INTRODUCTION

Prolonged inflammation underlies the pathogenesis of various diseases including cancer. Recent years have seen a paradigm shift in our understanding of the etiopathogenesis of inflammation whereby chronic inflammation may not result only from persistence of inflammatory mechanisms, but may also arise from an inadequate engagement of the resolution mechanisms. Resolution is an active process brought about by a series of cellular mechanisms that are activated to restore tissue homeostasis. Lipid specialized pro-resolving mediators (SPMs), which are the main effectors of resolution, derive from the effects exerted by lipoxigenases (ALOX5 and 15) on ω-6 arachidonic acid (AA), or on ω-3 eicosapentaenoic acid (EPA) and docosahexaenoic acid (DHA), which are essential polyunsaturated fatty acids (PUFA) (Fig. 1A). Recent evidence associates a deficit in pro-resolving pathways with the establishment of various inflammatory-related disorders (e.g., asthma, fibrosis, and autoimmune diseases). However, whether and how defects in resolution mechanisms affect cancer initiation and progression, and what signals control pro-resolving pathways in this context is unknown.

Formyl peptide receptors (FPR1, 2, and 3) are pattern recognition receptors (PRR) of the G-protein-coupled (GPCR) family that recognize both exogenous and endogenous “danger” signals, and trigger inflammation and immune responses. FPRs can also trigger inflammation resolution, depending on the environmental context and on the specific ligand. We recently showed that genetic ablation of FPR1 in AGS gastric cancer (GC) cells, which constitutively express high levels of the receptor, increased their angiogenic and tumorigenic potential. Accordingly, enforced expression of FPR1 in MKN45 GC cells, which constitutively express high levels of the receptor, dramatically impaired GC cell xenograft angiogenesis and growth in immunodeficient mice.

Since genetic deletion of FPR1 in GC cells increased angiogenesis and enhanced the response to pro-inflammatory cytokines, which is a phenotype suggestive of unresolved inflammation, we asked whether, at the gastric level, FPR1 might actively sustain pro-resolving pathways to inhibit GC angiogenesis and growth.

DISCUSSION

Here, we demonstrate that genetic or pharmacologic modulation of FPR1 in GC cells regulated ALOX5/15 expression and production of the SPMs Resolvin D1 (RvD1) and Lipoxin B4 (LXB4). SPM treatment of GC cells abated their angiogenic potential. Genetic deletion of ALOX15 or of the RvD1 receptor GPR32 increased the angiogenic and tumorigenic activity of GC cells thereby mimicking FPR1 loss. Deletion/inhibition of ALOX5/15 or GPR32 blocked FPR1-mediated anti-angiogenic activities, indicating that ALOX5/15 and GPR32 are required for FPR1’s pro-resolving action. An ω-3- or ω-6-enriched diet enforced SPM endogenous production in mice and inhibited growth of shFPR1 GC xenografts by suppressing their angiogenic activity. These data implicate that FPR1 and/or pro-resolving pathway components might be used as risk/prognostic markers for GC; ω-6/3-enriched diets, and targeting FPR1 or SPM machinery may be exploited for GC management.

ABSTRACT

Chronic inflammation can result from inadequate engagement of resolution mechanisms, mainly accomplished by specialized pro-resolving mediators (SPMs) arising from the metabolic activity of lipoxigenases (ALOX5/15) on ω-6 or ω-3 essential polyunsaturated fatty acids (PUFA). We previously demonstrated that formyl peptide receptor 1 (FPR1) suppresses gastric cancer (GC) by inhibiting its inflammatory/angiogenic potential. In this study, we asked whether FPR1 exploits inflammation resolution pathways to suppress GC angiogenesis and growth. Here, we demonstrate that genetic or pharmacologic modulation of FPR1 in GC cells regulated ALOX5/15 expression and production of the SPMs Resolvin D1 (RvD1) and Lipoxin B4 (LXB4). SPM treatment of GC cells abated their angiogenic potential. Genetic deletion of ALOX15 or of the RvD1 receptor GPR32 increased the angiogenic and tumorigenic activity of GC cells thereby mimicking FPR1 loss. Deletion/inhibition of ALOX5/15 or GPR32 blocked FPR1-mediated anti-angiogenic activities, indicating that ALOX5/15 and GPR32 are required for FPR1’s pro-resolving action. An ω-3- or ω-6-enriched diet enforced SPM endogenous production in mice and inhibited growth of shFPR1 GC xenografts by suppressing their angiogenic activity. These data implicate that FPR1 and/or pro-resolving pathway components might be used as risk/prognostic markers for GC; ω-6/3-enriched diets, and targeting FPR1 or SPM machinery may be exploited for GC management.
Here, we show that, in GC cells, FPR1 expression/activation levels directly correlate with ALOX expression, and with SPM Resolvin D1 (RvD1) and Lipoxin B4 (LXB4) (3) production (Fig. 1A). The increased angiogenic potential of GC cells lacking FPR1, previously reported,7 was reverted by RvD1 or LXB4 treatment. Genetic ablation of ALOX15 or of GPR32 (an RvD1 receptor)8 induced a pro-angiogenic phenotype in GC cells similar to that induced by FPR1 deletion. ALOXs and GPR32 are required for FPR1-mediated anti-angiogenic activity in GC cells. Consistently, administration of ω-3 or ω-6 PUFA-enriched diets, which enforces endogenous production of SPMs,9 inhibited xenograft growth of FPR1-silenced GC cells by abating their angiogenic activity.

