affected responses at the concentrations used (data not shown).

As shown in Figure 1, LPA significantly stimulates proliferation of DU145 cells. Compared to serum-free control cells, the number of viable cells was decreased by treatment with EPA, but not by DHA, at 48 and 72 hours. However, cell number was not reduced below the number initially seeded in the dish, and the number of non-viable cells was not increased, indicating that EPA inhibits proliferation and does not cause cell death. Cells achieved confluence after 72 hours in the presence of serum; cells maintained in medium without serum remained viable for 72 hours (data not shown). Both EPA and DHA block LPA-induced proliferation in DU145 cells. However, since EPA was consistently more efficacious than DHA in this and other assays (data not shown), subsequent studies focused on the effects of EPA.

Effects of EPA on LPA-induced signal transduction

We reasoned that inhibitory effects of EPA on proliferation would likely be
reflected in effects on growth factor-induced signal transduction. Since we had previously characterized signaling responses to LPA in DU145 cells [Gibbs et al., 2009], we tested whether EPA influences representative LPA responses. A 15-minute pre-treatment with EPA was used to screen for effects that might be receptor mediated.

The first response tested was activation of the Erk MAPK pathway, which plays an important role in LPA-induced proliferation in prostate cancer cells [Kue et al., 2002; Gibbs et al., 2009]. A time course was conducted in which cells were pre-incubated with 20 µM EPA for 15 minutes, then incubated with LPA for various times. As shown in Figure 2A and B, EPA inhibits LPA-induced Erk phosphorylation in DU145 cells.

We next examined tyrosine phosphorylation of FAK, another response elicited by LPA in prostate cancer cells [Gibbs et al., 2009]. EPA inhibits LPA-induced FAK phosphorylation over the 6-hour time course of the experiment (Figure 2C and D).

The effects of EPA on activation of p70S6K, which is activated via the PI3K/Akt/mTORC1 pathway, were investigated. Signaling through this pathway is frequently up-regulated in prostate cancer [Morgan et al., 2009]. As shown in Figure 3A, LPA stimulates the activating phosphorylation of p70S6K within 10 minutes, after which the response gradually declines over 6 hours. LPA also stimulates activating phosphorylation of p-Akt in these cells (data not shown). EPA suppresses LPA response throughout the time course. Figure 3B presents quantification, including the effects of EPA alone. EPA alone causes a transient partial activation of p70S6K (10 minutes). However, in marked contrast to LPA, EPA is not able to induce sustained activation of p70S6K. The ability of EPA alone to initiate transient signaling has the result that significant inhibitory effects of EPA on LPA signaling are not always observed
at early time points, when the stimulatory responses to EPA alone can obscure inhibitory effects on LPA signaling. In Figure 3B, the difference between LPA alone and LPA plus EPA was only significant at 60 minutes, as analyzed by ANOVA.

We noted that the inhibitory effects of EPA on activation of the PI3K/Akt/p70S6K pathway in prostate cancer cells, as observed in our study, contrast with the role reported for FFAR agonists in some other cancer cell models [Wu et al., 2013]. In a study by Wu and co-workers, the FFA1/FFA4 agonist GW9508 was shown to rapidly stimulate Akt activation when applied alone to serum-starved colon cancer cells. We therefore tested effects of LPA and EPA in Caco-2, a human colon cancer cell line that expresses FFA4 [Mobraten et al., 2013; Wu et al., 2013]. LPA has been shown to induce proliferation of Caco-2 cells [Yun et al., 2005]. Our data show that LPA increases activating phosphorylation of p70S6K in Caco-2 cells (Figure 3C); similar results were seen for phospho-Akt (data not shown). However, the response to EPA alone was similar to or greater than the response to LPA (e.g., ~4-fold increase in p-p70S6K at 30 minutes with EPA alone). Both magnitude and duration of the EPA response were in contrast to the weak effect of EPA alone in DU145 cells (Figure 3B). The significant p70S6K response to EPA in Caco-2 interfered with efforts to determine whether EPA exerts an inhibitory influence in these cells; i.e., the response to a combination of EPA and LPA was approximately additive (data not shown). The stimulatory effect of EPA alone in Caco-2 cells is generally consistent with the GW9508-induced activation of Akt reported previously for other colon cancer cell lines [Wu et al., 2013].

The final signaling response assessed in prostate cancer cells was induction of CCN1. This response requires transcription and translation and is therefore a later
response to growth factor action. Expression of CCN1 was first examined in the three most commonly used human prostate cancer cell lines, PC-3, LNCaP, and DU145. As shown in Figure 4A, all three cell lines express CCN1 when grown in the presence of serum; levels are highest in DU145 cells, and lowest in the androgen-independent cell line LNCaP. Although CCN1 is eventually secreted, cellular levels were assessed in our experiments. LPA induces robust expression of CCN1 in PC-3 cells (Figure 4B), a response that is blocked by the LPA receptor antagonist Ki16425. The time course of the effects of LPA on CCN1 levels is presented in Figure 4C, where maximal induction was observed at 4 hours. These studies established that CCN1 is expressed in human prostate cancer cells, and that its expression can be induced in by LPA. Figures 4D and 4E present data for DU145 cells, analogous to Figures 2 and 3. LPA induces CCN1 expression in DU145 cells by 1 hour; this response is blocked by 20 µM EPA. EPA alone has no effect on CCN1 levels (Figure 4F); this result was confirmed by a time course study where were treated with EPA for up to 24 hours (data not shown).

