Probiotic *Lactobacillus rhamnosus* GG (LGG) restrains the angiogenic potential of colorectal carcinoma cells by activating a proresolving program via formyl peptide receptor 1

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Formyl peptide receptors (FPR1, FPR2 and FPR3) are innate immune sensors of pathogens and commensal bacteria and have a role in colonic mucosa homeostasis. We identified FPR1 as a tumour suppressor in gastric cancer cells due to its ability to sustain an inflammation resolution response with antiangiogenic potential. Here, we investigate whether FPR1 exerts similar functions in colorectal carcinoma (CRC) cells. Since it has been shown that the commensal bacterium *Lactobacillus rhamnosus* GG (LGG) can promote intestinal epithelial homeostasis through FPR1, we explored the possibility that it could induce proresolving and antiangiogenic effects in CRC cells. We demonstrated that pharmacologic inhibition or genetic deletion of FPR1 in CRC cells caused a reduction of...
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1. Introduction

Chronic inflammation is a risk factor for colorectal carcinoma (CRC) onset [1], as strongly suggested by the increased predisposition to colon carcinogenesis of inflammatory bowel disease (IBD) patients [2].

In recent years, unresolved chronic inflammation has been associated with insufficient production of mediators, which are actively involved in the resolution of inflammation [3,4]. Molecules diverse in nature are able to actively participate in different moments of the resolution program [3]: (i) lipidic autacoids known as specialized proresolving mediators (SPMs) [4]; (ii) proteic mediators [e.g. annexin A1 (AnxA1), adrenergic hormone, chemerin peptides, and galectin-1] [3,5]; (iii) gaseous mediators (nitric oxide, hydrogen sulfide and carbon monoxide) [3]; (iv) the adenosine, a purine nucleoside generated by the dephosphorylation of adenine [6]; and (v) neuropeptides such as acetylcholine and other neuropeptides produced by vagus nerves [7].

Specialized proresolving mediators are lipidic mediators derived from omega-6 (ω-6) arachidonic (AA), or ω-3 eicosapentaenoic (EPA) and docosahexaenoic (DHA) essential polyunsaturated fatty acids (PUFA) through the activity of lipooxygenases 5 and 15 (ALOX5/15). The best-characterized SPMs are lipoxins (LXA4, LXB4), E- and D-series resolvins (RvEs and RvDs), protectins (PD) and maresins (MaR) [8]. They exert anti-inflammatory, antiangiogenic and proresolving effects subsequent to inflammatory conditions [4,8,9].

We recently described a novel function of SPMs in gastric cancer (GC) demonstrating that RvD1 and LXB4 suppress angiogenesis, thus inhibiting tumour growth. We also demonstrated that formyl peptide receptor 1 (FPR1), a member of the FPR family, controls SPM production in GC [9,10], functioning as a tumour suppressor [11,12]. FPRs are pattern recognition receptors (PRRs) known to balance inflammatory and proresolving responses by sensing host-derived and bacterial products [13,14].

Several reports point to a crucial protective role of proresolving pathways also in CRC carcinogenesis [15]. Indeed, it has been demonstrated that CRC is associated with a reduced intake of ω-3 PUFA [16] and that dietary supplementation of ω-3 PUFA exerts anticancer effects in CRC [17]. Furthermore, ALOX15 has been described as a tumour suppressor in CRC [18] and specific SPMs (i.e. RvD1, LXA4) have been demonstrated to exert antitumour activity in CRC models [19–24].

Intestinal inflammatory conditions are strongly influenced and in turn affect microbiota composition [1,25]. In the last years, several studies in CRC patients and experimental models demonstrated that colon tumorigenesis is associated with significant alterations of intestinal microbial composition termed as dysbiosis [26,27]: in CRC patients, specific bacterial species are over-represented compared with those in noncancerous patients and exert protumorigenic function(s), while other species are under-represented and exert tumour suppressive functions [26–28].

Lactobacillus rhamnosus GG (LGG) is a commensal bacterium used as probiotic in order to reverse intestinal dysbiosis [29]. Several preclinical studies point to its effects in reducing chronic inflammation linked to CRC development [30]: LGG has been demonstrated...
to regulate homeostasis and restitution following colonic wounds in mice [31–34]; in CRC models, LGG activates proapoptotic and antimetastatic responses [35,36], lowers inflammation and favours adaptive protective immune responses against cancer cells [37]. It has been shown that LGG activity in intestinal epithelial cells depends on the expression of FPR1 [32].

Since the gastrointestinal tract is continuously exposed to external insults, proresolving pathways are particularly important to balance inflammatory responses for its homeostasis [4]. Thus, we investigated the role of FPR1 in CRC cells in order to verify the possibility that it functions as a regulator of inflammation resolution, angiogenesis and cancer. Moreover, we evaluated whether homeostatic and anticancer effects of LGG in CRC models could depend on its ability to activate a proresolving response. In particular, we investigated the possibility that LGG could activate a proresolving and an antiangiogenic response in CRC cells by stimulating FPR1.

Our data confirm this hypothesis and highlight the possibility that FPR1 could be exploited in order to increase the proresolving and inhibit the angiogenic potential of CRC cells, also through the use of probiotics.

2. Materials and methods

2.1. Cell culture

The HCT116 and HT29 cell lines derived from colorectal carcinoma (CRC) were kindly provided by S. Scala (Istituto Nazionale Tumori-IRCCS-Fondazione G. Pascale, Napoli, Italy) and grown as elsewhere described [38]. To generate HCT116 cells stably expressing FPR1 shRNA, we used pools of 5 constructs (Qiagen, Valencia, CA, USA) containing 21-mer short hairpin RNAs (shRNA) directed to various coding regions of the target gene. Transfectants were selected in medium with 500 ng mL⁻¹ puromycin [10]. Human umbilical vein endothelial cells (HUVECs) from Cell Biologics (Chicago, IL, USA) were grown in human endothelial cell medium with the addition of VEGF, heparin, EGF, FGF, hydrocortisone, L-glutamine, antibiotic/antimycotic solution and FBS according to the manufacturer’s instructions (Cell Biologics) [39].

Cell treatments to verify (a) mRNA changes, (b) enzyme or receptor protein expression, (c) AnxA1 expression, (d) signalling pathway activation and (e) autacoid release were made in serum-free media and after a 12-h serum starvation. Instead, cell treatment for VEGF-A release and to collect cell culture supernatants to be used in capillary formation assay was performed in 1% FBS to improve the stability of VEGF-A. The same conditions were used when bacterial supernatants (SN) were used to treat CRC cells; the relative control sample of each bacterial strain SN was the same titration of the culture broth.