Figure 1. Relation between FPR1 expression and SPM biosynthesis machinery. (A) Schemes of arachidonic acid-AA, docosahexaenoic acid-DHA, and eicosapentaenoic acid-EPA metabolism. (B) AGS shFPR1 cl 15 cells produced significantly lower amounts of RvD1 and LXB4 compared to AGS shCTR cells, while MKN45 cells overexpressing FPR1 (MKN45 FPR1 cl 4) released higher amounts of RvD1 and LXB4 compared to empty vector transfected cells (MKN45 pcDNA), as evaluated by EIA assays. Data are represented as mean ± SD of five independent experiments. *p < .05 compared to the relative control. (C) Increased release of PGE2 and LTB4 from shFPR1 AGS cells and from MKN45 pcDNA cells compared to the relative control cells, assessed by EIA. Data are represented as mean ± SD of three independent experiments. *p < .05 compared to the relative control. (D) AGS shFPR1 cl 15 expressed significantly lower levels and MKN45 FPR1 cl 4 significantly higher levels of ALOX5, ALOX15A, and ALOX15B mRNAs compared to relative controls (dotted line), as assessed by real-time PCR. Data are represented as mean ± SD of three independent experiments. *p < .05 compared to the relative control. (E) ALOX5, ALOX15A, and ALOX15B protein levels were lower in AGS shFPR1 vs. AGS shCTR, and in MKN45 pcDNA vs. MKN45 FPR1 cells, as evaluated by cytofluorimetric analysis. One representative experiment out of three is shown.
Our data indicate that FPR1 signaling activates a pro-resolving program in GC cells that inhibits angiogenesis and growth.

Results

**FPR1 controls ALOX5 and ALOX15 expression and the production of SPMs in GC cells**

To study whether pro-resolving pathway components are involved in the FPR1-mediated anti-angiogenic and tumor suppressor activity of GC, we used our previously generated GC cell lines, namely, FPR1-silenced AGS (AGS shFPR1) and MKN45 ectopically expressing FPR1 (MKN45 FPR1) or their relative controls (AGS shCTR and MKN45 pcDNA).7

RvD1 and LXB4 levels were significantly lower in AGS shFPR1 cells than in AGS shCTR cells. Consistently, MKN45 cells overexpressing FPR1 released higher amounts of RvD1 and LXB4 than did empty vector-transfected cells (Fig. 1B). RvD1 and LXB4 syntheses were reduced in shFPR1 but not in shFPR2 or shFPR3 AGS cells (not shown), which indicates that FPR1 plays a non-redundant role in controlling SPM production, as already observed for

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**Figure 2.** Effects of FPR1 pharmacologic modulation on SPM biosynthesis. (A) EIA assays showing that fMLF (agonist to FPR1, 10^{-9} M) treatment of AGS significantly increases RvD1 and LXB4 release compared to untreated cells (dotted line). In contrast, CsH (inverse agonist to FPR1, 800 nM) treatment significantly reduced RvD1 and LXB4 release compared to controls (dotted line). Data are represented as mean ± SD of three independent experiments. *p < .05 compared to untreated cells. (B) fMLF induced, whereas CsH inhibited, ALOX5, ALOX15A, and ALOX15B mRNA expression in AGS and MKN45 cells. Data are represented as mean ± SD of three independent experiments. *p < .05 compared to untreated cells (dotted line). (C) fMLF induced, whereas CsH inhibited, ALOX5, ALOX15A, and ALOX15B protein expression in AGS cells, as evaluated by cytofluorimetric analysis. One representative experiment out of three is shown. (D) fMLF inhibited, whereas CsH induced, pro-angiogenic molecule mRNAs expression in AGS and MKN45 cells. Data are represented as mean ± SD of three independent experiments. *p < .05 compared to untreated cells (dotted line).
its tumor suppressor function. We evaluated the lipidomic profiles of controls and FPR1-depleted/overexpressing GC cell supernatants by liquid chromatography tandem-mass spectrometry (LC-MS/MS) (Fig. S1). Compared to controls, AGS shFPR1 cells produced significantly lower levels of SPMs, including 18-HEPE, a resolvin precursor, and LXB4 (Fig. S1A). MKN45 FPR1 cells released significantly higher levels of RvD3 and LXB4 than did control cells (Fig. S1A). LXA4 levels did not differ significantly among GC cells expressing different levels of FPR1 (Fig. S1B). SPM expression parallels the downregulation of pro-inflammatory eicosanoids. Consistently, AGS shFPR1 cells synthesized significantly higher amounts of LTB4 and PGE2 compared to shCTR cells, whereas MKN45 FPR1 cells released lower levels of PGE2 compared to empty vector transfected cells, as assessed by EIA (Fig. 1C).

The levels of the enzymes involved in SPM synthesis (ALOX5, ALOX15A, and ALOX15B) were significantly lower in FPR1-silenced GC cells than in shCTR cells, both at mRNA (Fig. 1D) and protein (Fig. 1E) level. The expression of SPM receptors GPR32, ChemR23, and BLT1 was lower in AGS shFPR1 cells than in controls (Fig. S1C and D). On the other hand, MKN45 FPR1 cells produced significantly higher levels of pro-resolving enzymes (Fig. 1D and E).
E) and of SPM receptors (Fig. S1C and D). These data were reproduced on one more clone and a mass population of both AGS shFPR1 and MKN45 FPR1 GC cells (Fig. S2).