The effects of EPA on responses to another growth factor, EGF, were also tested. Previous work from our lab has shown that EGF and LPA have similar effects on signal transduction in human prostate cancer cells [Gibbs et al., 2009]. Figure 5A illustrates the relative effects of serum, LPA, and EGF on proliferation of DU145 cells. Consistent with Figure 1C, LPA is a weaker mitogen than fetal bovine serum, which contains a mixture of growth factors. EGF acts similarly to LPA. Figure 5B demonstrates that EPA inhibits activation of Erk and FAK in EGF-treated cells, while Figure 5C shows that EPA also blocks induction of CCN-1 in response to EGF. Our group has previously shown that EGF can increase production of LPA, resulting in transactivation of LPA.
receptors, in ovarian cancer cells [Snider et al., 2010]. We therefore tested for receptor cross-talk in DU145 cells. As shown in Figure 5D, the LPA receptor antagonist Ki16425 suppresses proliferation induced by either LPA or EGF, suggesting that LPA receptors play a role in EGF response in DU145. Similar results were obtained in PC-3, another human prostate cancer cell line (data not shown). Thus, the inhibitory effects of EPA on EGF signaling could conceivably be an indirect consequence of EPA-induced suppression of LPA signaling.

Effects of FFAR pharmacologic agonists on growth factor response

The results presented above show that EPA can act rapidly (within 15 minutes) to inhibit mitogenic responses. This rapid action suggests a receptor-mediated response. The roles for FFARs were therefore investigated.

Two FFARs have been shown to bind long-chain n-3 FAs as agonists: FFA1/GPR40 and FFA4/GPR120. FFA4 is widely expressed, including in both normal and cancerous prostate tissue according to a database search [www.oncomine.org], and thus seemed a likely candidate in prostate cancer cells. RT-PCR was performed to determine whether this receptor is expressed in human prostate cancer cell lines. The results of RT-PCR experiments are shown in Figure 6A. Caco-2 cells, which have previously been shown to express FFA4 [Mobraten et al., 2013], were used as a positive control. The data show that DU145 cells express mRNA for FFA4. PC-3, another androgen-independent human prostate cancer cell line, expresses lower levels of FFA4 mRNA. Interestingly, both cell lines also express FFA1 mRNA, with PC-3 expressing much higher levels than DU145. The expression of FFA4 protein was confirmed in both cell lines by immunoblotting, as shown in Figure 6B.
Based on these results, we tested whether TUG-891, a recently described pharmacologic agonist that is selective for human FFA4 [Simpukade et al., 2012; Hudson et al., 2013], exerts effects similar to those of EPA. TUG-891 has 1000-fold higher affinity for FFA4 than for FFA1. We also tested for effects of GW9508, an agonist that conversely is 1000-fold selective for FFA1 over FFA4 [Briscoe et al., 2006].

We first performed a concentration-effect ("dose-response") study examining the effects of FFAR agonists on cell proliferation. Neither agonist affects DU145 cell proliferation when added alone to serum-starved cells (data not shown). Figure 7A shows that EPA inhibits LPA-induced proliferation of DU145 cells with an IC\textsubscript{50} of 5.7 µM. The FFA4-selective agonist TUG-891 also inhibits the ability of DU145 cells to proliferate in response to LPA. The IC\textsubscript{50} for this effect is 73 nM, which is consistent with published EC\textsubscript{50} values of ~100 nM for TUG-891 for effects mediated by human FFA4 [Hudson et al., 2013]. The relative potencies of EPA, DHA, GW9508, and TUG-891 are also consistent with previously published values. TUG-891 was developed as a selective FFAR inhibitor, and has higher potency than the endogenous n-3 FAs [Hudson et al., 2013]. GW9508, the FFA1-selective agonist, inhibited proliferation with an IC\textsubscript{50} of 950 nM. This value is much higher than the published EC\textsubscript{50} of GW9508 at FFA1 (~50 nM), but is consistent with its EC\textsubscript{50} for FFA4 (~2 µM) [Briscoe et al., 2006]. In Figure 7B, concentration-effect curves were obtained for inhibition of serum-induced proliferation by TUG-891, GW9508, EPA, and DHA; in these experiments DU145 cells were maintained continuously in serum. It was noted that both agonists had approximately 10-fold higher IC\textsubscript{50} values against serum as compared to LPA. In contrast, GW9508 had a lower IC\textsubscript{50} value with serum than with LPA. There are at least two possible
explanations for the differences in IC$_{50}$ values between LPA and serum. First, serum contains proteins, lipids, and fatty acids that potentially interfere with the delivery and/or action of the FFAR agonists; this is especially likely to be the case for EPA and DHA, which are fatty acids that bind to plasma proteins. Second, serum contains multiple growth factors, some of which may act through mechanisms that are not inhibited by FFA4 agonists, or that are more sensitive to FFA1 agonists. In any case, the FFA4 agonist TUG-891 was the most potent inhibitor of proliferation induced by either LPA or serum. In summary, while the concentration-effect data do not completely exclude a role for FFA1, they are most consistent with a role for FFA4.

