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Myofibroblast-Derived SFRP1 as Potential Inhibitor of Colorectal Carcinoma Field Effect

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Abstract

Epigenetic changes of stromal-epithelial interactions are of key importance in the regulation of colorectal carcinoma (CRC) cells and morphologically normal, but genetically and epigenetically altered epithelium in normal adjacent tumor (NAT) areas. Here we demonstrated retained protein expression of well-known Wnt inhibitor, secreted frizzled-related protein 1 (SFRP1) in stromal myofibroblasts and decreasing epithelial expression from NAT tissues towards the tumor. SFRP1 was unmethylated in laser microdissected myofibroblasts and partially hypermethylated in epithelial cells in these areas. In contrast, we found epigenetically silenced myofibroblast-derived SFRP1 in CRC stroma. Our results suggest that the myofibroblast-derived SFRP1 protein might be a paracrine inhibitor of epithelial proliferation in NAT areas and loss of this signal may support tumor proliferation in CRC.

Introduction

Communication between the epithelial and stromal compartments is fundamental in the modulation of normal epithelial homeostasis which becomes dysregulated during carcinogenesis due to genetic and epigenetic alterations [1]. Regarding these compartments myofibroblasts (α-SMA+, fibroblast-like cells) represent ultimate members of the information flow [2]. Myofibroblasts become primarily mesenchymal elements during the development of colorectal cancer (CRC) and may play a crucial role in the process of field cancerization [3–5]. The theory of field cancerization describes the formation of a genetically and epigenetically altered, but histologically normal tumor area [6–8]. These genetic and epigenetic changes could contribute to the altered epithelial homeostasis, characterized by increased cell proliferation and predispose to the development of cancer in morphologically normal adjacent tumor (NAT) areas [9]. In some cases, between tumor and NAT areas, a transitional area (TA) was identified, which displayed a different degree of dysplasia (i.e. altered crypt morphology, elongation, pseudostratification, loss of cell polarity and nuclear polymorphism). Although several studies have already described molecular abnormalities in association with field cancerization in epithelial tumors including CRC, the exact role of stroma in this process is still unclear [3,10].

Here, we aim to examine the potential role of stroma-derived Wnt inhibitor secreted frizzled-related protein 1 (SFRP1) in CRC field cancerization.

SFRP1 inhibits proliferation and induces apoptosis by directly binding to Wnt-1 and Wnt-5 ligands via preventing the activation of Wnt receptors (i.e. Frizzled receptors (FRZ) and low-density lipoprotein receptor-related protein-5 and 6 (LRP-5, LRP-6)) [11]. This system is dysregulated in around 90% of sporadic CRC patients due to aberrant canonical Wnt signaling, including mutation of cytoplasmic β-catenin degradation complex (DC) proteins, such as Adenomatous Polyposis Coli (APC) and Axin [12–14]. In 5% of CRC cases, β-catenin is mutated and does not undergo proteosomal degradation via failed phosphorylation by GSK3β (member of DC) [15]. Mutation of the Wnt pathway results in inappropriate nuclear β-catenin migration, accumulation and T-cell factor (TCF)/lymphocyte enhanced factor (LEF) activation [11,13]. This irregular TCF/LEF activation is independent of Wnt receptor activation; however, changes in the homeostasis of cell lines bearing an APC mutation as a result of the effect of different Wnt inhibitors have been described [12,16,17]. Based on the methylation analysis of macrodissected samples, it has been described that in colorectal carcinogenesis SFRP1 promoter is epigenetically silenced [12,18–20].
In this study, we aim to examine the protein expression and methylation patterns of myofibroblast-derived SFRP1 in NAT and CRC tissues, and to demonstrate the effect of SFRP1 protein on HCT116 CRC cell line as a potential model of paracrine (stromal) inhibition of the Wnt pathway in colorectal carcinoma.

**Materials and Methods**

**Ethics statement**

The study was conducted according to the declaration of Helsinki and approved by Semmelweis University Ethics Committee and the governmental Regional and Institutional Committee of Science and Research Ethics (TUKÉB, Nr69/2008). Written informed consent was obtained from all patients included in the study.