To approach the FPR1-SPMs association from a pharmacologic standpoint, we treated AGS and MKN45 cells with fMLF (10⁻⁹ M), which is an FPR1 agonist, or with cyclosporine H (CsH 800 nM), which is an FPR1 inverse agonist¹² and verified their ALOX5/15 expression, SPM release, and synthesis of pro-angiogenic mediators (VEGF-A, -B, -C, -D, Ang1, and CXCL1). The release of RvD1 and LXB4 was significantly higher in AGS treated with fMLF (12 h) than in untreated cells. In contrast, RvD1 and LXB4 release was significantly lower in cells treated with CsH (12 h) than in controls (Fig. 2A). fMLF induced, whereas CsH reduced, ALOX5, ALOX15A, and ALOX15B mRNA expression in AGS and MKN45 cells (Fig. 2B). These observations were confirmed at the protein level (Fig. 2C). mRNA levels of pro-angiogenic molecules were significantly reduced by fMLF and induced by CsH treatment (Fig. 2D).

**SPMs inhibit the production of pro-angiogenic factors and restore the expression of ALOX5/15 and of SPM receptors in GC cells**

To assess whether SPMs (RvD1 and LXB4) block the pro-angiogenic activity of GC cells, we treated or not AGS cells with RvD1 (1 nM) or LXB4 (1 nM), and evaluated the levels of pro-angiogenic factor mRNAs. RvD1 and LXB4 suppressed pro-angiogenic factor mRNA expression in GC cells (Fig. 3A). RvD1 treatment of GC cells significantly reduced VEGF-A release both in basal conditions and upon WKYMVm stimulation (pro-inflammatory FPR2/3 ligand) (Fig. 3B)⁷ Similar results were obtained with LXB4 (not shown). This effect was significantly more efficient in GC cells expressing low levels of FPR1 (AGS shFPR1 and MKN45 pcDNA) (Fig. 3A and B). Interestingly, RvD1 treatment restored, and in some cases increased above basal levels, the expression of ALOX5, ALOX15A, ALOX15B, and the SPM receptors (GPR32, ChemR23, and BLT1) at

![Figure 4](https://example.com/figure4.png)

**Figure 4.** ALOXs and GPR32 involvement in GC angiogenic response. (A) Tumor growth curves of AGS shCTR, shALOX5 (three clones) and shALOX15 (three clones) xenografts in immunodeficient mice. *p < .05 compared to shCTR xenografts. (B) Representative images and quantification (five fields/sample) of the proliferation index (Ki-67), vessel density (CD31), and apoptotic rate (Cleaved Caspase 3), assessed by immunohistochemistry, of shCTR, shALOX5, and shALOX15 cell xenografts harvested 28 d post-inoculation. *p < .05 compared to shCTR xenografts. (C) ALOX15 and ALOX5 mRNA expression levels of 295 patients affected by gastric adenocarcinoma stratified for disease-free and overall survival status. *p < .05 between the two groups. (D) Tumor growth curves of AGS shCTR and shGPR32 xenografts (average of three clones) cells in immunodeficient mice *p < .05 compared to shCTR xenografts. (E) Representative images of the vessel density (CD31), assessed by immunohistochemistry, of shCTR and shGPR32 AGS cell xenografts harvested 28 d post-inoculation.
mRNA (Fig. 3C) and protein level (Fig. 3D and E) thereby triggering a positive feed-forward loop. These responses of AGS shCTR to RvD1 were significantly less efficient than that of AGS shFPFR1 cells (Fig. 3C). Similar results were obtained with LXb4 (not shown).

**ALOX15- and GPR32-silencing increase angiogenesis and tumorigenesis of GC cells thereby mimicking the FPR1-silencing phenotype**

To evaluate whether SPM pathway components are necessary to control the angiogenic activity of GC cells, we stably transfected AGS cells with vectors expressing shRNAs targeting ALOX5, ALOX15, GPR32, or control non-targeting shRNAs (shCTR). ALOX5-silencing induced a specific reduction of ALOX5 mRNA levels in AGS cells. However, AGS shALOX5 cells overexpressed ALOX15A and ALOX15B mRNAs (Fig. S3A). Transfection of shRNAs targeting ALOX15 significantly reduced ALOX15A and ALOX15B levels in AGS cells, but did not modify ALOX5 mRNA (Fig. S3A). The effects of shRNAs were confirmed at the protein level (Fig. S3B). shALOX5 and, to a lesser extent shALOX5, impaired AGS cell growth and survival, as did FPR1 silencing (not shown). shALOX5 and shALOX15 AGS cells (three clones/each) constitutively produced lower amounts of RvD1 and increased amounts of PGE2 and LTb4 versus controls, as did shFPFR1 cells. LXb4 release was reduced in shALOX5, but not in shALOX5 cells (Fig. S3C). GPR32 and ChemR23 protein expressions were lower in AGS shALOX5 and shALOX15 cells than in controls (not shown). AGS shALOX5 and shALOX15 cells constitutively synthesized higher levels of VEGF-C and VEGF-D mRNAs and released increased amounts of VEGF-A than did controls (Fig. S3D).

