Targeting cancer stem cells: Modulating apoptosis and stemness
Çolak, S.

Citation for published version (APA):
Çolak, S. (2016). Targeting cancer stem cells: Modulating apoptosis and stemness

General rights
It is not permitted to download or to forward/distribute the text or part of it without the consent of the author(s) and/or copyright holder(s), other than for strictly personal, individual use, unless the work is under an open content license (like Creative Commons).

Disclaimer/Complaints regulations
If you believe that digital publication of certain material infringes any of your rights or (privacy) interests, please let the Library know, stating your reasons. In case of a legitimate complaint, the Library will make the material inaccessible and/or remove it from the website. Please Ask the Library: http://uba.uva.nl/en/contact, or a letter to: Library of the University of Amsterdam, Secretariat, Singel 425, 1012 WP Amsterdam, The Netherlands. You will be contacted as soon as possible.
Chapter 6

Human colonic fibroblasts regulate stemness and chemotherapy resistance of colon cancer stem cells

S. Colak & J.P. Medema

Published in Cell Cycle, 2014
Abstract

There is increasing evidence that cancers are heterogeneous and contain a hierarchical organization consisting of cancer stem cells and their differentiated cell progeny. These cancer stem cells are at the core of the tumor as they represent the clonogenic cells within a tumor. Moreover, these cells are considered to contain selective therapy resistance, which suggests a pivotal role in therapy resistance and tumor relapse. Here we show that differentiated cells can re-acquire stemness through factors secreted from fibroblasts. This induced CSC state also coincides with re-acquisition of resistance to chemotherapy. Resistance induced in newly formed CSCs is mediated by the anti-apoptotic molecule BCLXL and inhibition of BCLXL with the BH3 mimic ABT-737 sensitizes these cancer cells towards chemotherapy. These data point to an important interplay between tumor cells and their microenvironment in the regulation of stemness and therapy resistance.
Introduction

Over the last decade it has become increasingly clear that a variety of cancers is hierarchically organized and cancer stem cells (CSCs) can be found at the top of this hierarchy. CSCs are, in contrast to their differentiated daughter cells, highly tumorigenic and besides their self-renewal capacity CSCs can differentiate in more differentiated cell types. In colorectal cancer CSCs were identified by two independent groups that made use of an antibody that recognize the cell surface molecule CD133. More recent data provided for evidence for other CSC markers in colon, such as LGR5 expression and Wnt pathway activity. Especially the latter was surprising as this occurred even in tumors where all cells contained a mutated APC, indicating that not all cells have the same level of Wnt pathway activation, despite this mutation. When primary colon cancer spheroid lines were dissociated and transduced with a Wnt activity reporter that directs the expression of enhanced green fluorescent protein (TOP-GFP), cells expressing high TOP-GFP levels (TOP-GFP^hi^) are shown to be the CSCs. In contrast, cells with low Wnt pathway activity (TOP-GFP^lo^) are not tumorigenic upon xenotransplantation and express markers of differentiated colon cells. Using these TOP-GFP spheroid cultures we have designed flow cytometer based assays in which cell death in CSCs and their differentiated progeny can be measured simultaneously and importantly under the same conditions. This assay showed that differentiated cells are killed by chemotherapy, while CSCs were resistant to all therapies that we have tested. Resistance was due to a disturbed apoptotic balance in CSCs in favor of anti-apoptotic molecules. This suggests that targeting this balance could be an effective therapy. However, we have recently also shown that signals emanating from the microenvironment are crucial in the regulation of stemness and can direct the acquisition of stemness features in more differentiated tumor cells. This suggests that the microenvironment is a pivotal player in the hierarchy within a tumor and could therefore also regulate chemotherapy sensitivity. Here we show that factors derived from myofibroblasts can not only dedifferentiate non-CSCs into CSCs, but also direct a resistance towards chemotherapy. Furthermore, this fibroblast induced resistance can be reverted by inhibition of BCLXL with the BH3 mimetic ABT-737.

Results

Factors secreted from fibroblasts can induce dedifferentiation of non-CSC

Myofibroblasts make up an important part of the microenvironment in colon cancers and direct CSC features. To study the effect of myofibroblast-secreted factors
on CSCs, myofibroblast-conditioned medium (MFCM) was derived from human colon 18Co fibroblasts. These fibroblasts model cancer-associated fibroblasts (CAFs) and similar observations can be obtained using CAFs (not shown). Spheroid cultures, containing CSCs and more differentiated tumor cells, were grown in the absence or presence of this MFCM and CSC markers CD133 and LGR5 were measured using flow cytometry. This revealed a clear increase in cells expressing these CSC markers (Figure 1a). In line with the idea that CSC numbers increased in these cultures, a concomitant decrease in the expression of differentiation markers Ck20 and Mucin2 was observed after MFCM exposure (Figure 1b). Increased stem cell marker expression in combination with decreased differentiation marker expression suggest a MFCM-induced increase in the stem cell fraction in our spheroid cultures. However, marker expression by itself is not sufficient to determine stem cell capacity, which in vitro can only be quantified using a limiting dilution assay. Spheroid cultures were therefore sorted in a limiting dilution fashion of either the CSCs (GFPhi) or the differentiated cells (GFPlow) cells. Confirming previous data, only CSCs were clonogenic while differentiated cells failed to grow. However, when differentiated cells were sorted in plates containing MFCM, clonogenicity was restored to levels of CSCs (Figure 1c), confirming the idea that fibroblast secrete factors that promote dedifferentiation of differentiated cells into CSCs. Previously, we observed that this reversal even resulted in the re-acquisition of tumor-initiating potential confirming the idea that MFCM can re-install cancer stemness in more differentiated cells.5