Next, we asked whether the effects of TUG-891 on proliferation were reversible. TUG-891, a pharmacologic agent that is not a lipid, is not incorporated into phospholipid and acts reversibly [Hudson et al., 2013b]. In Figures 7C and 7D, DU145 cells were incubated with TUG-891 and serum to inhibit proliferation, after which the TUG-891 was removed and serum incubation was continued. Serum-induced proliferation resumes after TUG-891 is removed. These results demonstrate reversibility, confirm that TUG-891 is not toxic to DU145 cells, and indicate that the FFA4 agonist must be present continuously to inhibit proliferation. We next tested whether the effects of EPA are reversible. Since exogenous EPA has been shown to be readily incorporated into phospholipid in prostate cancer cells in culture [Gu 35 al., 2013a], effects of n-3 FAs on membrane fluidity, phospholipid composition, or generation of n-3 metabolites from FAs incorporated into phospholipid would be anticipated to be only slowly reversible. However, the reversibility of the inhibitory effects of EPA was observed to be similar to those of TUG-891 (Figures 7D and 7D). The rapid reversibility is more consistent with a
receptor-mediated effect of EPA.

The effects of the FFA4 agonist TUG-891 on LPA-induced signal transduction were examined. Figure 7E shows that EPA and TUG-891 have significant inhibitory effects on p70S6K activation when LPA response is measured at 10 minutes. Thus, a selective FFA4 agonist can suppress responses to LPA.

In Figure 8, we tested whether the effects of FFA4 agonists extend to another prostate cancer cell line. The effects of EPA and TUG-891 were tested on proliferation induced by serum, LPA, or EGF in both DU145 and PC-3 cells. The top panels show that both EPA and TUG-891 significantly inhibit serum-induced proliferation in both cell lines. In these experiments, the cells were serum-starved prior to addition of the FFAR agonist and serum. The inhibitory effect of the agonists at the doses used is partial, which is consistent with the data shown in Figure 7B. The remaining panels show that EPA and TUG-891 are both able to completely block proliferation in response to LPA (middle panels) and EGF (bottom panels) in both cell lines. Thus, TUG-891 and EPA inhibit proliferation in two different human prostate cancer cell lines.

FFA4 protein levels, as detected by immunoblotting, were not significantly altered by treatment of DU145 or PC-3 cells with either 20 µM EPA or 1µM TUG-891 for up to 4 hours (data not shown). These results suggest that FFA4 is not subject to agonist-mediated proteolytic down-regulation under the conditions tested.

**Effects of FFAR agonists on cell migration**

In order to evaluate effects of FFAR agonists on another response relevant to cancer cell biology, we examined cell migration. A previous report showed that n-3 FAs inhibit migration of PC-3 cells [Brown et al., 2006]. As shown in Figure 9, LPA and EGF
both enhance migration of DU145 and PC-3 cells, as assessed in a modified Boyden chamber assay. The response to either agonist was approximately 70% of the maximal response elicited by 10% serum in both cell lines (data not shown). EPA and TUG-891 significantly inhibit migration induced by either LPA or EGF, in both DU145 and PC-3 cells.

Effects of FFA4 knockdown

As another approach to test whether FFA4 mediates the inhibitory effects of FFAR agonists, siRNA was used to inhibit FFA4 expression in DU145 cells. siRNAs are normally administered to cells in the continuous presence of growth factors (i.e., not serum starved). Fortuitously, as shown previously (see Figure 7B), FFAR agonists can inhibit proliferation of cycling DU145 cells. First, we optimized conditions for the FFA4 knockdown. As shown in Figure 10A, the siRNA treatment was sufficient to achieve a maximal decrease in FFA4 protein levels at 24 hours. The extent of knockdown was 67.2 ± 7.0% (n=5) at 24 hours, as measured in the absence of FFA4 agonist. Next, we tested whether FFAR agonists can inhibit serum- or LPA-induced proliferation in cells deficient in FFA4. In these experiments, serum or LPA was added at the same time as the FFA4 siRNA. FFA4 knockdown alone, in the absence of FFA4 agonists, had no effect on proliferation in response to either serum or LPA (data not shown). The results presented in Figures 10B and 10C demonstrate that EPA and TUG-891 significantly inhibit serum-induced proliferation at 48 and 72 hours in DU145 and PC-3 cells treated with scrambled siRNA (negative control), but have no effect in cells treated with FFA4 siRNA. Similarly, in Figures 10D-G, we tested effects of EPA and TUG-891 on LPA- and EGF-induced proliferation with FFA4 knockdown in both DU145 and PC-3 cells. Cells
were serum-starved for the LPA and EGF experiments, although the medium contained low levels of Opti-MEM, which contains proprietary growth factors. Time course studies for both LPA and EGF yielded similar results at 24, 48, and 72 hours; for clarity, only data for the 48-hour time point are presented. The results presented in Figure 10 show that FFA4 knockdown eliminates the inhibitory effects of EPA and TUG-891 on proliferation.

Similar studies were performed to test for effects of FFA4 knockdown on migration of DU145 and PC-3 cells. As shown in Figure 11, inhibitory effects of TUG-891 and EPA on migration induced by LPA, EGF, and serum were absent when FFA4 was knocked down.

It is interesting to note that FFA4 knockdown suppressed effects of FFA4 agonists on proliferation in Figure 10, even though FFA4 was expressed by the cells at the beginning of the incubation. In other words, the FFA4-mediated signaling that occurs initially, prior to FFA4 knockdown, is not sufficient to block proliferation occurring over a longer time course. These results, along with the reversibility results shown in Figure 7, suggest that the inhibitory effects of FFA4 agonists require ongoing stimulation of FFA4, in the context of the continuous presence of growth factor receptor agonists.
Discussion

This study presents unexpected findings showing that omega-3 fatty acids and FFA4 agonists can inhibit growth factor-induced signaling. To our knowledge, ours is the first study to examine the roles of FFA4 in prostate cancer cells.