Treatments of CRC cells were made with fMLF at 1 nM, the concentration at which it binds specifically to FPR1 [13]. The SPMs were used again at 1 nM, the same optimal concentration used to inhibit angiogenesis in the GC model [10].

The functional involvement of GPR32 or MAPK signalling in CRC proresolving responses was evaluated by preincubating cells for 30 min with an anti-GPR32 neutralizing antibody (Ab) (1 μg mL⁻¹) (GeneTex, Irvine, CA, USA) or the MAPK signalling inhibitor U0 126 (10 μM) (Cell Signaling, Danvers, MA, USA), respectively, before proceeding with the specific treatment.

2.2. Bacterial culture and supernatant preparation

The bacterial strains used in this study were as follows: Lactobacillus rhamnosus (Hansen) Collins et al. (LGG) (ATCC 53103), obtained from the ATCC (Manassas, VA); Escherichia coli ATCC 13762 (E. coli), used as control of nonprobiotic bacteria; and Bifidobacterium bifidum (B. bifidum), an anaerobic lactic acid bacterium isolated from the ProBiota Bifido (Seeking Health, Bellingham, WA, USA). Bacterial suspensions, at 0.1 optical density (OD) at 600 nm, were inoculated in broth medium and grown in slight motion at 37 °C overnight in aerobic or anaerobic condition, in order to obtain a number of colony-forming unit (cfu) of ~10⁶/mL, determined by plate counting on medium agar plates. In detail, LGG suspension was inoculated in De Man, Rogosa and Sharpe (MRS) broth and MRS agar medium (Becton Dickinson, Franklin Lakes, NJ, USA) in aerobic condition at 37 °C. E. coli ATCC 13762 was cultured in Tryptic Soy (TS) broth and TS agar (Oxoid, Basingstoke, Hampshire, UK) in aerobic condition at 37 °C. B. bifidum was cultured in anaerobic condition at 37 °C, using MRS broth medium supplemented with 0.25% cysteine/HCl (Sigma-Aldrich, St. Louis, MO, USA). Bacterial supernatant (SN) of each strain was prepared by centrifugation of the overnight cultures in the specific growth medium at 4000 g and 4 °C for 10 min and stored in single-use aliquots at −80 °C until needed.

2.3. Flow cytometry

Cells were incubated (30 min at 4 °C) with specific or isotype control Abs. ALOX5, ALOX15A and
ALOX15B Abs were from Santa Cruz Biotechnology (Dallas, TX, USA), anti-GPR32 from Acris (Herford, Germany), anti-BLT1 from LSBio (Seattle, WA, USA), and anti-ChemR23 and anti-FPR1 from R&D Systems (Minneapolis, MI, USA). Cells were analysed with a FACS Calibur cytometer using the CELLQUEST software (BD Biosciences, Lakes, NJ, USA). When necessary, we performed cell membrane permeabilization using the Cytofix/Cytoperm Kit (BD Biosciences). The receptors followed as indicators of resolution responses were the same modulated by FPR1 in the GC model [10]. The concentration used for flow cytometric staining was that indicated by manufacturers. AnxA1 staining was performed using a primary anti-AnxA1 goat polyclonal Ab (1:500) (R&D Systems) followed by the staining with a secondary anti-goat Ab Alexa Fluor 488 (Invitrogen, Waltham, MA, USA). The secondary antibody alone was used as a negative matched control.

2.4. ELISA and EIA
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, USA) [39]. Cell culture supernatants were assayed, undiluted for autacoid evaluations and diluted 1:5 for VEGF-A release.

2.5. Real-time PCR
Total RNA was isolated and retrotranscribed according to the manufacturer’s instructions (Qiagen) as previously described [40]. Real-time quantitative PCR was performed on iCycler (Bio-Rad, Hercules, CA, USA) using the PE SYBR Green PCR kit (Applied Biosystems, Foster City, CA, USA) as reported elsewhere [41]. Normalization was performed using β-actin mRNA levels. Primer sequences are listed in Table S1.

2.6. Protein studies
Protein extractions and immunoblotting experiments to evaluate signalling pathway activation were performed according to standard procedures [11]. Anti-phospho-MAPK, anti-phospho-Akt, and anti-phospho-STAT3 Abs (1:1000) were from Cell Signaling Technology. Antitubulin was from Sigma-Aldrich (1:10 000) (St. Louis, MO, USA), and secondary anti-mouse and antirabbit Abs coupled to HRP were from Bio-Rad (1:3000).

The expression of angiogenesis-related proteins in CRC cell culture supernatants was determined using the Human Angiogenesis Array Kit (R&D Systems) according to the manufacturer’s instructions. The array allows the detection of 55 angiogenesis-related proteins by specific capture antibodies spotted onto a nitrocellulose membrane. The data from developed X-ray film were digitalized and quantified using the ImageJ analysis software [42].

2.7. Tubule formation
The formation of network-like structures by HUVECs (Cell Biologics) on an extracellular matrix (ECM)-like 3D gel consisting of Matrigel® (BD Biosciences) was performed as previously described and validated [43]. HUVECs (5 x 10⁴) were seeded on a Matrigel matrix in the presence of cell culture supernatants. After incubation, HUVECs underwent differentiation into capillary-like tube structures. Tubule formation was defined as a structure exhibiting a length four times its width. Network formation was observed using an inverted-phase contrast microscope (Zeiss, Oberkochen, Germany). Representative fields were taken [43], and the number of branching points counted in five fields was presented as mean ± SD of three experiments.

2.8. Wound-healing assay
For wound-healing assays, confluent monolayers of HCT116 cells were treated with mitomycin (2 μg·mL⁻¹ for 2 h) (Sigma-Aldrich). The cell monolayers were scraped in three straight lines for each 60-mm dish with a p10 pipet tip. Cell migration was assessed as previously described [44]. Confluent monolayers were incubated for 30 min with an anti-GPR32 neutralizing Ab (1 μg·mL⁻¹) (GeneTex) or CsH (800 nM) (Sigma-Aldrich) and then treated with LGG SN or control broth (1:30 titration) for 12 h before assessing cell migration.