To evaluate their tumorigenic potential, we injected AGS shCTR cells or ALOX-silenced clones subcutaneously in athymic mice. AGS shCTR and shALOX5 tumors reached a volume of about 1 mm³ at day 7, and regressed thereafter (Fig. 4A). In contrast, AGS shALOX15 cells formed larger non-regressing tumors, i.e., 2.8 mm³ (Fig. 4A), again replicating the phenotype of FPR1-silenced cells. The tumor growth advantage of shALOX15 cells was due to their higher vessel density compared to controls (Fig. 4B). No differences were detected in Ki-67+ and cleaved caspase 3+ cells in shCTR and shGPR32 xenografts (not shown). The low levels of GPR32 and ALOXs, evaluated far from the necrotic areas, among shCTR, shALOX5, and shALOX15 cell xenografts (Fig. 4B). Our in vivo observations are corroborated by data, publicly available in the cBioPortal for Cancer Genomics database (http://www.cbioportal.org),13,14 reporting the RNAseq analysis of 295 human GC samples. These data show that enhanced mRNA expression of ALOX15, but not of ALOX5, was significantly associated with overall survival status and disease-free status in GC patients (Fig. 4C).

To analyze the effects of GPR32 silencing on the GC cell phenotype, we selected various AGS shGPR32 clones. The low levels of GPR32 were associated with a significant reduction of growth ability and with an increased apoptotic rate of GC cells in culture (not shown). The reduced level of GPR32 in AGS GC cells caused a significant reduction of ALOX5 and ALOX15A/B expression levels (Fig. S3E) and an increased expression of pro-angiogenic mediators at mRNA (Fig. S3E) and protein (Fig. S3F) level vs. shCTR cells. RvD1 did not reduce angiogenic factor mRNAs in shGPR32 cells (Fig. S3G). When xenotransplanted into nude mice, AGS shGPR32 formed significantly larger tumors than formed by shCTR cells (Fig. 4D). Consistently, the vessel density of shGPR32 xenografts was significantly higher than that of controls (Fig. 4E). No differences were detected in Ki-67+ and cleaved caspase 3+ cells in shCTR and shGPR32 xenografts (not shown).

**FPR1-mediated suppression of angiogenesis in GC cells requires ALOX5, ALOX15, and GPR32**

To assess whether FPR1-mediated suppression of angiogenesis requires ALOX5 and/or ALOX15, we stimulated with fMLF (10⁻⁹ M) GC cells in which ALOX expression/activity was abolished. As shown in Fig. 5A, fMLF significantly reduced the mRNA levels of pro-angiogenic factors (VEGF-B, VEGF-C, and CXCL1) in AGS shCTR, but not in shALOX5 and shALOX15 AGS cells. Similar results were obtained in AGS cells in which ALOXs were pharmacologically inhibited by 10 μM nordihydroguaiaretic acid-(NDGA) (Fig. 5B).16 These observations suggest that ALOX5 and ALOX15 are required to trigger the anti-angiogenic activity of FPR1 in GC cells.

To verify the requirement of the RvD1 pathway for FPR1-mediated anti-angiogenic activity, we stimulated parental GC cells with fMLF in the presence or absence of a GPR32-neutralizing antibody. fMLF (10⁻⁹ M) induced ALOX5 and ALOX15A mRNA expression and concomitant VEGF-B and CXCL1 mRNA downregulation in AGS. These effects were reverted by the GPR32-blocking antibody (5 μg/mL) (Fig. 5C). Similar results were obtained in MKN45 cells (Fig. 5A). Consistently, fMLF did not induce ALOXs or suppress pro-angiogenic mediators in AGS shGPR32 cells (Fig. 5D), although shCTR and shGPR32 cells released comparable amounts of RvD1 both in basal conditions and upon fMLF stimulation (not shown). We then assessed the involvement of FPR2, the other RvD1 receptor.8 fMLF stimulation of AGS shFPFR2 cells induced ALOX5 and ALOX15A mRNA expression, and VEGF-B and CXCL1 downregulation, and these effects were reverted by the anti-GPR32 blocking antibody (Fig. 5E). Taken together, these results indicate that RvD1, by binding GPR32, but not FPR2, mediates FPR1’s anti-angiogenic effects in GC cells.

We then asked which signaling pathways were involved in the anti-angiogenic response of GC cells to RvD1 and whether differences can be detected in FPR1-lacking cells compared to controls. We found that Akt, MAPK, JNK, and p38 signaling pathways were involved in GC cells treated with RvD1 (1 nM) for different times (15 min–12 h) without significant differences in the activation kinetic and strength between AGS shCTR and shFPFR1 cells, or MKN45 pcDNA and FPR1 cells (Fig. 5F). In contrast, RvD1 treatment induced a significant STAT3 and SRC activation only in AGS shFPFR1 and MKN45 pcDNA cells expressing lower levels of FPR1 compared to the relative control cells (Fig. 5F). RvD1-mediated reduction of VEGF-
A release was reverted by pre-treatment of AGS shFPR1 cells with the two STAT3 inhibitors FLLL31 (10 μM) and 5–15DPP (15 μM) (Fig. 5F). Similar results were not detectable in AGS shCTR cells (Fig. 5F). STAT3 inhibition reverted RvD1 effects on CXCL1 and Ang1 mRNAs levels (Fig. S4B). Instead, when MAPK [U0 126 (25 μM)] or Akt [LY294002 (15 μM)] inhibitors were used, no effects on VEGF-A release were observed (Fig. S4C).