Most of the experiments described herein utilized LPA as the mitogenic agonist. The pro-mitogenic and pro-migratory activities of LPA are mediated by GPCRs in prostate cancer cells [Hao et al., 2007; Gibbs et al., 2009]. Several mechanisms could potentially be responsible for the inhibitory effect of FFA4 agonists on LPA response in prostate cancer cells.

First, activation of FFA4 could potentially activate signal transduction pathways that act downstream to "functionally antagonize" LPA action. For example, agents that increase cAMP levels have been shown to inhibit Erk activation in several model systems [Stork and Schmitt, 2002], and cAMP inhibits LPA-induced RhoA activation in PC-3 cells [Chen et al., 2005]. However, the broad spectrum of the inhibitory action of EPA and FFA4 agonists argues against this type of mechanism. The signal transduction responses examined in our study, Erk, p70S6K, and FAK, are activated through signaling pathways that are largely separate, although cross-talk between pathways does occur. Erk is activated through the Ras/Raf/MEK pathway. The Akt/mTORC1 pathway initiates activation of p70S6K. FAK is subject to regulation by several pathways, including integrins, but the activating phosphorylation that was assessed in our study is performed by c-Src. Therefore, the demonstrated ability of FFA4 agonists to suppress all of these pathways simultaneously suggests a more general mechanism that lies upstream of the protein phosphorylation cascades.
A second type of inhibitory mechanism involves cross-talk between GPCRs. For example, activation of FFA4 could heterologously desensitize LPA receptors, so that subsequent responses to LPA are inhibited. Consistent with this idea, it has been noted that FFA4 agonists cause prominent homologous desensitization of FFA4 via β-arrestin binding [Watson et al., 2012; Dranse et al., 2013; Burns et al., 2014]. The fact that EPA and TUG-891 also inhibit responses to EGF, in our studies, could argue against this mechanism. However, our data demonstrate that LPA antagonists can inhibit response to EGF, suggesting a significant role for EGF-induced transactivation of LPA receptors in EGF response. Thus, desensitization of LPA receptors could also inhibit responses to EGF. Another type of negative cross-talk can occur that involves heterodimerization of GPCRs [Milligan, 2004]. This phenomenon has not been demonstrated to date for FFA4, but is conceivable. LPA receptors have been shown to heterodimerize with other GPCRs in prostate cancer cells [Ward et al., 2011]. Cross-talk that involves changes in receptor localization is also conceivable, since FFA4 is efficiently internalized in response to agonists [Watson et al., 2012]. Further work will be needed to explore these potential mechanisms of inhibition.

The experimental protocol used in our studies facilitated the acquisition of unique insights into the action of FFA agonists. Since we were interested in growth factor action, we routinely use serum-starved cells in which growth factor-induced stimulated signal transduction events, and inhibition of these signals, can be more readily assessed. However, subsequent studies showed that FFA4 agonists also inhibit proliferation of cells that are already actively growing in serum (Figures 8 and 10). Also, since we examined the kinetics of signaling responses, which are best studied in serum-
starved cells, we were able to discern that FFA4 agonists have transient stimulatory effects as well as inhibitory effects. Future studies can examine the extent to which these effects occur in the tumor microenvironment in vivo.

Three previously published studies have indicated that n-3 FAs can suppress signal transduction via activation of FFA4; all of these studies examined inflammatory responses. One group showed that n-3 FAs inhibit inflammasome activation in macrophages [Yan et al., 2013], while another showed that a 1-hour pre-incubation with n-3 FAs inhibits IL-1β response in Caco-2 colon cancer cells [Mobraten et al., 2013]. A third group reported that DHA (24-hour pre-incubation) inhibits lipopolysaccharide responses mediated via Toll-like receptor 4, which include Akt/JNK phosphorylation [Li et al., 2013]. While these results are not directly analogous to the growth inhibitory responses examined in the current study, it is now clear that FFA4 can mediate inhibitory effects on signaling in multiple cellular models.

In contrast to the inhibitory effects discussed above, a recent report indicated that FFA4 plays an oncogenic role in human colon cancer, enhancing angiogenesis as well as migration [Wu et al., 2013]. With regard to signal transduction, Wu and co-workers examined direct effects of GW9508 on two colon cancer cell lines, demonstrating activation of PI3K/Akt and NFκB. As was the case in our prostate cancer cell experiments, Wu and colleagues tested the effects of FFA4 agonists on the proliferation and migration of serum-starved cells. Our data (Figure 3C) similarly show that the FFA4 agonist EPA activates p70S6K in Caco-2 colon cancer cells. The results with colon cancer cells stand in contrast to the results reported herein for prostate cancer cells. Specifically, the FFA1/FFA4 agonist GW9508 is mitogenic for colon cancer cells.
[Wu et al., 2013], but not for prostate cancer cells (Figure 7). GW9508 (10 µM) stimulates migration of colon cancer cells treated for 72 hours prior to the migration assays [Wu et al., 2013], while TUG-891 inhibits migration of prostate cancer cells (Figure 9). It will be important to further explore the mechanisms underlying the opposing effects of FFA4 agonists on proliferation in different types of cancer cells. These differences may reflect variations in the expression levels of FFA4, other GPCRs, G-proteins, or GPCR effectors.