2.9. Statistical analysis
Values from groups were compared using the paired Student t test. P < 0.05 was considered statistically significant. Clinic–pathologic parameters in relation to FPR1 or FPR2 expression were plotted using the cBioPortal database. Coexpression data were obtained according to the cBioPortal online instructions: a log-rank test was provided to identify the significance of Spearman’s correlation coefficient between the mRNA expression z-scores (RNASeq V2 RSEM) [39].
3. Results

3.1. FPR1 pharmacologic stimulation sustains a proresolving response in colorectal carcinoma (CRC) cells

We recently described that FPR1 stimulation induces a proresolving program that relies on the expression of enzymes involved in SPM production (ALOX5, ALOX15A, and ALOX15B), the release of specific SPMs (RvD1 and LXB4), and the expression of SPM receptors (GPR32, ChemR23, and BLT1) in gastric cancer (GC) cells [11].

To determine whether FPR1 activates a proresolving program also in colon cancer, we selected two human colorectal carcinoma (CRC) cell lines, HT29 and HCT116 cells. The expression of FPR1 in HT29 and HT29 cells was assessed by FACS analysis (Fig. S1A). To pharmacologically modulate FPR1, we treated CRC cells with fMLF, an agonist to FPR1 [13], or with cyclosporine H (CsH), an inverse agonist to FPR1 [13], and we analysed the impact of these treatments on SPM pathway.

We found that, in HT29 and HCT116 cells, fMLF \((10^{-9} \text{ M} - 3\text{-h treatment})\) significantly increased, whereas CsH (800 nm – 3-h treatment) significantly decreased, the mRNA expression of enzymes (ALOX5, ALOX15A and ALOX15B) and receptors (GPR32, ChemR23 and BLT1) involved in SPM synthesis and recognition (Fig. 1A). In accordance with these observations, the protein levels of SPM enzymes (ALOX5, ALOX15A and ALOX15B) and receptors (GPR32, ChemR23 and BLT1) were significantly induced or reduced, compared with relative controls, in HT29 and HCT116 cells treated for 6 h with fMLF or CsH, respectively, as assessed by cytofluorimetric analysis (Fig. 1B,C).

The activation of SPM synthesis during resolution of inflammation counterbalances the production of proinflammatory lipidic mediators (e.g. prostaglandins and leukotrienes) [8,39]. In CRC, a crucial proinflammatory and protumorigenic role has been described to an anti-inflammatory function of LXA4 [23,46]. Thus, we decided to verify whether FPR1 pharmacologic modulation induces changes in PGE2 and LXA4 release, together with the two SPMs modulated by FPR1 in GC cells (i.e. RvD1 and LXB4) [11].

To this purpose, we treated CRC cells with fMLF \((10^{-9} \text{ M})\) or with CsH (800 nm) for 12 h and looked for RvD1, LXA4, LXB4, and PGE2 release. We found that fMLF significantly increased, while CsH significantly decreased SPM (RvD1, LXA4, and LXB4) release in both HT29 and HCT116 cells (Fig. 1D). Consistently, the treatment of HT29 and HCT116 cells with fMLF \((10^{-9} \text{ M} - 12\text{ h})\) significantly reduced, whereas CsH (800 nm – 12 h) significantly increased, PGE2 release (Fig. 1D).

These data indicate that FPR1 is able to activate in CRC cells an inflammation resolution program, by promoting the induction of SPMs (RvD1, LXA4, and LXB4), SPM enzymes (ALOX5, ALOX15A and ALOX15B) and SPM receptors (GPR32, ChemR23 and BLT1).

3.2. FPR1 pharmacologic modulation controls the angiogenic potential of CRC cells

We previously found that FPR1 ablation/pharmacological inhibition caused a drop in the endogenous levels of SPMs and a concomitant increase in the angiogenic potential of GC cells. We also found that SPMs control the production of angiogenic mediators in GC cells, since the exogenous administration of SPMs (RvD1 or LXB4) to FPR1-depleted GC cells could suppress their increased angiogenic potential [11].

To investigate whether FPR1 acts as a negative modulator of angiogenesis also in CRC cells, we treated HT29 and HCT116 cells with fMLF \((10^{-9} \text{ M})\) or with CsH (800 nm) for 3 h and measured the mRNA expression levels of several proangiogenic mediators (VEGF-A, VEGF-B, VEGF-C, VEGF-D, Ang1, and CXCL1). We observed that fMLF induced a reduction of mRNA levels for VEGF-A, VEGF-C and CXCL1 in HT29 cells and for VEGF-B, VEGF-C, Ang1 and CXCL1 in HCT116 cells (Fig. 2A). Consistently, CsH treatment significantly increased mRNA expression of proangiogenic molecules in both HT29 and HCT116 cells (Fig. 2A). Moreover, the release of VEGF-A was significantly lower in HCT116 treated for 12 h with fMLF \((10^{-9} \text{ M})\) and significantly higher in the same cells treated with CsH (800 nm) compared to relative controls (Fig. 2B). Similar results were obtained in HT29 cells (not shown).

To corroborate our results in CRC cultures, we verified the mRNA coexpression data present in the publicly available eBioPortal for Cancer Genomics database (http://www.ebiportal.org) [47,48]. Data on 594 colorectal adenocarcinoma revealed that FPR1 mRNA levels significantly and directly correlated with mRNA expression levels of the proresolving factors ALOX5 and ALOX15B (Fig. 2C). Consistently, FPR1 mRNA levels inversely correlated with mRNA levels of two key angiogenic mediators [VEGF-A and angiopoietin 1 (Ang)] (Fig. 2C).
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(A) mRNA Fold Change

(B) Cell Counts

(C) Mean Fluorescence Intensity

(D) RvD1 Change (%)