**ω-6 and ω-3 diets reduce the tumorigenic potential of shFPR1 GC cells in vivo**

Dietary interventions based on differential PUFA ω-3 or ω-6 intake affect AA-, EPA-, and DHA metabolism and the derived SPM endogenous production in mice (Fig. 1A). To assess the effects exerted by ω-6- or ω-3-enriched diets on GC cell growth, we xenotransplanted AGS shCTR or shFPR1 cells in

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**Figure 5.** ALOXs and GPR32 are required for FPR1 tumor suppressor role. (A) VEGF-B, -C, and CXCL1 mRNA synthesis was inhibited in shCTR, but not in shALOX5 or shALOX15 AGS cells upon fMLF (10⁻⁹ M) treatment. Data are represented as mean ± SD of three independent experiments. *p < .05 vs. the relative untreated cells (dotted line). (B) Pre-treatment of AGS cells with NDGA (ALOX inhibitor) reverted the ability of fMLF to inhibit the mRNA synthesis of pro-angiogenic molecules. Data are represented as mean ± SD of three independent experiments. *p < .05 compared to fMLF treated cells. (C) In AGS cells, fMLF induced ALOX5 and ALOX15A mRNA overexpression and concomitant VEGF-B and CXCL1 mRNA downregulation. These effects were reverted by a neutralizing GPR32 antibody. An isotype-matched antibody was used as a control. Data are represented as mean ± SD of three independent experiments. *p < .05 compared to isotype-matched control. (D) fMLF significantly induced ALOX5, ALOX15A, and ALOX15B mRNA levels and significantly reduced the mRNA levels of pro-angiogenic mediators in AGS shCTR, but not in shGPR32 cells. Data are represented as mean ± SD of three independent experiments. *p < .05 compared to untreated cells (dotted line). (E) A neutralizing GPR32 antibody inhibited fMLF-induced ALOX5 and ALOX15A mRNA overexpression and concomitant VEGF-B and CXCL1 mRNA downregulation in AGS shFPR2 cells. An isotype-matched antibody was used as a control. Data are represented as mean ± SD of three independent experiments. *p < .05 compared to untreated cells (dotted line). (F) Activation kinetics of AKT, MAPK, JNK, p38, STAT3, and SRC in AGS shCTR, AGS shFPR1, MKN45 pcDNA, MKN45 FPR1 cells assessed by western blot for their phosphorylated forms. Two STAT3 inhibitors (5-15 DPP and FLLL31) reverted the anti-angiogenic effects of RvD1 in AGS shFPR1 cells. Data are represented as mean ± SD of three independent experiments. *p < .05 compared to untreated cells (NT). **p < .05 compared to RvD1 treated cells.
immunocompromised mice randomly fed a classic, an \( \omega-6/\omega-3 \) balanced (grape seed-colza oils 50/50%, CTRL), an \( \omega-6 \) enriched (grape seed oil 100%, \( \omega-6 \)), or an \( \omega-3 \) enriched (colza/fish oils 80/20%, \( \omega-3 \)) diet, 2 weeks before and 4 weeks after xenotransplantation. No differences were detected between animals fed a classic diet and those fed a balanced diet (not shown) demonstrating that a 5% PUFA-enriched diet did not change the tumorigenic potential of GC cells. As shown previously,\(^7\) AGS shCTR cells formed small tumors that regressed in mice fed a balanced, \( \omega-3 \)-, or \( \omega-6 \)-enriched diet, without statistical differences among groups (Fig. 6A). In contrast, AGS shFPR1 cells formed tumors that progressively increased in size (Fig. 6A).\(^7\) Both \( \omega-3 \) and \( \omega-6 \)-enriched diets significantly suppressed AGS shFPR1 xenograft growth. However, growth inhibition was significant at day 7 post-injection in \( \omega-6 \)-fed mice vs. day 14 in \( \omega-3 \)-fed mice (Fig. 6A). The administration of an \( \omega-3 \) or \( \omega-6 \)-enriched diet to AGS shFPR1 xenotransplanted mice abolished statistical differences in tumor growth curves between shFPR1 and shCTR cells.

Vessel density was higher in shFPR1 xenografts than in shCTR xenografts (Fig. 6B); \( \omega-6 \) and \( \omega-3 \) diets reverted this difference (Fig. 6B). Vessel density was unchanged in shCTR xenografts irrespective of the diet administered (Fig. 6B). Both \( \omega-6 \) and \( \omega-3 \) diets significantly reduced Ki-67 staining of shFPR1, and, to a lesser extent, of shCTR excised tumors (Fig. 6C). Diet composition did not affect apoptotic rate as assessed by cleaved caspase-3 evaluation (not shown).