**mRNA microarray analysis of biopsy and laser microdissected stroma samples**

Endoscopically obtained biopsy samples from CRC (stage II, moderately differentiated tumors from sigmoid colon and rectum; n = 49) areas and paired histologically normal colonic mucosa (n = 49) were taken during routine colonoscopy and stored in RNALater Reagent (Qiagen Inc, Germantown, US) at −80°C until further processing. Total RNA was extracted and Affymetrix microarray analysis was performed as described before [21]. In the laser capture microdissection study, surgically removed NAT (n = 6) and CRC (n = 6) samples were used, which were embedded in TissueTek OCT compound (Sakura Finetek, Japan). Series of 6 μm sections were mounted onto PALM Membrane Slide 1.0 PEN (Carl Zeiss, Bernried, Germany) at −20°C and were stored at −80°C. Slides were fixed at 70% and absolute ethanol, then stained with cresyl violet acetate (Sigma-Aldrich, St. Louis, USA).

**Cell culture, proliferation and apoptosis assays**

Human CRC cell line HCT116 (from István Peták M.D., Ph.D., 1st Department of Pathology and Experimental Cancer Research, Semmelweis University, Budapest, Hungary) were cultured in DMEM (Sigma-Aldrich, Irvine, UK) supplemented with L-glutamine, 10% fetal bovine serum (FBS, Sigma-Aldrich, Irvine, UK) and 1% penicillin-streptomycin mixture (Sigma-Aldrich, Irvine, UK), then grown at 37°C in an atmosphere of 5% CO₂ and 95% humidity.

For the assays, cells were seeded at a density of 70,000 cells/cm² in collagen-treated 24-well culture plates (Greiner Bio One, Frickenhausen, Germany) and treatments were carried out in duplicate (using three wells for each concentration). Cells were treated for 48 hours with SFRP1 full length recombinant protein (Abcam, ab64445, Cambridge, UK) at concentrations of 0.1 μg/ml and 1.0 μg/ml. During the treatment fetal bovine serum deprived cultures were used to avoid the interaction with SFRP1.

The treated and control cells were harvested and fixed for 30 minutes at room temperature in 70% ethanol (−20°C) and stored at −20°C until further analysis. DNA was extracted with alkaline buffer (200 mM di-sodium phosphate, pH 7.8, adjusted with 200 mM citric acid) supplemented with 100 μg/ml RNase A (Sigma-Aldrich, Irvine, UK) followed by 10 μg/ml propidium iodide (Sigma-Aldrich, Irvine, UK) staining and incubated for 15 minutes at room temperature. 10,000–20,000 events were measured per sample by FACSscan flow cytometer (Becton Dickinson FACScan, CA, USA), and the analyses were performed by Winlist software (Verity Software House).

**Immunohistochemistry**

For immunohistochemistry normal biopsy samples (n = 20), surgically removed CRC (n = 35) and colonic tissues containing NAT and CRC areas (n = 14) were used. Samples were fixed in formaldehyde, embedded in paraffin and 4 μm thick sections were cut. Following deparaffinization and rehydration, microwave-based antigen retrieval was performed in TRIS EDTA buffer (pH 9.0) (900 W/10 minutes, then 340 W/40 minutes). SFRP1 protein expression was detected using anti-SFRP1 polyclonal antibody (ab4193, Abcam, Cambridge, UK, 1:500 dilution) with diaminobenzidine - hydrogen peroxidase - chromogen-substrate system (Cytomation Liquid DAB + Substrate Chromogen System, K3468, Dako, CA, USA). This antibody showed high specificity against SFRP1 protein which was confirmed by Western blot (http://www.abcam.com/sfrp1-antibody-ab4193.html). Scoring of SFRP1 protein expression representing the intensity of the immunohistochemical reaction (both in myofibroblasts and epithelial cells) was made on the basis of the following criteria: the scoring value was −2 for no reaction, 0 for weak, 1 for moderate, and 2 for strong cytoplasmic protein expression.