HCT116

HT29

Mean Fluorescence Intensity

 gel

Vehicle

Stimulus

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Fig. 1. Effects of formyl peptide receptor 1 (FPR1) pharmacologic modulation on specialized proresolving mediator (SPM) biosynthetic machinery of colorectal carcinoma (CRC) cells. (A) ALOX5, ALOX15A, ALOX15B, GPR32, ChemR23 and BLT1 mRNA fold change in HT29 and HCT116 cells treated with fMLF (10−9 M) or CsH (800 nM) for 3 h. Data are represented as mean ± SD of five independent experiments. *P < 0.05 compared with the control (dotted line) by Student’s t test. (B) ALOX5, ALOX15A, ALOX15B, BLT1 and ChemR23 protein expression levels (mean fluorescence intensity), assessed by cytofluorimetric analysis, in HT29 cells treated with fMLF (10−9 M) or CsH (800 nM) for 6 h. A representative experiment of three independent experiments is shown. (C) ALOX5, ALOX15A, ALOX15B, GPR32, ChemR23 and BLT1 protein expression levels (mean fluorescence intensity), assessed by cytofluorimetric analysis, in HCT116 cells treated with fMLF (10−9 M) or CsH (800 nM) for 6 h. Data are represented as mean ± SD of three independent experiments. *P < 0.05 compared with the control (NT) by Student’s t test. (D) Proresolving and proinflammatory autacoid (RvD1, LXA4, LXB4, PGE2) release in HCT116 and HT29 cells treated or not with fMLF (10−9 M) or CsH (800 nM) for 12 h. Baseline values of each mediator were in HCT116: RvD1 118 ± 18 pg/106 cells, LXB4 34 ± 5 pg/106 cells, LXA4 485 ± 58 pg/106 cells and PGE2 98 ± 12 pg/106 cells. Baseline values of each mediator were in HT29: RvD1 132 ± 24 pg/106 cells, LXB4 32 ± 4 pg/106 cells, LXA4 462 ± 48 pg/106 cells and PGE2 86 ± 11 pg/106 cells. Data are represented as mean ± SD of changes over baseline levels obtained in five independent experiments. *P < 0.05 compared with the control by Student’s t test.

Finally, in order to search for information on the clinic–pathologic role of FPRs and/or proresolving pathways in CRC, we queried the eBioPortal for Cancer Genomics database (http://www.cbioportal.org): mRNA levels of FPR1 showed a statistc trend of association with the Overall Survival Status (P = 0.11) and a direct and statistically significant association with the disease-free months (Fig. S2). No statistically significant association with the two parameters was found for FPR2, suggesting that FPR1 in CRC plays a nonredundant role similar to that observed in GC [11].

These data support the hypothesis that FPR1 is responsible for a proresolving and antiangiogenic response in CRC cells.

3.3. FPR1 genetic ablation reduces the proresolving activities and increases the angiogenic potential of CRC cells

To confirm the results presented in 3.1 and 3.2 paragraphs, we generated HCT116 cells stably transfected either with FPR1-targeting short hairpin RNAs (HCT116 shFPR1) or with nontargeting short hairpin RNAs (HCT116 shCTR). We identified various clones expressing low levels of the receptor (Fig. S1B).

By real-time PCR, we found that genetic ablation of FPR1 significantly reduced mRNA level of proresolving enzymes (ALOX15A and ALOX15B) and receptors (GPR32, ChemR23) compared with control cells (Fig. 3A). Moreover, HCT116 shFPR1 cells exhibited an increase of mRNA levels for angiogenic mediators compared with controls (Fig. 3A).

The genetic ablation of FPR1 significantly decreased the protein levels of ALOX5, ALOX15A, ALOX15B, GPR32 and ChemR23, but not BLT1 compared with HCT116 shCTR cells, as assessed by cytofluorimetric analysis (Fig. 3B). Moreover, the amount of RvD1, LXB4 and LXA4 released was significantly reduced in HCT116 shFPR1 compared with that in shCTR cells (Fig. 3C). Furthermore, FPR1 silencing in HCT116 cells caused the release of higher levels of PGE2 compared with that of controls (Fig. 3C). In accordance, these cells displayed increased constitutive release of VEGF-A (Fig. 3D).

The formation of capillary-like tube structures in the extracellular matrix by endothelial cells (ECs) is a classical method to measure angiogenesis in vitro [49]. To investigate whether differences in FPR1 expression/activation control functional angiogenic properties of CRC cells, we studied the ability of CRC cell conditioned media (CM) to induce human umbilical vein endothelial cell (HUVEC) network formation on a Matrigel substrate. In particular, we evaluated tubule formation in vitro in response to CM from HCT116 cells silenced or not for FPR1 (shCTR vs shFPR1). As shown in Fig. 3E, HUVECs plated in the presence of CM from HCT116 shCTR cells formed only a few tube structures at 12 h. On the contrary, when the endothelial cells were plated in the presence of CM from HCT116 shFPR1, a significantly higher number of formed tube structures were observed compared with that induced by shCTR CM (Fig. 3E).

To assess whether the increased proangiogenic potential of FPR1-silenced HCT116 cells could be due to the defective SPM biosynthesis of these cells, we added back LXA4 (1 nm), RvD1 (1 nm) or LXB4 (1 nm) to HCT116 shFPR1 and shCTR cells for 3 h and evaluated their proangiogenic activity. By means of real-time PCR, we observed that LXA4 had no effects on proangiogenic mediator expression (Fig. S3). Thus, although it has been described that LXA4 exerts a strong anti-inflammatory potential in CRC [23,46], our results demonstrate that it has no effect on the modulation of CRC cell angiogenic potential.
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(A) mRNA Fold Change

(B) VEGF-A release (pg/10^6 cells)

(C) mRNA Expression, RSEM (Batch normalized from Illumina HiSeq RNASeqV2): FPR1 (19q13.41)

Spearman: 0.40 (p = 7.76e-22)
Pearson: 0.29 (p = 2.95e-11)

Spearman: 0.55 (p = 2.83e-43)
Pearson: 0.13 (p = 3.85e-3)

Spearman: -0.19 (p = 7.055e-6)
Pearson: -0.06 (p = 0.168)

Spearman: -0.04 (p = 0.0585)
Pearson: 0.08 (p = 0.3851e-3)
Fig. 2. Effects of formyl peptide receptor 1 (FPR1) pharmacologic modulation in colorectal carcinoma (CRC) cells on angiogenic response. (A) VEGF-A, VEGF-B, VEGF-C, VEGF-D, Ang1 and CXCL1 mRNA fold change in HT29 and HCT116 cells treated with fMLF (10^{-9} M) or CsH (800 nM) for 3 h. Data are represented as mean ± SD of five independent experiments. *P < 0.05 compared with the control (dotted line) by Student’s t test. (B) VEGF-A release in HCT116 cells treated with fMLF (10^{-9} M) or CsH (800 nM) or the relative controls for 12 h. Data are represented as mean ± SD of five independent experiments. *P < 0.05 compared with the control by Student’s t test. (C) Correlation between the mRNA expression levels of the indicated markers in 594 patients affected by colorectal adenocarcinoma. Spearman’s factor, Pearson’s factor, and the relative p are indicated.