To investigate the effect of dietary intervention on mouse serum lipid composition, we profiled fatty acids by LC-MS/MS. Concentrations of AA were significantly higher in mice fed an \( \omega-6 \) diet than in those fed a balanced or an \( \omega-3 \) diet (Table 1). Consistently, EPA ad DHA concentrations were higher in \( \omega-3 \)-fed mice than in the other two groups of mice (Table 1). The concentrations of LXB4 were significantly higher in the sera of \( \omega-6 \)-fed mice than in controls. Concentrations of RvD1, RvD3, and RvE2 were significantly higher in the serum of mice fed an \( \omega-3 \) diet than in controls. Interestingly, the \( \omega-3 \) diet significantly reduced PGE2, LTB4, and TXB2 serum concentrations vs. controls (Table 1). Lipid serum concentrations did not differ among mice fed the same diet but xenotransplanted with shCTR or shFPR1 AGS cells (not shown).

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**Figure 6.** Effects of \( \omega-6 \) and/or \( \omega-3 \) increased consumption on GC growth/angiogenesis in a xenotransplantation mouse model. (A) Tumor growth of AGS shCTR and shFPR1 xenografts in CD1 nu/nu mice fed an \( \omega-6/\omega-3 \) balanced (grape seed-colza oils 50/50%, CTRL), an \( \omega-6 \) enriched (grape seed oil 100%, \( \omega-6 \)), or an \( \omega-3 \) enriched (colza/fish oils 80/20%, \( \omega-3 \)) diet. \( *p < .05 \) vs. CTRL diet. (B) Vessel density (CD31) assessed by immunohistochemistry of shCTR and shFPR1 cell xenografts harvested 28 d post-inoculation from mice fed the three diets as in panel A. Representative images and the relative quantifications (five fields/sample) are shown. \( *p < .05 \) vs. the relative CTRL diet. (C) Proliferation index (Ki-67) assessed by immunohistochemistry of shCTR and shFPR1 cell xenografts harvested 28 d post-inoculation from mice fed with the three diets. Representative images and the relative quantifications (five fields/sample) are shown. \( *p < .05 \) vs. the relative CTRL diet.
To assess whether the effects exerted by an ω-3 or ω-6 diet on the proliferation rate of GC xenografts was a direct event or whether it was indirectly due to differences in tumor vascularization levels, we studied the effects of RvD1 (1 nM), LXB4 (1 nM), AA (20 μM), DHA (20 μM), and EPA (20 μM) on AGS shCTR and shFPR1 cell proliferation in vitro. We found no difference in BrdU incorporation between treated and untreated cells at the concentrations of lipids above mentioned (Fig. S5). AA, DHA, and EPA significantly reduced GC cell viability when used at higher concentrations (50 μM), as already reported by others. However, these concentrations were never reached in vivo in diet-fed mice.2,22 (Table 1).

**Discussion**

The role of FPR1 in cancer is complex. In certain cancer types, FPR1 exerts tumor-promoting activities by stimulating the motility and growth of cancer cells. However, in mouse liver and colon, FPR1- or FPR2-genetic deletion abolishes epithelial homeostasis, increases inflammation after injury and, in colonic mucosae, increases tumorigenesis. Recent studies showed that decreased FPR1 concentrations in tumor-infiltrating antigen-presenting cells compromise chemotheraphy-induced antitumor immunity. In human GC, high FPR1 expression, evaluated by immunohistochemistry, has been correlated with advanced tumor stage and poor overall survival. However, an FPR1 polymorphism linked to reduced FPR1 activity has been associated with an increased risk of stomach cancer in humans.

Here, we provide evidence that FPR1, expressed by GC cells, plays a tumor suppressor role by constitutively activating an inflammation resolution program that includes the expression of ALOX5/15, SPMs (RvD1 and LXB4), and SPM receptors (BLT1, ChemR23, and GPR32). FPR1 loss/inhibition suppresses this program and induces an inflammatory/angiogenic phenotype in GC cells. Interestingly, the GC-associated FPR1 polymorphism was also associated with periodontitis, a disease whose pathogenesis has been linked to defective pro-resolving pathways and in which SPMs exert protective effects.

We demonstrate that the anti-angiogenic potential of SPMs, already reported in other contexts, was more evident in cells expressing low levels of FPR1, despite their lower levels of SPM receptors (BLT1, GPR32, and ChemR23), vs. controls. This effect might be due to constitutively high levels of RvD1 and LXB4 produced by control cells, which could saturate/desensitize their receptors thereby rendering these cells less responsive to exogenous SPM stimulation.

Various studies have recognized that ALOXs exert tumor-promoting or tumor-suppressive activities. Our study introduces a new concept in this field, namely, that another component of pro-resolving pathways, i.e., the Resolvin D1 receptor GPR32, can exert tumor suppressor activity in GC. The LX4 receptor remains unidentified; consequently, we were unable to investigate its role in GC. We demonstrate that the ALOX enzymes and GPR32 are induced by FPR1 and necessary for FPR1-mediated SPM production and anti-angiogenic activity in GC. In vivo, vessel density and tumor growth were higher in shALOX5 and shGPR32, but not in shALOX5, xenografts than in controls, despite their comparable angiogenic potential in culture. This discrepancy could be explained by the compensatory upregulation of ALOX15 expression in shALOX5 cells that enabled the production of other SPMs, including LXB4. Thus, qualitative or quantitative differences in SPM production between shALOX5 and shALOX5 cells and/or their interaction with tumor stroma might result in a different angiogenic potential in vivo. Be that as it may, in line with our data, an RNAseq analysis showed that higher ALOX15, but not ALOX5, mRNA expression was significantly associated with better overall survival of GC patients (TCGA, http://www.cbioportal.org).