To identify the stromal source of SFRP1 protein α-SMA/SFRP1 dual fluorescent staining was used in samples that contained NAT tissues and CRC areas as well (n = 9). The slides were incubated with anti-α-SMA (Dako, CA, USA) 1:1 solution with Alexa Fluor 488 (Invitrogen, Eugene, CA, USA) and anti-SFRP1 (ab4193, Abcam, Cambridge, UK) 1:500 dilution with Alexa Fluor 568 (Molecular Probes, Eugene, OR, USA) and 1% penicillin-streptomycin mixture (Sigma-Aldrich, St. Louis, USA) followed by 10,000–20,000 events were measured per sample by FACScan flow cytometer (Becton Dickinson FACScan, CA, USA), and the analyses were performed by Winlist software (Verity Software House).

**DNA methylation analysis of laser microdissected α-SMA positive myofibroblasts and epithelial cells**

Surgically removed stage II, moderately differentiated CRC (n = 8) and paired NAT (n = 8) samples (3 slides/sample) were snap-frozen in liquid nitrogen and stored at −80°C until further use. 1,000 myofibroblasts/slides (6 samples; 18 slides) were collected based on their α-SMA positivity, while epithelial cells were readily recognizable without specific staining (H&E stain; 10 samples; 30 slides). To analyze the methylation status of the SFRP1 gene promoter, laser capture microdissected samples were bisulfite converted using the EZ DNA Methylation-Direct Kit™ (Zymo Research, Irvine, USA) according to the manufacturer’s protocol. A region on the promoter was amplified using unbiased BS-PCR primers that do not differentiate between methylated and unmethylated bisulfite modified DNA. (F: 5'-GGAAAGAGA-TAAGGGGAGAAAAAGAA-3'; R: 5'-ATTTCATAAAATTATGAAATCTATATCCACTGC-3'). These primers were designed using PyroMark Assay Design Software 2.0. They amplify a region 230 bp upstream of the transcription startpoint that contains 6 CpG sites (Figure S1). The specificity of the primers was tested in silico by the BiSearch software.

PCR cycling and HRM were performed on a LightCycler 480 instrument (Roche, Basel, Switzerland). The 25 μl reaction mix measured per sample by FACSscan flow cytometer (Becton Dickinson FACScan, CA, USA), and the analyses were performed by Winlist software (Verity Software House).
consisted of a final concentration of 1x AmpliTaq Gold 360 PCR Master Mix (Life Technologies, Carlsbad, USA), 1x ResoLight HRM Dye (Roche, Basel, Switzerland), 200 nM forward and reverse primers and 1 ng/well bisulfite converted DNA template. Cycling conditions were as follows: 95°C (10 min), 95°C (30 s), 60°C (30 s) decreased by 0.5°C in every cycle, 72°C (30 s) for 10 touchdown PCR cycles followed by, 95°C (30 s), 56°C (30 s) and 72°C (30 s) in 40 PCR cycles. On completion of the PCR thermal cycling, for the HRM analysis the samples were denatured at 95°C for 1 min, cooled down to 40°C and held for 1 min, then continuously warmed up to 95°C at a rate of 0.03°C/second during the melting curve fluorescence acquisition. For calibration of the analysis, 0% and a 100% artificially methylated control DNA samples were used (EpiTect PCR Control DNA Set, Qiagen, Hilden, Germany). To identify characteristic melting profiles of DNA products the LightCycler 480 Instrument Tm Calling analysis software Module was used.