RvD1 and LXBa were able to reduce the expression of proangiogenic mRNAs, in HCT116 shFPR1 and, to a lesser extent, in HCT116 shCTR cells. Consistently, as shown in Fig. 3D, we found that RvD1 and LXBa significantly reduced VEGF-A protein release in both HCT116 shFPR1 and, to a lesser extent, HCT116 shCTR cells.

Finally, RvD1 and LXBa displayed the ability to restore the expression of ALOXs and SPM receptor mRNAs in HCT116 shFPR1 cells (Fig. S4A). The RvD1- and LXBa-induced upregulation of SPM enzymes and receptors was also confirmed at the protein level by FACS analysis (Fig. S4C, D).

These results demonstrate that FPR1 exerts an antiangiogenic effect in CRC cells through the modulation of SPM production.

3.4. **Lactobacillus rhamnosus GG (LGG) supernatant sustains resolvin and antiangiogenic responses in CRC cells**

Previous studies showed that *Lactobacillus rhamnosus* GG (LGG), a colonic commensal bacterium and one of the most used probiotic strains [30], exerts a homeostatic function in intestinal mucosa [31–33], decreases levels of procarcinogenic metabolites, reduces chronic inflammation associated with intestinal neoplastic transformation and inhibits the proliferation of malignant cells [30,35,36]. Some of these LGG functions were associated with its ability to interact with FPR1 [32]. Thus, we hypothesized that LGG protective effect in normal intestinal mucosa and LGG anticancer activities in CRC cells could be linked to a prosolving and antiangiogenic response activated by the bacteria.

In order to verify our hypothesis, we treated HT29 and HCT116 cells with LGG supernatant (SN) (1 : 30 titration) for 3 h, or with the same dilution of the culture broth, and evaluated, by real-time PCR, the expression levels of prosolving pathways’ components and of proangiogenic markers. The 1 : 30 titration was chosen as optimal one after testing different dilutions (1 : 10–1 : 100) of LGG SN in cell culture media in order to balance pH changes induced by LGG SN and its functional activity (not shown). We found that LGG SN induced, in both CRC cells, a significant increase of ALOX15A, ALOX15B, GPR32, ChemR23 and BLT1 mRNA levels (Fig. S5A) and a statistically significant decrease of proangiogenic mediator mRNA levels (VEGF-C, VEGF-D, Ang1 and CXCL1) (Fig. S5B) when compared to control (culture broth).

We confirmed these observations at the protein level: the treatment of HT29 cells with LGG SN for 6 h significantly increased ALOX5, ALOX15A, ALOX15B, BLT1 and ChemR23 protein levels (Fig. 4A). Similar results were obtained in HCT116 cells (not shown). Consistently with the induction of enzymes, LGG SN (12 h treatment) induced a significant increase in SPM release (RvD1, LXBa, LXA4) and a significant decrease in PGE2 levels in both HT29 and HCT116 cells (Fig. 4B).

To confirm the antiangiogenic effect of LGG in CRC cells, we evaluated, by an ELISA assay, VEGF-A release in HCT116 and HT29 cells treated for 12 h with LGG SN or broth as a control. We found that LGG SN was able to significantly reduce VEGF-A release in both CRC cells compared with the control broth (Fig. 4C). Furthermore, we evaluated the expression of several other angiogenic proteins using dedicated antibody arrays incubated with CM from HT29 or HCT116 cells treated for 12 h with LGG SN or the relative control (broth) [39]. LGG SN (1 : 30 titration) treatment in HT29 cells downregulated, with changes superior to 10%, the levels of amphiregulin, artemin, coagulation factor III, DPP IV, EGF-VEGF, endothelin 1, GDNF, IGFBP-1, IGFBP-2, TGF-beta1, MCP-1, MMP8, MMP9, pentraxin 3, PD-ECGF, PDGF-AA, PDGF-AB and BB, PiGF, Serpin E1, TIMP-1, TIMP-4, Thrombospondin 1, uPA and VEGF (Fig. 4D).

We then evaluated the ability of LGG SN to modulate CRC cell functional angiogenic potential. HUVECs plated on wells coated with Matrigel with the addition of CM from HCT116 or HT29 cells treated with control culture broth (1 : 30 titration) formed a characteristic capillary-like network at 12 h. On the contrary, when the cells were plated on Matrigel with the addition of CM from HCT116 or HT29 treated...
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(A) HCT116 shFPR1

Gene Expression

(B) HCT116

Mean Fluorescence Intensity

(C) HCT116

Proinflammatoryarker Release

(D) HCT116

VEGF-A Release

(E) HCT116

Branching points / 5 fields

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with LGG SN (1 : 30 titration), a significantly lower number of tube structures were observed (Fig. 4E).

Finally, to verify whether the antiangiogenic potential of LGG SN depends on SPM activity, we used a GPR32 neutralizing antibody (Ab) that inhibits the activity of its ligand RvD1. For this purpose, HCT116 cells were stimulated with LGG SN in the presence or absence of the anti-GPR32 (1 μg·mL⁻¹) Ab, and their antiangiogenic potential was evaluated. As shown in Fig. 6, at the concentration used for this experiment, the neutralizing anti-GPR32 Ab was able to significantly reduce the LGG-mediated inhibition of the angiogenic potential of CRC cells. The anti-GP32 effect was partial, as expected, due to its ability to block only RvD1 effects and not that of other SPMs (Fig. 6).

These data suggest that LGG is able to induce a proresolving response and a following antiangiogenic effect in CRC cells.

### 3.5. The proresolving and antiangiogenic properties of LGG are not common to other commensal bacteria

In order to verify the specificity of action of LGG, we asked whether *Escherichia coli* (*E. coli*), as example of a commensal nonprobiotic strain [50], or *Bifidobacterium bifidum* (*B. bifidum*), as other lactic acid probiotic strain [51], could sustain a proresolving and antiangiogenic response in CRC cells.