It is interesting to note that DU145 and PC-3 cells differ in many respects with regard to signal transduction. Their profiles of LPA receptor expression and response are different [Gibbs et al., 2009]. PTEN, a tumor suppressor that attenuates activation of the Akt pathway, is expressed in DU145 but not in PC-3 cells [Fraser et al., 2011]. Nonetheless, FFAR agonists have similar anti-proliferative actions in both cell lines. Since FFA4 is expressed in normal as well as transformed prostate cancer cells, we speculate that FFA4 agonists can suppress growth of both pre-cancerous and transformed prostate epithelial cells. This hypothesis requires further investigation.

In our study, CCN1 induction was used to assess a later response to LPA action. CCN1, which is induced in response to various stimuli including growth factors, plays roles in proliferation, apoptosis, and migration of prostate cancer cells [Franzen et al., 2009; Schmitz et al., 2013]. CCN1 expression is induced by LPA in benign prostatic hyperplasia cells [Sakamoto et al., 2004]. While several studies have implicated CCN1 in prostate tumor progression, including studies of DU145 cells [Sun et al., 2008], the roles of this interesting agonistic protein remain to be fully defined. It is possible that CCN-1 induction is important for the second phase of the biphasic signaling response to
LPA, which likely requires the production of additional pro-mitogenic factors. Our results establish an experimental model in which CCN1 production can be inhibited as well as stimulated.

Our demonstration that FFA4 agonists can inhibit growth factor response in an acute and ongoing manner has implications for the administration of n-3 FA supplements or FFAR agonists with the intention of preventing cancer. If the major mechanism for the beneficial effects of n-3 FAs involves changes in phospholipid composition, as previously proposed, then the dosage regimen may be less critical. However, if the mechanism involves modulation of signal transduction pathways and requires the continuous presence of agonist, as our results indicate, then it may be important to administer the agonists more frequently. It will also be important to determine the underlying mechanism. For example, signaling events mediated by arrestins are less susceptible to desensitization, and therefore have a more sustained duration, than those mediated by G proteins [Luttrell and Getsy-Palmer, 2010]. Our results show that continuous engagement of FFA4 is required for inhibitory effects, which persist as long as the agonist is present; these findings suggest arrestin involvement. Translational studies will be needed to determine whether FFA4 agonists suppress mitogenic signaling in tissues, and to define the dosage regimen required for such an effect.

The discovery that a pharmacologic FFA4 agonist can mimic the anti-proliferative effects of an n-3 FA is particularly pertinent to proposed therapeutic approaches for the purposes of preventing or treating cancer. First, the fish oil preparations that are typically used for n-3 FA supplementation are comprised of a mixture of FAs, primarily
EPA and DHA. While these compounds are efficacious as FFA4 agonists, a pharmacologic agent such as TUG-891 can be targeted much more specifically to FFA4, and its pharmacodynamic properties can be more easily defined. Second, the ability of n-3 FA metabolites (“resolvins”) to inhibit inflammation is of considerable interest [Zhang and Spite, 2012]. While inflammation promotes tumor formation, the direct effects of the numerous n-3 FA metabolites on cancer cells are under-studied. Since small molecule FFAR agonists are not metabolized to resolvins, utilization of these agonists will facilitate studies of the relative roles of FFARs to tumor progression. Finally, fish oil supplements are not well tolerated by all individuals. Dranse and colleagues [2013] have provided a thoughtful discussion of dietary versus pharmacologic approaches to targeting FFARs [2013]. FFA4 agonists are already of great interest with respect to the treatment of metabolic diseases [Holliday et al., 2011]. The potential benefit of these agents with respect to cancer provides exciting new avenues for therapeutic intervention.

Acknowledgments

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Authorship contributions

Participated in research design: Meier, Hopkins, Liu, Zhang

Conducted experiments: Liu, Hopkins, Zhang, Quisenberry, Fix, Galvan

Contributed new reagents or analytic tools: --

Performed data analysis: Hopkins, Liu Meier, Zhang, Fix, Galvan

Wrote or contributed to the writing of the manuscript: Meier, Liu, Hopkins, Zhang, Fix
References


C163-C170.


Footnotes

a. This work was supported by the College of Pharmacy at Washington State University, and by a Collaborative Translational Seed Grant [UL1RR025014] to Z.Z. from the Institute of Translational Health Sciences at the University of Washington. C.R.B. was the recipient of a Nutrition and Exercise Physiology Undergraduate Research Fellowship from the WSU College of Pharmacy. M.M.H., a member of the Graduate Program in Pharmaceutical Sciences in the College of Pharmacy, also received support from the NIH Protein Biotechnology Training Program at WSU [T32 GM008336]. B.M.G. was the recipient of a Summer Undergraduate Research Fellowship funded by the American Society for Pharmacology and Experimental Therapeutics and by the College of Pharmacy. Mandi M. Hopkins and Ze Liu contributed equally to this manuscript.

b. Parts of this study comprised the MS dissertation in Nutrition and Exercise Physiology for Ze Liu at Washington State University (2013). Abstracts discussing aspects of this work were presented at the 2013 Southeastern Regional Lipid Conference, and at the 2013 and 2014 Experimental Biology Meeting.

c. Reprint requests should be addressed to: Kathryn E. Meier, PhD, PO Box 1495, College of Pharmacy, Washington State University, Spokane, WA 99210-1495. Email address: kmeier@wsu.edu
Figure legends