Results

Reduced stromal SFRP1 mRNA expression in CRC compared to NAT

Whole biopsy (macrodissected) samples from NAT and CRC regions were compared. Macrodissected NAT samples showed significantly (P<0.001) higher SFRP1 mRNA expression than CRC samples (Figure 1/A). Using laser microdissected tissue compartments, higher (although not statistically significant) SFRP1 mRNA expressions were found in the NAT epithelium compared to CRC epithelium (Figure 1/B). However, a significant (P<0.001) difference was observed in the expression of
transcripts (202036_s_at and 202037_s_at) between stromal samples from NAT and CRC. The average gene expression values in CRC was one-third (0.38 and 0.36 respectively) of the average expression values observed in NAT samples (Figure 1/C, Table 1). Abnormal Wnt signaling in parallel with reduced SFRP1 was analyzed with Affymetrix whole genome microarrays. Based on 38 normal and 38 CRC samples (GSE37364), genes with significantly different (logFC > |2|, p < 0.001) expression levels in the tumors with reduced SFRP1 expression compared to healthy controls were listed in Table S1.

<table>
<thead>
<tr>
<th>analyzed cells</th>
<th>NAT</th>
<th>Tumor</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>mRNA expression</td>
<td>stroma (LCM)</td>
<td>logFC_{CRC-N} = -2.23</td>
<td>n = 38</td>
</tr>
<tr>
<td>DNA methylation</td>
<td>α-SMA myofibroblasts (LCM)</td>
<td>0%</td>
<td>60%</td>
</tr>
<tr>
<td>protein expression</td>
<td>α-SMA+ SFRP1+/α-SMA cells (LCM)</td>
<td>85.37%±12.6%</td>
<td>27.65%±18.2%</td>
</tr>
</tbody>
</table>

n=3/3 | n=3/3 |

n=9/9 | n=9/9 |

LCM: laser capture microdissection.

SFRP1 transcripts (202036_s_at and 202037_s_at) between stromal samples from NAT and CRC. The average gene expression values in CRC was one-third (0.38 and 0.36 respectively) of the average expression values observed in NAT samples (Figure 1/C, Table 1). Abnormal Wnt signaling in parallel with reduced SFRP1 was analyzed with Affymetrix whole genome microarrays. Based on 38 normal and 38 CRC samples (GSE37364), genes with significantly different (logFC > |2|, p < 0.001) expression levels in the tumors with reduced SFRP1 expression compared to healthy controls were listed in Table S1.

Figure 2. SFRP1 protein expression in α-SMA+ myofibroblasts in NAT, TA and CRC areas in representative photomicrographs. SFRP1 protein expression (red immunofluorescent staining) in whole slide (A) which includes NAT (white arrow), TA (yellow arrow) and CRC (red arrow). Both in NAT region (in 9/9 cases; B) and TA (in 4/9 cases; C) α-SMA+ myofibroblasts (E, F) showed retained SFRP1 expression (85.73±12.61%) (B, C; merged pictures: H, I). Around most CRC glands α-SMA+ myofibroblasts (green immunofluorescent staining; G) showed reduced (P<0.001) SFRP1 expression (27.65%±18.27%; in 9/9 cases) (D; merged picture: J).
Stromal and epithelial SFRP1 protein expression in normal, NAT, TA and CRC

Strong stromal SFRP1 protein expression (score +2) was found in normal, NAT and TA tissues, which localized mainly in the pericryptal region. These stromal SFRP1+ cells were identified as α-SMA+ myofibroblasts in NAT and in TA by dual fluorescent immunohistochemistry. Both α-SMA and SFRP1 showed strong (score +2) or moderate (score +1), diffuse, cytoplasmic staining in the elongated stromal cells. In NAT regions (9/9 cases) pericryptal α-SMA+ cells showed strong SFRP1 protein expression (85.73±12.61%; all α-SMA and SFRP1 double positive cells/all α-SMA positive cells in stroma; Figure 2/B, E, H, Table 1). Interestingly pericryptal α-SMA+ myofibroblasts retained strong or moderate SFRP1 protein expression in TAs (this area was identifiable in 4/9 cases), but the frequency of α-SMA/SFRP1 double positive cells was lower (72.32±2.32%) than in NAT tissues (Figure 2/C, F, I). In most of the analyzed CRC samples heterogeneous SFRP1 expression was detected. The number of α-SMA+ cells increased significantly, but the SFRP1 protein expression was significantly reduced (27.65±18.27%) in representative tumoral areas (in 9/9 cases) as compared to NAT (Figure 2/D, G, J, Table 1).