To this aim, we treated HT29 and HCT116 cells with *E. coli* or *B. bifidum* SN (1 : 30 titration) and verified their effects on ALOX expression, RvD1 and VEGF-A release. By flow cytometric analysis, we observed that neither *E. coli* nor *B. bifidum* SN were able to increase the level of ALOX5, ALOX15A and ALOX15B proteins in HCT116 cells, while LGG did it (Fig. 5A). Similar results were obtained also in HT29 cells (not shown). Consistently, LGG SN significantly induced RvD1 release, while *E. coli* and *B. bifidum* SN did not (Fig. 5B) both in HT29 and in HCT116 CRC cells. Finally, we verified the effects of *E. coli* or *B. bifidum* SN on VEGF-A release. Neither *E. coli* nor *B. bifidum* SN were able to reduce VEGF-A release from HT29 and HCT116 cells (Fig. 5C).

Altogether, these data support the evidence that the activation of a proresolving and antiangiogenic response in CRC cells is not general and common to all the commensal or to all the lactic acid bacteria.

### 3.6. *Lactobacillus rhamnosus* GG (LGG)-mediated proresolving response requires FPR1

It has been reported that LGG depends on the expression of FPR1 for its activity in colon cells [32,52]. To provide evidence that LGG-mediated proresolving and antiangiogenic responses in CRC cells require FPR1, we treated HCT116 shFPR1 and shCTR with LGG SN or with the control broth and we evaluated the production of proresolving and proangiogenic factors. LGG SN (1 : 30 titration—3 h) promoted a significant increase of ALOX5, ALOX15A and ALOX15B mRNA levels in shCTR but not in shFPR1 CRC cells (Fig. 6A). Similar results were obtained for the receptors GPR32, ChemR23 and BLT1 (Fig. 6A). Consistently, LGG SN induces the reduction of proangiogenic mediators (VEGF-A, -B, -C) only in shCTR but not in shFPR1 CRC cells (Fig. 6A).

To determine whether LGG depends on FPR1 also for its induction of SPM release, we treated shCTR and shFPR1 HCT116 cells with LGG SN for 12 h. We found that HCT116 shFPR1 treated with LGG SN did not display the increase in RvD1, LXA₄ and LXB₄ release observed in control cells (Fig. 6B). As
expected, LGG SN caused a significant reduction in PGE2 release in shCTR but not in shFPR1 HCT116 cells (Fig. 6B).

To confirm that also the antiangiogenic effect of LGG in CRC cells depends on FPR1, we evaluated the VEGF-A release in HCT116 shFPR1 and shCTR cells in the presence or absence of LGG SN after 12 h. As shown in Fig. 6C, VEGF-A release was higher in HCT116 shFPR1 than in controls; LGG SN treatment was able to significantly reduce VEGF-A release in shCTR but not in shFPR1 HCT116 cells (Fig. 6C).
Taken together, these data demonstrate that the proresolving and antiangiogenic activities of LGG require FPR1.

It has been described that one of the most important homeostatic functions of LGG is mediated by its ability to sustain restitution of injured intestinal epithelial monolayers by interacting with FPR1 [31,52]. Here, we verified whether the ability of LGG to induce CRC epithelial cell restitution through the activation of FPR1 is dependent, at least in part, on the activation of proresolving pathways. To this aim, we performed a wound-healing assay on HCT116 in the presence or absence of LGG SN, CsH (800 nM) or a neutralizing anti-GPR32 Ab (1 μg/mL). As shown in Fig. 6D, LGG SN elicits a significant movement of HCT116 cells following injury, which was completely abolished by CsH, confirming the dependence of LGG effects on FPR1 (Fig. 6D).

Interestingly, the neutralizing antibody against GPR2 was able to significantly reduce (about 30%) the LGG SN-induced migration, suggesting that the effects of LGG SN on FPR1 in terms of wound closure imply, at least in part, the production of proresolving mediators (i.e. RvD1).

In other experimental models, the activation of a proresolving response goes through the induction, not only of lipidic SPMs but also of proresolving mediators of a different type (e.g. AnxA1) [53]. In order to verify the possibility that LGG and FPR1 could induce AnxA1 expression in CRC cells, we treated both HT29 and HCT116 cells with LGG SN (1 : 30 titration), fMLF (10^{-9} M) or the three SPMs [RvD1 (1 nM), LXB4 (1 nM) and LXA4 (1 nM)] for 12 h and verified the expression levels of AnxA1, a potent endogenous proresolving and immunomodulatory protein [5].

FACS analysis shows that both fMLF and LGG SN induced an increase in the protein expression of AnxA1 in CRC cells (Fig. S7). Similarly, the treatment with RvD1, LXB4 and LXA4 increased AnxA1 protein expression levels in CRC cells (Fig. S7). These observations suggest that FPR1 activation sustains a proresolving response that includes AnxA1 and that SPMs could further stimulate AnxA1 expression in a feed-forward loop [53].

These results confirm previous observations indicating that LGG mediates a homeostatic function in colonic mucosa requiring FPR1 [31,32]. Moreover, our data indicate that this FPR1 function is maintained also in CRC cells, and is dependent on its ability to sustain a proresolving response.

3.7. FPR1 activation of proresolving program requires MAPK activation

We then explored the signalling pathways, which are involved in the proresolving response of CRC cells to FPR1 activation mediated by LGG or fMLF.

To this aim, we treated HCT116 cells with fMLF (10^{-9} M) or LGG SN (1 : 30 titration) and the relative controls (not-treated or broth alone, respectively) and harvested at different time points. fMLF has been demonstrated to classically activate the ERK, PI3K/Akt and STAT3 pathways [54,55]; thus, we verified the expression levels of phospho-MAPK, phospho-Akt, and phospho-STAT3 in order to evaluate the activated forms of these proteins.

As shown in Fig. 7A, in HCT116 cells a significant activation of MAPK and STAT3 was observed in response to fMLF, while no significant activation levels of Akt were detected. LGG SN, when compared to the broth alone at the same titration (1 : 30),
activated MAPK but not STAT3 signalling (Fig. 7A).

Thus, the common proresolving and antiangiogenic properties of fMLF and LGG SN could be due to MAPK activation.

To verify this hypothesis, we treated HCT116 cells with fMLF (10^{-6} m) or LGG SN (1 : 30 titration) and the relative controls (not-treated or broth alone, respectively) in the presence or absence of U0 126, a selective inhibitor of mitogen-activated protein kinase kinase [56]. Figure 7B shows that fMLF and LGG SN significantly upregulated ALOX15A, ALOX15B and ChemR23 expression levels and that the preincubation of cells with U0 126 partially or completely reverted these effects. Consistently, VEGF-B and VEGF-C were significantly reduced by fMLF and LGG SN treatments and these effects were not detectable in cells pre-treated with U0 126 (Fig. 7C).