**Figure 1:** Effects of EPA and DHA on proliferation of DU145 cells.

DU145 cells were serum-starved overnight and then incubated in duplicate wells for the indicated times in the absence or presence of LPA (10 µM) or 10% fetal bovine serum in the absence or presence of 20 µM EPA or DHA. Live cells were counted. Each data point represents mean ± SEM for values from triplicate experiments. Panel A shows the time course of effects of serum, DHA, and EPA alone. The * indicates p values for control vs serum, the # indicates p values for control vs EPA; control vs DHA had no significant difference. Panel B shows the time course of the effects of LPA, LPA plus DHA, and LPA plus EPA. The * indicates p values for LPA vs LPA+EPA, the # indicates p value for LPA vs LPA+DHA. Panel C presents the results for the 48 hour incubation. All panels show data derived from the same three experiments. Data for panels A and B were analyzed by two-way ANOVA followed by Tukey’s multiple comparisons test; data for panel C were analyzed by one-way ANOVA followed by Tukey’s multiple comparisons test.

**Figure 2:** Effects of EPA on LPA-induced phosphorylation of Erk and FAK in DU145 cells.

Panel A: Serum-starved DU145 cells were incubated with or without 20 µM EPA for 15 minutes. LPA (10 µM) was then added for the indicated times. Whole-cell extracts, equalized for protein, were immunoblotted for phospho-Erk and for actin (loading control). Panel B: Immunoblotting results were quantified from two separate experiments. Panel C: Serum-starved DU145 cells were incubated with or without 20 µM EPA for 15 minutes. LPA (10 µM) was then added for the indicated times.
cell extracts, equalized for protein, were immunoblotted for phospho-FAK and for actin. The actin blot was the same as shown for Panel A, since panels A and C were from the same experiment. Panel D: Immunoblotting results were quantified from two separate experiments. For panels B and D, results were normalized to actin, and then to the untreated control. Each value represents mean ± SEM from duplicate experiments. Data for panels B and D were analyzed by two-way ANOVA followed by Tukey’s multiple comparisons test. Asterisks denote values that are significantly different from untreated control.

Figure 3: Effects of EPA on LPA-induced phosphorylation of p70S6K in DU145 and Caco-2 cells.

Panel A: Serum-starved DU145 cells were incubated with or without 20 µM EPA for 15 minutes. LPA (10 µM) was then added for the indicated times. Whole-cell extracts, equalized for protein, were immunoblotted for phospho-p70S6K and for actin (loading control). Panel B: Immunoblotting results were quantified from experiments in which cells were incubated for the indicated times with EPA, LPA, or EPA plus LPA. Results were normalized to actin, and then to the untreated control. Each value represents mean ± SEM from duplicate experiments, except for EPA alone, which was from a single experiment. Asterisks denote values that are significantly different from untreated control. Panel C: Serum-starved Caco-2 cells were incubated with or without 20 µM EPA or 10 µM LPA for the indicated times. Whole-cell extracts, equalized for protein, were immunoblotted for phospho-p70S6K and for actin (loading control). Immunoblotting results were quantified from three separate experiments. Results were normalized to actin, and then to the untreated control. Each value represents mean ±
SEM from duplicate experiments. When there were two separate experiments for the same time point, the line on the graph was drawn through the mean of the data points.

**Figure 4:** Effects of LPA and EPA on CCN1 expression in prostate cancer cells.

Panel A: Whole cell extracts were prepared from PC-3, DU145, and LNCaP cells growing in serum. The extracts, equalized for protein, were immunoblotted for CCN1 and actin (loading control). Panel B: Serum-starved PC-3 cells were incubated for 4 hours with or without 10 µM LPA in the absence and presence of 10 µM Ki16425, an LPA receptor antagonist. Whole-cell extracts were immunoblotted for CCN1 and for total FAK (loading control). Panel C: Serum-starved PC-3 cells were incubated with 10µM LPA for the indicated times. Whole-cell extracts were immunoblotted for CCN1 and actin (loading control). Panel D: Serum-starved DU145 cells were incubated with or without 20 µM EPA for 15 minutes. LPA (10 µM) was then added for the indicated times. Whole-cell extracts, equalized for protein, were immunoblotted for CCN1 and for actin (loading control). Panel E: Immunoblotting results, from experiments carried out as in panel D, were quantified. Results were normalized to actin, and then to the untreated control. Each value represents mean ± SD from duplicate experiments. Asterisks indicate values that are significantly different from the untreated control (p<0.001). Panel F: The effects of EPA were quantified in experiments in which EPA alone was included as a control. Cells were incubated for 60 minutes with either LPA (10 µM) or EPA (20 µM), or with EPA for 15 minutes prior to incubation with LPA for an additional 60 minutes. Each data point represents mean ± SEM from duplicate experiments.
Figure 5: Effects of EPA on EGF responses in DU145 cells.

In Panel A, the effects of LPA (10 µM), EGF (10 nM), and fetal bovine serum (10%) on cell proliferation were assessed in a time course experiment, as described for Figure 1; results of two separate experiments done in duplicate are shown. In Panel B, the effects of 10 nM EGF on activating phosphorylations of Erk and FAK were examined by immunoblotting, as described for Figures 2 and 3. In Panel C, serum-starved DU145 cells were incubated with and without 20µM EPA for 15 minutes, followed by 10 nM EGF for 1 hour. Each data point represents mean ± SD from duplicate experiments. In Panel D, the effects of EGF and LPA on cell proliferation were assessed after 48 hours, in the presence and absence of 10 µM Ki16425, an LPA receptor antagonist. Each data point represents mean ± SD of values from duplicate experiments, each performed with duplicate samples of cells.