Several studies have demonstrated that ω-3 PUFA plays a direct anti-tumor or an adjuvant role in various cancers by modulating cell proliferation or apoptosis in the high micromolar range. We believe that, in our system, the effects of an ω-3 or ω-6 diet on xenograft growth were predominantly due to differences in vessel density rather than to a direct effect on cell proliferation. In fact, in the low micromolar range, which corresponds to the concentrations we found in vivo upon dietary modification, we found that PUFA suppressed the angiogenic response but did not directly affect GC cell proliferation in culture. In the present study, vessel density in shCTR xenografts did not differ among mice fed different diets, although cell proliferation was slightly, albeit significantly, reduced in mice fed an ω-3 or ω-6 diet. We ascribe this phenomenon to the very low in vivo angiogenic potential and to the massive presence of necrotic areas in these tumors. However, it should be noted that since PUFA administration induces the production of many intermediate metabolites that exert anti-angiogenic and anti-proliferative effects in vivo, we cannot exclude the possibility that in the xenograft microenvironment, these intermediates, together with PUFA, could reach a concentration sufficient to induce xenograft growth inhibition by directly modulating cell proliferation.

| Table 1. Lipidomic profile of mice fed ω-6 and/or ω-3 enriched diets. |
|------------------|------------------|------------------|
| **DIET** | **CTRL** | **ω-6** | **ω-3** |
| AA | 6353.6 ± 1444.1 | 16991 ± 925.8* | 5677.2 ± 1295 |
| PGD2 | 281 ± 43.9 | 2373 ± 33.3 | 327 ± 75.1 |
| PGE2 | 183.5 ± 45.1 | 172 ± 66.7 | 92.7 ± 20.1* |
| PGFla | 76 ± 33 | 57 ± 23 | 473 ± 14.7 |
| TXB2 | 740.8 ± 64.1 | 911.4 ± 147.1 | 466 ± 148.7 |
| LTB4 | 740.8 ± 141.2 | 775 ± 145.9 | 355.8 ± 109.8 |
| LXA4 | 5748.6 ± 258.5 | 50126 ± 185.1 | 5017 ± 230.5 |
| LXB4 | 134 ± 20.9 | 3187 ± 33.4* | 183.4 ± 21.5 |
| EPA | 1476 ± 280.6 | 21678 ± 671.3 | 5028.1 ± 462.8* |
| RvE1 | 108 ± 29.9 | 326 ± 17.1 | 130.8 ± 26.4 |
| RvE2 | 325 ± 532.2 | 3797 ± 474.2 | 88627 ± 576.1 |
| RvE3 | 213 ± 44.6 | 1623 ± 42.5 | 249.7 ± 73.6 |
| DHA | 8886 ± 1586.2 | 102202 ± 1364.6 | 17447 ± 572.1* |
| RvD1 | 458 ± 19.5 | 523 ± 47.5 | 957 ± 12.5* |
| RvD2 | 466 ± 14.6 | 475 ± 49 | 86 ± 29.4 |
| RvD3 | 2664.7 ± 365.9 | 3252 ± 446.5 | 5101.1 ± 380.3* |
| RvD5 | 30 ± 5.9 | 20 ± 8.2 | 46.7 ± 18.2 |
| RvD6 | 56.7 ± 15.8 | 40.7 ± 17.4 | 98.2 ± 25.6 |
| PDI | 56 ± 6.3 | 39.5 ± 6.3 | 56.3 ± 28 |

Lipid concentration in serum of mice fed the indicated diets (10 mice/group). Values are reported as nM.

*p < .05 vs. the CTRL diet group.
Pattern recognition receptors trigger inflammation, but can also promote its resolution. This is particularly important for the gastrointestinal tract, which, being continuously exposed to food- and microbiota-derived antigens, is subject to a chronic low-level of mucosal inflammation that must be counteracted to avoid tissue damage. In injured mouse colonic mucosa, FPR1 promotes tissue restitution by recognizing the endogenous N-terminal-derived annexin A1 (Ac2-26) peptides or specific commensal microbiota strains. Our data suggest that FPR1 is constitutively active in GC cells, which implies the presence of an endogenous ligand endowed with pro-resolving activity. A possible candidate ligand is peptide Ac2-26 that derives from AnxA1, which, as we previously showed, is constitutively expressed in GC cells.

In conclusion, we describe a novel molecular mechanism in GC, actively controlled by FPR1, that links PUFA metabolism and SPM biosynthesis with angiogenesis. This raises the possibility of new prognostic tools and therapeutic interventions for GC. It is feasible that FPR1 and/or components of pro-resolving pathways may represent novel risk factors or prognostic markers of GC. Administration of SPMs or pharmacologic stimulation of FPR1, ALOX15, or GPR32 could suppress GC angiogenesis and growth. Lastly, as SPM production can be largely determined by diet, increasing ω-3 or ω-6 consumption might be a tool in the management of GC.

Materials and methods

Cell culture

The AGS and MKN45 cell lines derived from poorly differentiated human gastric adenocarcinoma were grown as previously described. To generate AGS cells stably expressing ALOX5, ALOX15, or GPR32 shRNA, we used pools of five constructs (Qiagen, Valencia, CA, USA) containing 21-mer hairpin RNAs (shRNA) directed to various coding regions of each target gene. Transfectants were selected in medium with 500 ng/mL puromycin.