These experiments demonstrated that the proresolving and antiangiogenic responses of fMLF and LGG require MAPK signalling activation.
4. Discussion

The homeostasis of intestinal mucosa is tightly regulated by mechanisms able to perceive bacterial species distinguishing between pathogenic and commensal ones, triggering an inflammatory antibacterial and a tolerogenic proresolving response, respectively [1, 57]. Formyl peptide receptors (FPRs), a family of pattern recognition receptors, can recognize several bacterial products and trigger either inflammation or its resolution [13, 14], being optimal candidates to the role of central regulators of intestinal mucosa homeostasis [14, 32, 58].

It has been reported that intestinal epithelial cells sense commensal bacteria using various receptors, including FPRs; as a consequence of this, an increase in barrier function, and improved resolution of epithelial wounds are observed [31, 52, 58, 59]. For instance, it has been demonstrated that the commensal bacteria Lactobacillus rhamnosus GG (LGG), by activating FPR1, influence intestinal epithelial homeostatic signalling and sustain epithelial cell motility enhancing wound restitution [31, 52]. Furthermore, several studies point to an important role of intestinal microbiota, not only in the physiology of intestinal mucosa but also in eliciting a protective antitumour response by both acting directly on cancer cells and modulating the immune response to them [26, 28, 30, 60].

We recently showed that the genetic ablation of FPR1 caused an increase in proinflammatory, angiogenic and tumorigenic potential in gastric cancer (GC) cells [10, 11]. We further showed that these functions of FPR1 are mediated by its ability to sustain the expression and function of proresolving pathways [11]. We also demonstrated that Toll-like receptor 7 (TLR7) displays similar proresolving and antiangiogenic properties in non-small-cell lung cancer cells [39]. Here, we demonstrated, for the first time, that FPR1 exerts a similar function also in the intestinal mucosa. Taken together, our results sustain the hypothesis that specific PRRs could exert homeostatic and proresolving functions in different tissues, which need to balance injuries and inflammatory insults.

More in detail, we showed that FPR1 activation mediated by formyl peptides, which are the main natural ligands to FPR1 [13], or by LGG SN, induces the expression of enzymes and receptors involved in proresolving responses, and the release of a significantly higher amount of SPMs (RvD1, LXB4, LXA4) at the expense of proinflammatory lipid mediators (PGE2). Although we focused our attention specifically on lipidic proresolving autacoids, the proresolving response in CRC cells mediated by FPR1 could be more general, since we also verified that fMLF and LGG are able to induce the expression in CRC cells of AnxA1, a proteic proresolving mediator [53].

The physiology of this type of response is of paramount importance, if one takes into account that SPMs act in vitro at an optimal concentration in the low nanomolar range. Indeed, we observed that the antiangiogenic properties of SPMs are detectable at 0.5–1 nm concentration. By means of EIA, we detected basal concentration of RvD1 released by 10^6 of two distinct CRC cells around 300 pm; LXB4 was constitutionally released at ~ 80 pm, while LXA4 was around 1.5 nm. The stimulation of CRC cells with LGG increased RvD1 release of around 40%, LXB4 of 20% and LXA4 of 30%, thus allowing to reach in cell culture media, and presumably in the gut, optimal and active concentrations of each SPM. Furthermore, it should be considered that while we identified 1 nm as the optimal concentration of each SPM alone to inhibit angiogenesis, in the gut the interaction of LGG product(s) with intestinal epithelial cells allows the contemporary production of several SPMs, which could act in synergy.

Although we have still not identified the bacterial products secreted in the LGG SN and responsible for FPR1 stimulation, it is likely that LGG-derived formylated peptides are the mediators of the observed effects. However, we cannot exclude the presence in LGG SN of other FPR1 agonist(s). Whatever the case, the activity of secreted FPR1 ligands in LGG SN suggests that, in the physiology of the gut, LGG could exert its homeostatic/protective effects on intestinal mucosa by releasing several factors in the extracellular space.

Consistently with previous observations demonstrating that LGG sustains intestinal epithelial restitution via an FPR1-mediated ERK activation [31, 52], we observed that LGG SN and fMLF shared the ability to activate MAPK signalling in CRC cells. We also verified that the proresolving response activated by LGG in CRC cells is dependent on the activation of this signalling.

In the GC model, the levels of SPMs inversely correlated with the number of proangiogenic mediators produced by cancer cells [11]. Consistently, also in the colorectal carcinoma (CRC) model, SPM production inversely correlated with the angiogenic potential of CRC cells. In particular, we observed that FPR1 activation mediated by fMLF or LGG SN significantly reduced the production of several angiogenic mediators in CRC cells. Furthermore, our experiments also demonstrated that the increased angiogenic potential of CRC cells lacking FPR1 is due to the deficit of...
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(A) mRNA Fold Change

- ALOX5
- ALOX15A
- ALOX15B
- GPR32
- ChemR23
- BLT1
- VEGF-A
- VEGF-B
- VEGF-C
- VEGF-D
- Ang1
- CXCL1

(B) LGG SN-induced ALOX5 Change (%)

(C) LGG SN-induced VEGF-A release (pg/10^6 cells)

(D) LGG SN-induced PGE2 Change (%)

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The different effects observed between RvD1/LXB4 and LXA4 suggest that probably in dependence of the mediators and on the tissue district, each SPM could exert a typical or at least predominant activity among that ascribed to SPMs (e.g. control of inflammation, limiting tissue damage, promoting resolution, attenuating fibrosis and inhibiting angiogenesis) [61].

In a mouse hepatocarcinoma cell line [62] and in a model of inflammation-induced pathological neovascularization of the cornea [63], LXA4 has been reported as antiangiogenic. However, in the colon experimental model, it has been described to date that LXA4 could both protect against acute injury [46] and suppress CRC development [23] by specifically regulating intestinal mucosa inflammation: LXA4 inhibits inflammatory mediator expressions [23,46], and reduced proinflammatory monocyte and neutrophil infiltration in tumours favouring lymphocyte activation [23,46].

Obviously, these findings deserve a more in-depth study in order to identify the possible different mechanisms of action (e.g. signalling pathways, receptor expression levels, cell metabolic asset) justifying the antiangiogenic properties of LXA4 in other tumours and not in CRC model and the differences in action compared with RvD1 and LXB4.