Figure 6: Expression of FFA4 in prostate cancer cell lines.

In Panel A, RT-PCR reactions were carried out using total mRNA prepared from DU145, PC-3, and Caco-2 cells to synthesize first-strand cDNA. The cDNA was then used for PCR using primers specific for human FFA4, FFA1, or Rplp0 (loading control). Products were separated by agarose gel electrophoresis and visualized under UV light. Results shown are representative of three separate experiments, each conducted in triplicate. In panel B, immunoblotting was performed for FFA4 protein and actin (loading control) using whole-cell extracts, equalized for protein, prepared from DU145 and PC-3 cells. Results shown are representative of two separate experiments.

Figure 7: Dose-response for the effects of EPA, TUG-891, and GW9508 on DU145 cell proliferation.
Panel A: Serum-starved DU145 cells were incubated for 48 hours with and without 10 µM LPA in the absence and presence of the indicated concentrations of EPA, DHA, TUG-891, or GW9508. The number of cells achieved in response to LPA in the absence of other additions was defined as 100% response; the number of cells present in the absence of LPA was defined as 0% response. Each point represents mean ± SEM of values obtained from 2-6 experiments, each of which was done using duplicate wells of cells for each experimental condition. Panel B: DU145 cells (not serum-starved) were incubated for 48 hours with and without 10% fetal bovine serum in the absence and presence of the indicated concentrations of EPA (closed triangles), DHA, TUG-891, or GW9508. Each data point represents mean ± SEM of values obtained from 2-3 experiments, each of which was done using duplicate wells of cells for each experimental condition. For both panels, data were fit by nonlinear regression, using GraphPad Prism software, with 0 and 100% response values defined by negative and positive controls within each experiment. Panels C and D: DU145 cells (C) and PC-3 cells (D) were serum-starved, treated with or without 1 µM TUG-891 (C) or 20 µM EPA (D) for 15 minutes, and then treated with or without 10% serum for 24 hours. Cells were then washed, incubated with fresh medium without TUG-891 or EPA (but with serum as indicated), and counted (time “0”). The incubation continued, with live cells being counted every 24 hours cells from 2-4 experiments. Data analysis was done using two-way ANOVA, followed by Tukey’s multiple comparisons test. Panel E: Serum-starved DU145 cells were incubated with or without 20 µM EPA or 1 µM TUG-891 for 15 minutes. LPA (10 µM) or vehicle was then added for 10 minutes. Whole-cell extracts, equalized for protein, were immunoblotted for p-p70S6K and actin. Immunoblots for
p70-S6K were quantified, normalized to actin, and then normalized to the untreated control. Each data point represents mean ± SD of values from duplicate dishes of cells. The results shown are representative of three separate experiments.

**Figure 8:** Effects of TUG-891 on proliferation of DU145 and PC-3 cells.

Proliferation assays were conducted using serum-starved PC-3 (left panels) or DU145 (right panels) cells as described in Figure 1. Cells were incubated with or without fetal bovine serum (10%), LPA (10 µM), or EGF (10 nM) in the absence and presence of EPA (20 µM) or TUG-891 (1 µM) for the indicated times. Each data point represents the mean ± SEM of values from duplicate experiments, each of which used duplicate wells of cells for each experimental condition. The control, EPA alone, and TUG-891 alone data are repeated in the three graphs for each cell line, since all data were from the same series of experiments. Asterisks represent the p values for LPA vs LPA+EPA and # represents the p values for LPA vs LPA+TUG-891; * is p<.05, ** is p<.01, *** is p<.001, and **** is p<.0001. There was no significant difference between control and LPA+EPA or LPA+TUG-891 at any time point. Data analysis was done using two-way ANOVA, followed by Tukey’s multiple comparisons test.

**Figure 9:** Effects of EPA and TUG-891 on migration of DU145 and PC-3 cells.

Serum-starved DU145 and PC-3 cells were seeded in serum-free medium at 2.5 X 10^4 cells per insert in the upper chambers of 8-µm transwell inserts. Serum-free medium was added to the lower wells. Cells were then treated with 10% FBS, 20 µM EPA, 1 µM TUG-891, 10 µM LPA, or 10 nM EGF, either alone or in combination. Following a 6-hour migration period, the insert membranes were fixed and stained using methanol and
crystal violet. Cells that invaded the lower chambers were counted by microscopy. Each bar represents the mean ± SEM of values obtained from duplicate experiments, each of which used duplicate wells of cells for each experimental condition. Data analysis was done using one-way ANOVA, followed by Tukey’s multiple comparisons test.