In normal samples epithelial SFRP1 expression was moderate (score: +1), lower than it was found in myofibroblasts (score: +2). Decreased SFRP1 expression was observed in 57.14% of NAT samples (8 of 14 samples) in crypts which localized closely to the tumor (Figure 3/B) and TAs (Figure 3/C). Parallel with the disappearance of epithelial SFRP1 protein expression, subepithelial myofibroblasts showed retained protein expression in these areas (Figure 3/B, C). NAT regions which localized more distantly from CRC showed similar SFRP1 protein expression patterns than normal samples (Figure 3/A). Decreased epithelial SFRP1 protein expression (score: 0) was detected in 71.42% of CRC samples (34 of 48+35 samples).

SFRP1 recombinant protein induces apoptosis in HCT116 cells

We analyzed the apoptotic effect of SFRP1 recombinant protein in HCT116 (APC wild type, β-catenin mutant) cells by flow cytometry. As the anti-proliferative and apoptotic effect of SFRP1 is well-known [12,23], we examined these functions of SFRP1 at low concentrations (exogenously administered recombinant human SFRP1 (rhSFRP1; Abcam, Cambridge, UK); 0.1 μg/ml and 1.0 μg/ml) as a potential model of paracrine signaling. Comparative analysis of treated versus non-treated specimens showed measurable (both not statistically significant) increase of apoptosis in a dose-dependent manner (control: 10.66±0.7; 0.1 μg/ml: 12.29±0.88; 1.0 μg/ml: 16.13±1.68) as measured 48 hours after a single dose of rhSFRP1 protein (Figure S2).

SFRP1 promoter hypermethylation in the NAT area and CRC

The methylation status of α-SMA immunopositive stromal myofibroblasts was examined in laser microdissected cells from NAT and CRC samples. The BS-PCR products of 100% methylated control with a higher melting temperature (melting peak: 79.6°C) and unmethylated control with a lower melting temperature (melting peak: 77.1°C) (Figure 4/A) were used. Myofibroblasts from NAT showed only one melting peak in the unmethylated temperature range, while myofibroblasts from CRC had an additional peak at higher temperature indicating a partially hypermethylated status of SFRP1 promoter in CRC (Figure 4/B, Table 1). Double peaks on the melting temperature (Tm) curves of CRC samples denoted their heterogeneity considering their methylation status, which was probably due to the presence of...
The temperature (Tm) of 100% methylated standard was 79.6°C, while the Tm of 0% methylated standard was 77.1°C. The melting curve of unmethylated products appeared, suggesting that SFRP1 is hypermethylated in NAT samples which localized closely to the crypts of most NAT samples. SFRP1 transcription was found in morphologically normal gland epithelium in TA, which was characterized by altered crypt morphology (e.g. elongation, pseudostratification, loss of cell polarity and nuclear polymorphism). On the contrary, epigenetic silencing of SFRP gene contributes to the development of several disorders including cancer [39–41]. Consistent with this duality, SFRP1 showed a biphasic effect on β-catenin stabilization elicited by Wingless, increasing β-catenin protein levels at low SFRP1 concentrations, but inhibiting it at high concentrations [42]. In different cellular contexts, SFRP1 has been shown either to increase or decrease β-catenin stabilization [43,44]. Furthermore, another study suggested that SFRP1 could stimulate the Wnt/calcium pathway via Frizzled-2 independently of endogenous Wnts [45].

Regarding field cancerization, methylation of hMLH1, CDKN2A/P16 and SFRP1 has been recently indicated to be associated with malignant transition in endometrial cancer [46].