We demonstrated that the proresolving and antiangiogenic program in CRC cells could be induced by supernatants obtained by LGG cultures. Our data reinforce the idea that probiotic species contribute to an enhanced repair of mucosal wounds [32,33] and to a protective antitumour response, which imply not only the already demonstrated antiproliferative effect [35] but also, as here presented, an antiangiogenic response on cancer cells.

These speculations are corroborated by the evidence that the properties observed for LGG are not shared with other commensal nonprobiotic bacteria, as demonstrated by our experiments on CRC cells treated with *E. coli* and by other evidence in the literature [32,33,64]. Furthermore, we did not observe the same effects of LGG on CRC cells neither when we used a different lactic acid probiotic strain, as *B. bifidum*. We are aware that we did not identify a specific factor/protein produced by LGG and activating FPR1; however, our experiments with *E. coli* and *B. bifidum* and previous evidence suggest that the proresolving and homeostatic functions can be ascribed only to some bacterial strains [64].

Several experimental observations in the literature suggest that SPMs or diet supplement of their precursors (ω3/6 PUFA) could integrate CRC treatment because of their ability to counteract intestinal carcinogenesis [15]. In addition, specific commensal bacteria have been identified and described as able to limit colon tumorigenesis by acting on cancer cells or on the protumorigenic inflammatory microenvironment [65]. However, to date no connection between commensal microbiota, a pattern recognition receptor sensing specific probiotics and the activation of proresolving pathway has been made. We provide direct evidence of such concept, by showing that LGG can activate FPR1 sustaining the expression and function of proresolving pathways, which, in turn, suppress angiogenesis.

**Fig. 6.** Dependence of *Lactobacillus rhamnosus* GG (LGG) supernatant (SN) effects on formyl peptide receptor 1 (FPR1) expression in colorectal carcinoma (CRC) cells. (A) ALOX5, ALOX15A, ALOX15B, GPR32, ChemR23, BLT1, VEGF-A, VEGF-B, VEGF-C, VEGF-D, Ang1 and CXCL1 mRNA fold change in HCT116 cells silenced for FPR1 (HCT116 shFPR1, three clones) or in control cells transfected with nontargeting short hairpin RNAs (shCTR cells, a mass population) upon treatment for 3 h with *Lactobacillus rhamnosus* GG (LGG) supernatant (SN)—1 : 30 titration. Data are represented as mean ± SD of five independent experiments. *P < 0.05 compared with the control (broth—dotted line) by Student’s *t* test. 6§<0.05 compared with the relative control by Student’s *t* test. (B) Proresolving and proinflammatory autacoid (RvD1, LXA4, LXB4, PGE2) release over control in HCT116 shCTR (a mass population) or shFPR1 (three clones) upon treatment for 12 h with LGG SN—1 : 30 titration or the culture broth for 12 h. Data are represented as mean ± SD of changes over baseline levels obtained in five independent experiments. *P < 0.05 compared with the broth control by Student’s *t* test. 6§ <0.05 compared with the relative control by Student’s *t* test. (C) VEGF-A release in HCT116 shCTR (a mass population) or shFPR1 (three clones) cells treated with LGG SN—1 : 30 titration or the culture broth for 12 h. Data are represented as mean ± SD of five independent experiments. *P < 0.05 compared with shCTR broth by Student’s *t* test. (D) Wound-healing assay of HCT116 cells in the presence of LGG SN (1 : 30 titration) or the same dilution of culture broth for 12 h. Cells were pretreated or not for 30 min with CsH (800 nM) or a neutralizing anti-GPR32 antibody (1 μg·mL⁻¹). Representative photograms and a quantitative evaluation of the wound closure are shown. Scale bar 50 μm. Values represent the average of triplicate experiments ± SD. *P < 0.05 compared with broth alone by Student’s *t* test. §P < 0.05 compared with the relative control by Student’s *t* test.
5. Conclusions

Our results consolidate the hypothesis that a correlation exists linking proresolving pathways’ deficit and cancer in humans. Our data also highlight the possibility that innate immune receptors, including FPR1, could be the key regulators of the balance between microbiota recognition, inflammation regulation and neoplastic transformation. By defining the molecular mechanisms linking lipid metabolism and inflammation resolution with FPR1 in gastrointestinal (GI) tract, our data will allow the comprehension of the
general mechanisms involved in tumour cell growth following their angiogenic switch and will open the possibility to identify new prognostic markers and a novel therapeutic approach for cancers of the GI tract.

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Conflict of interest
The authors declare no conflict of interest.

Author contributions
FL participated in the design of the study, carried out experiments, drafted the manuscript, and read and approved this manuscript. MM, DS, CP, VC, GM, ES and PS carried out experiments, and read and approved this manuscript. RMM conceptualized and designed the study, analysed and interpreted the data, wrote the manuscript and designed the study, carried out experiments, analysed and interpreted the data, wrote the manuscript and gave final approval of the manuscript.

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Data accessibility
All data generated or analysed during this study are included within the article or available from the corresponding author on reasonable request.

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Supporting information

Additional supporting information may be found online in the Supporting Information section at the end of the article.

Fig. S1. Formyl peptide receptor 1 (FPR1) expression levels in colorectal carcinoma (CRC) cells.

Fig. S2. Formyl peptide receptor 1 (FPR1) correlation to colorectal carcinoma (CRC) patients’ characteristics.

Fig. S3. Absence of antiangiogenic effect of LXA4 in colorectal carcinoma (CRC) cells.

Fig. S4. Antiangiogenic and proresolving effects of RVd1 and LXB4 in colorectal carcinoma (CRC) cells.
Fig. S5. Effects of *Lactobacillus rhamnosus* GG (LGG) supernatants LGG (SN) on specialized proresolving mediator (SPM) biosynthetic machinery and angiogenic potential of colorectal carcinoma (CRC) cells.

Fig. S6. Antiangiogenic effects of *Lactobacillus rhamnosus* GG (LGG) supernatants (SN) in the presence of a neutralizing anti-GPR32 antibody.

Fig. S7. Annexin A1 (AnxA1) induction upon formyl peptide receptor 1 (FPR1) activation or specialized proresolving mediator (SPM) stimulation of colorectal carcinoma (CRC) cells.

Table S1. List of primers.

Data S1. Supplementary legends.