**Figure 10:** Effects of FFA4 knockdown on inhibitory effects of EPA and TUG-891 on cell proliferation.

Panel A: DU145 cells were incubated with scrambled siRNA (negative control) or FFA4 siRNA as described in Methods. Whole-cell extracts were immunoblotted for FFA4 and actin (loading control). The percent knockdown (“KD”) was calculated after quantification by densitometry, and normalization to actin. Panels B and C: DU145 cells (B) and PC-3 cells (C) were incubated with scrambled siRNA (“Scr”) or FFA4 siRNA (“F4i”), or without siRNA (“control”) as described in Methods. Simultaneously, cells were treated with and without 20 µM EPA or 1 µM TUG-891, in the continued presence of 10% serum. Panels D-G: DU145 cells (D and E) and PC-3 cells (F and G) were treated as for Panels B and C, except that serum was removed for 24 hours prior to the addition of scrambled or FFA4 siRNA, without or without 20 µM EPA, 1 µM TUG-891, 10 µM LPA, 10 nM EGF, or vehicle as indicated. For Panels B-G, data points represent the mean ± SEM of values from duplicate experiments, each conducted in duplicate (n=4). Asterisks indicate values that were significantly different (two-way ANOVA followed by Tukey’s multiple comparisons test) from the untreated control, as well as from the scrambled siRNA control, for the time point tested.
Figure 11: Effects of FFA4 knockdown on inhibitory effects of EPA and TUG-891 on cell migration.

Serum-starved DU145 cells (Panels A and B) and PC-3 cells (Panels C and D) were incubated with scrambled siRNA (“Scr”) or FFA4 siRNA (“F4i”), or without siRNA (“control”) as described in Methods. After 24 hours, cells were seeded in a Boyden chamber and treated with and without 20 µM EPA or 1 µM TUG-891, with or without 10 µM LPA or 10 nM EGF. Migration experiments were carried out for 6 hours. Data points represent the mean ± SEM of values from duplicate experiments, each conducted in duplicate (n=4). Asterisks indicate values that were significantly different (two-way ANOVA followed by Tukey’s multiple comparisons test) from the untreated control, as well as from the scrambled siRNA control, for the time point tested.
Figure 1.

A

Cell Number (x 10,000)

Time (hrs)

Control

Serum

DHA

EPA

B

Cell Number (x 10,000)

Time (hrs)

Control

LPA

LPA+DHA

LPA+EPA

C

Cell Number (x 10,000)

Control

Serum

EPA

DHA

LPA

LPA+DHA

LPA+EPA

*p < 0.01

*p < 0.001

*p < 0.0001

# p < 0.01

## p < 0.001

### p < 0.0001

#### p < 0.0001
Figure 2.

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B

Time (min)

phospho-Erk (normalized to control)

C

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D

Time (min)

phospho-FAK (normalized to control)
Figure 3.

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B

DU145

![Graph showing time (min) vs. phosphoryl-p70S6K normalized to control for DU145 cells under different conditions.](image)

C

Caco-2

![Graph showing time (min) vs. phosphoryl-p70S6K normalized to control for Caco-2 cells under different conditions.](image)
Figure 4.

A

PC-3  LNCaP  DU145

CCN1  actin

B

PC-3

FAK  CCN1

Control  LPA  K16425  K16425+LPA

C

PC-3

CCN1  actin

Time (hrs)  0  0.5  1  2  4  8  16  24

D

DU145

LPA (min)  EPA (20µM)  0  10  30  60  120  240  360

CCN1  actin

E

DU145

CCN1 (normalized to control)

Time (min)  0  60  120  180  240  300  360

F

DU145

CCN1 (normalized to control)

Control  EPA  LPA  LPA+EPA

p<0.05
Figure 5.

A

Cell Number (x 100,000)

Time (hrs)

Control
Serum
LPA
EGF

B

Control  EGF  EGF+EPA

p-Erk

p-FAK

actin

C

CCN1 (normalized to actin)

Control  EGF  EGF+EPA

p<0.05

D

Cell Number (x 100,000)

Control  LPA  EGF

p<0.01  p<0.01

Control  +Ki16425
Figure 6.

A

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This article has not been copyedited and formatted. The final version may differ from this version.
**Figure 7.**

A. LPA

![Graph showing LPA-induced Proliferation](image)

- EPA
- DHA
- TUG-891
- GW9508

B. Serum

![Graph showing Serum-induced Proliferation](image)

- EPA
- DHA
- TUG-891
- GW9508

C. TUG-891

![Graph showing Cell Number](image)

- Serum+TUG-891
- LPA+TUG-891
- EGF+TUG-891

D. EPA

![Graph showing Cell Number](image)

- Serum+EPA
- LPA+EPA
- EGF+EPA

E. Graph showing p<0.01 and p<0.01

- Control
- LPA
- EPA
- LPA+EPA
- TUG891
- LPA+TUG891
Figure 8.

PC-3
Serum

DU145
Serum

LPA

LPA

EGF

EGF
Figure 9.

A

DU145

Migrated Cells

Control  EPA  TUG891  LPA  LPA+EPA  LPA+TUG891  EGF  EGF+EPA  EGF+TUG891

B

PC-3

Migrated Cells

Control  EPA  TUG891  LPA  LPA+EPA  LPA+TUG891  EGF  EGF+EPA  EGF+TUG891
Figure 10.

A  

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</table>
%KD:    | 92.9| 61.9|

B  

DU145 with serum

Control | Scr | Scr+EPA | Scr+TUG891 | F4i | F4i+EPA | F4i+TUG891

C  

PC-3 with serum

Control | Scr | Scr+EPA | Scr+TUG891 | F4i | F4i+EPA | F4i+TUG891

D  

DU145 with LPA

E  

DU145 with EGF

F  

PC-3 with LPA

G  

PC-3 with EGF
Figure 11.

A. DU145 with LPA

B. DU145 with EGF

C. PC-3 with LPA

D. PC-3 with EGF