In our study, the apoptotic effect of SFRP1 protein was demonstrated by administering low doses of rhSFRP1 on HCT116 CRC cell line. In HCT116 cells rhSFRP1 protein caused a measurable increase in apoptosis. We investigated the role of SFRP1 regarding the stroma-epithelium interaction in CRC and NAT areas. SFRP1 protein is a well-known intercellular inhibitor of Wnt pathway which regulates the epithelial proliferation as autocrine and/or paracrine signal [12,23]. We identified α-SMA+ myofibroblasts as the main source of stromal SFRP1 protein in NAT regions, in parallel with unmethylated SFRP1 promoter sequences. We also observed moderate to strong SFRP1 protein expression in α-SMA+ myofibroblasts in TA, which was characterized by altered crypt morphology (e.g. elongation, pseudostratification, loss of cell polarity and nuclear polymorphism). On the contrary, epigenetic silencing of epithelial SFRP1 appeared in both areas and this could serve as an early sign of field cancerization.

According to our hypothesis, in these areas field cancerization caused epithelial proliferation, which was partly inhibited by the less dysregulated stroma, indicated by the lower frequency of SFRP1 hypermethylation in myofibroblasts. Thus, decreased epithelial SFRP1 protein expression may partly be replaced by stromal sources (i.e. by α-SMA+ myofibroblasts). In contrast to the DNA repair gene O6-methylguanine-DNA methyltransferase (MGMT) [10], which has been found to be methylated in the macroscopic field of some CRCs, in our study epithelial SFRP1 was not methylated by Wnt binding to a member of the Frizzled receptor family and co-receptors (LRP5 or LRP6) [26]. This results in the recruitment of Dishevelled to Frizzled and Axin to phosphorylated LRP5/6, leading to the dissociation of a β-catenin degradation complex. In the absence of Wnt this complex mediates the sequential phosphorylation of β-catenin, causing its ubiquitination and proteosomal degradation. Wnt stimulation allows the accumulation of hypophosphorylated β-catenin in the cytosol and its translocation into the nucleus, where it binds to TCF/LEF and promotes the expression of Wnt/β-catenin target genes [26]. Constitutive activation of this pathway is commonly present in many types of cancer [24,25]. Non-canonical Wnt-signaling pathways (planar cell polarity or increased intracellular calcium concentration) are transduced by Frizzleds and/or other Wnt receptors or co-receptors [26,27]. Several non-canonical Wnt signaling mechanisms have been reported to inhibit the β-catenin pathway by decreasing β-catenin/TCF association with DNA or increasing β-catenin turnover [28–30].

SFRPs comprise a family of five proteins in mammals that were first identified as antagonists of the Wnt/β-catenin pathway during embryonic development [31–33]. SFRPs possess a remarkable range of biological activities, including tumor suppression [11,34]. This is also strengthened by epigenetic silencing of SFRP gene expression in a wide variety of cancers [35,36], and supported by the observation that restoration of expression is suppressive of the tumor phenotype [37,38]. By contrast, SFRP overexpression has been observed in some of the same malignancies [39–41]. Consistent with this duality, SFRP1 showed a biphasic effect on β-catenin stabilization elicited by Wingless, increasing β-catenin protein levels at low SFRP1 concentrations, but inhibiting it at high concentrations [42]. In different cellular contexts, SFRP1 has been shown either to increase or decrease β-catenin stabilization [43,44]. Furthermore, another study suggested that SFRP1 could stimulate the Wnt/calcium pathway via Frizzled-2 independently of endogenous Wnts [45].

Figure 4. Promoter hypermethylation of SFRP1 gene in BS-PCR HRM. SFRP1 is hypermethylated in α-SMA positive myofibroblasts in CRC. A: Melting peaks of SFRP1 BS-PCR products. Melting temperature (Tm) of 0% methylated standard sample was 77.1°C, while Tm of 100% methylated standard was 79.6°C. B: There was only one unmethylated melting peak detectable for NAT samples (green curves). In CRC samples (red curves) an additional melting peak belonging to hypermethylated products appeared, suggesting that α-SMA positive myofibroblasts are heterogenous concerning SFRP1 promoter methylation.

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cells with unmethylated promoters among hypermethylated promoters. Decreased epithelial SFRP1 protein expressions were found in crypts of most NAT samples which localized closely to the tumor. This decreased protein expression was related to epigenetic silencing of SFRP1 gene. Partially SFRP1 promoter hypermethylation was found in morphologically normal gland epithelium in 40% (2 of 5 samples) of NAT samples (Figure S3).