RNA isolation and real-time PCR

Total RNA was isolated and retrotranscribed according to the manufacturer’s instructions (Promega, Madison, WI, USA). Real-time quantitative PCR was performed on the CFX96 system (Bio-Rad, Hercules, CA, USA) using the PE SYBR Green PCR kit (Applied Biosystems, Grand Island, NY, USA). The target-specific primers used for real-time PCR are listed in Table S1. No-reverse transcribed mRNA samples served as a negative control. The expression levels of each target were calculated relative to that of control cells, arbitrarily considered equal to 1. Results were normalized to β-actin mRNA levels.

Flow cytometric analysis

Cells were incubated (30 min at 4°C) with specific or isotype control antibodies (Abs). ALOX5, ALOX15A, and ALOX15B Abs were from Santa Cruz Biotechnology (Dallas, TX, USA), anti-GPR32 was from Acris (Herford, Germany), anti-BLT1 from LSBio (Seattle, WA, USA), and anti-ChemR23 from Mybiosource (San Diego, CA, USA). Cells were analyzed with an FACS Calibur cytofluorimeter using CellQuest software (BD Biosciences, Mississauga, ON, Canada). When necessary, we performed cell membrane permeabilization using the Cytofix/Cytoperm kit (BD Biosciences).

Protein studies

Protein extractions, immunoblotting, and immunoprecipitation were carried out according to standard procedures. Anti-phospho-MAPK, -Akt, -p38, -JNK, -STAT3, -SRC antibodies (Abs) were from Cell Signaling Technology (Danvers, MA, USA). Anti-tubulin was from Sigma-Aldrich (St. Louis, MO, USA), and secondary anti-mouse and anti-rabbit Abs coupled to HRP were from Bio-Rad.

Xenografts in mice

Each group of 10 mice (4-week-old female CD1 nu/nu mice, Charles River, Wilmington, MA) was inoculated subcutaneously with shCTR, shFPR1, shALOX5, shALOX15, shGPR32 AGS cells (1×10⁶ cells/mouse). Tumor diameters were measured at regular intervals with a caliper. Tumor volumes (V) were calculated with the formula: V = A×B²/2 (A = axial diameter; B = rotational diameter). This study was conducted according to Italian regulations for experimentation on animals; the study protocol was approved by the Italian Ministry of Health. Paraffin-embedded tumors were analyzed by immunohistochemistry with anti-Ki-67 antibody from Biocare Medical (Concord, CA), anti-cleaved caspase 3, and anti-CD31 from R&D Systems (Minneapolis, MN).

Fatty composition of diets

The three isocaloric diets containing 5% fat (w/w) were designed as previously described. Pellets were prepared by Mucedola (Milan, Italy). We used grape seed and colza oils (50/50%) to prepare the pellets for the control diet (CTRL), grape seed oil (100%) for the ω-6 (ω-6), and colza oil/fish oils (80/20%) for an ω-3 fatty acid-enriched diet (ω-3). The oils were obtained from Sigma-Aldrich (St. Louis, MO). Pellets were stored under vacuum at −20°C. The diets were changed twice a week in each animal cage to avoid oxidative degradation of lipids.

ELISA and EIA assays

VEGF-A contents in culture supernatants were measured in duplicate determinations with a commercially available ELISA (R&D Systems). RvD1, LTB4, PGE2, and LXB4 contents in culture supernatants were measured in triplicate determinations with a commercially available EIA (Cayman Chemical, Ann Arbor, MI).

LC-MS/MS instrumentation and conditions

To extract the lipids, 120 μL samples (mouse serum or cell culture supernatants), together with deuterated internal standards,
were loaded onto solid phase extraction Waters Oasis HLB cartridges (60 mg sorbent, 30 μm particle size), and eluted with 2 mL methanol and 2 mL ethyl acetate into polypropylene tubes containing 6 μL of a glycerol solution (30% in methanol). The SPE eluates were dried and residues were then reconstituted in 100 μL methanol. 5 μL of methanol extract were analyzed by using a 4000QTrap mass spectrometer (Applied Biosystems) coupled to a 1,100 nanoHPLC system (Agilent Technologies, Waldbronn, Germany). The lipids were separated by using a micro-C18 column (10 cm x 1.0 mm, 5μ). The mobile phase was generated by mixing eluent A (0.1% acetic acid) and eluent B (acetonitrile/isopropanol 50/50) and the flow rate was 30 nL/min. Starting condition was 20% to 95% B in 5 min. Tandem mass spectrometry was performed using a turbo ion spray source operated in a negative mode (curtain gas 20psi, GS 1/2 50/50psi, ion spray voltage 60 V; Dwell 25 ms and temperature of 550°C), and the multiple reaction monitoring mode was used to detect a unique product ion arising from collision-induced fragmentation of the protonated parent compound as reported in Table S2.

**Statistical analysis**

Values from groups were compared by using the paired Student t test or the Duncan test. A p value < 0.05 was considered statistically significant.

**Disclosure of potential conflicts of interest**

The authors whose names are listed above inform the Editors-in-Chief, Deputy Editors, Senior Editors, or Scientific Editors that they have NO relationships that they believe could be construed as resulting in an actual, potential, or perceived conflict of interest with regard to this manuscript.

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