Discussion

Wnt signaling is a major regulator of a variety of cellular processes during embryonic development and promotes tissue homeostasis in the adult. Wnts are secreted lipid-modified glycoproteins regulating a wide range of cellular behavior including differentiation, proliferation, migration, survival, polarity and stem cell self-renewal [24]. Altered Wnt signaling may contribute to the development of several disorders including cancer [25]. The canonical/β-catenin pathway is the most extensively studied Wnt signaling mechanism, which is triggered by Wnt binding to a member of the Frizzled receptor family and co-receptors (LRP5 or LRP6) [26]. This results in the recruitment of Dishevelled to Frizzled and Axin to phosphorylated LRP5/6, leading to the dissociation of a β-catenin degradation complex. In the absence of Wnt this complex mediates the sequential phosphorylation of β-catenin, causing its ubiquitination and proteosomal degradation. Wnt stimulation allows the accumulation of hypophosphorylated β-catenin in the cytosol and its translocation into the nucleus, where it binds to TCF/LEF and promotes the expression of Wnt/β-catenin target genes [26]. Constitutive activation of this pathway is commonly present in many types of cancer [24,25]. Non-canonical Wnt-signaling pathways (planar cell polarity or increased intracellular calcium concentration) are transduced by Frizzleds and/or other Wnt receptors or co-receptors [26,27]. Several non-canonical Wnt signaling mechanisms have been reported to inhibit the β-catenin pathway by decreasing β-catenin/TCF association with DNA or increasing β-catenin turnover [28–30].
hypermethylation was only present in areas closely localized to the tumor (NAT).

Although in most CRC cases stromal myofibroblasts exhibited hypermethylation and low protein expression of SFRP1, we hypothesize that the remaining myofibroblasts with intact SFRP1 protein expression can decrease the proliferation rate of Wnt mutant CRC cells.

To conclude, our present study indicates that SFRP1 protein expression is conserved in +SMA+ myofibroblasts and partially disappears in the epithelium of NAT and TA. In contrast, we found low SFRP1 immunopositivity stroma-associated myofibroblasts in CRC. Both epithelial and myofibroblast-derived SFRP1 expression is under epigenetic regulation. We presume that the paracrine, anti-proliferative effect of myofibroblast-derived SFRP1 may inhibit field cancerization in NAT and TA areas, and so cancer-associated myofibroblasts may support the uncontrolled tumor cell proliferation by an epigenetically silenced Wnt inhibitory signal.

Supporting Information

Figure S1 Position of the BS-PCR product (black) on the SFRP1 gene promoter. The transcript of SFRP1 is represented by the blue line; CpG sites are red lines. Figure modified from UCSC genome browser (Human, GRCh37/hg19 Assembly). (TIF)

Figure S2 Apoptotic effect of recombinant SFRP1 protein on HCT116 cell line. rhSFRP1 protein caused measurable (both not statistically significant) increase in apoptosis (48 hours after a single dose). (TIF)

Figure S3 Promoter hypermethylation of SFRP1 in epithelial cells in NAT area. A: Melting peaks of SFRP1-BS-PCR products. Melting temperature of the 0% methylated standard sample was 77°C, while Tm of the 100% methylated standard was 79.5°C. B: SFRP1 was hypermethylated in 2/5 epithelial cells (40%) in NAT area (red line) indicated by their melting peaks at higher temperature. (TIF)

Table S1 Genes with significantly different (logFC > [2], p < 0.001) expression levels in tumors with reduced SFRP1 expression compared to healthy controls. (DOCX)

Acknowledgments

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Author Contributions

Conceived and designed the experiments: GV AVP. Performed the experiments: GV AVP AK BP IF. Analyzed the data: GV AVP AK BP IF BW GM FS TK EM SS BM ZT. Contributed reagents/materials/analysis tools: TK EM SS BM ZT. Wrote the paper: GV AVP AK BP IF BW GM FS TK EM SS BM ZT. Statistical analysis: BW.

References