**Conditioned medium increases BCLXL protein expression and induces resistance to chemotherapy**

Recently, we have described an assay that allows for the study of chemotherapy sensitivity in differentiated cells and CSCs simultaneously.6 Using this assay we have shown that a differential sensitivity exists between CSCs, which are resistant and differentiated cells, which are chemosensitive. To study if factor secreted by myofibroblasts were also capable of inducing therapy resistance in differentiated cells, spheroid cultures were grown in control or in MFCM and subsequently treated with chemotherapy. In control medium, differentiated cells were sensitive to oxaliplatin when compared with CSCs. However, when pretreated for only 24 hours with MFCM oxaliplatin-induced cell death was significantly blocked, suggesting that, similar to the induction of clonogenicity, also therapy resistance can be restored by the microenvironment (Figure 2a). To further investigate the mechanism of chemotherapy resistance the anti-apoptotic molecule BCLXL was analyzed, as we have previously shown that colorectal
MFCM induces stemness and therapy resistance

CSCs are dependent on BCLXL for survival and therapy resistance. Upon MFCM exposure of spheroid cultures BCLXL protein expression was increased (Figure 2b and c), which is sufficient to block oxaliplatin-induced cell death in spheroid cultures.

**ABT-737 reverts MFCM-induced chemotherapy resistance**

Inhibition of apoptosis is one of the hallmarks of cancer. Therefore there are small molecule inhibitors in clinical trials that inhibit anti-apoptotic BCL2 molecules. ABT-737 inhibits BCLXL, BCL2 and BCLW and displays anti-tumorigenic activity.13, 14

---

**Figure 1: MFCM dedifferentiate differentiated cells to CSCs.** Spheroid cultures grown in control medium (red) or MFCM (brown) for 24h and were a) stained with stem cell marker Lgr5, CD133 or b) qrt-pcr was performed on differentiation markers Ck20 and Mucin2 (Muc2). c) Limiting dilutions experiments were performed and clonogenic fraction was calculated in CSC (TOP-GFP<sup>hi</sup>), Differentiated cell (TOP-GFP<sup>lo</sup>), and differentiated cells deposited in MFCM (Diff + MFCM). Significance is shown as ***p<0.001
To investigate if inhibition of these anti-apoptotic molecules is sufficient to sensitize cells that were exposed to MFCM cells were treated with ABT-737. Interestingly, differentiated cells that were exposed to MFCM were resistant to oxaliplatin. However, this road block in chemotherapy-induced apoptosis was lifted by ABT-737 (Figure 2d). This suggested that anti-apoptotic BCL2 proteins are crucial in MFCM induced-resistance and inhibition of these proteins using small molecule inhibitors is sufficient to sensitize to chemotherapy. Moreover, our data provide compelling evidence that factors secreted by fibroblast are able to induce CSC features in differentiated tumor cells, which include therapy resistance. These factors therefore are interesting candidates for therapeutic targeting in colorectal cancers.

**Discussion**

Despite the fact that only CSCs and not the more differentiated cancer cells are highly tumorigenic in xenotransplantation assays, differentiated cells can indirectly become tumorigenic by dedifferentiation.\(^5\), \(^7\), \(^8\), \(^15\), \(^16\) Here we show that human colonic fibroblast secrete factors that increases the stem cell fraction in spheroid cultures. In addition, increased anti-apoptotic BCLXL protein expression and resistance towards chemotherapy were observed. This MFCM induces resistance could be overcome by the BH3 mimetic ABT-737. Inhibition of BCLXL by ABT-737 is therefore likely to be sufficient to revert the effects of myofibroblasts. There is evidence that depletion of colorectal CSCs does not block tumor growth.\(^17\) This is analogous to normal intestinal stem cells. In the healthy intestine stem cells have been identified using LGR5 as marker.\(^18\) Depletion of LGR5-positive stem cells in the intestinal epithelium using a diphtheria toxin receptor driven by the LGR5-promoter, does not disturb homeostasis as would be expected after killing a stem cell population. Distinct stem cells, namely Bmi1-expressing stem cells were able to compensate for the loss of Lgr5-expressing cells.\(^19\) Besides highly proliferative LGR5-positive stem cells, there are also quiescent LGR5 expressing cells, also called label-retaining cells, that express Paneth and enteroendocrine cell markers. These cells are able to repopulate intestinal epithelial cells when there is intestinal damage, suggesting that these label-retaining cells are reserve stem cell population in the intestine.\(^20\)

In addition, high Notch ligand expressing cells are described in the mouse intestine, which represent progenitor cells derived from the LGR5-stem cells. These cells are already specified to a secretory lineage, but can dedifferentiate and replenish the pool of LGR5-positive stem cells.\(^21\) The same phenomenon of a bidirectional relationship
Figure 2: BCLXL is required for MFCM induced resistance towards oxaliplatin

a) Primary spheroid culture transduced with TOP-GFP construct (Co100) was exposed to MFCM or control medium for 24h. Subsequently, cells were treated for 24h with oxaliplatin and Annexin V/7AAD staining was performed. CSCs and differentiated cells were defined by gating on TOP-GFP<sup>hi</sup> and TOP-GFP<sup>lo</sup> expressing cells, respectively. MFCM treatment decreases oxaliplatin induced cell death.

b) Western blot analysis of BCLXL protein in spheroid cultures grown in control or MFCM for 24h. Increased BCLXL expression in spheroid culture treated with MFCM compared to control medium. Control western blots for ERK1/2 is shown in lower panel.

c) Quantification of BCLXL protein intensity relative to ERK1/2 protein expression. Average of three independent experiment is shown.

d) Co100 spheroid culture was exposed to MFCM for 24h. Cells were treated with 100nM ABT-737 in combination with or without oxaliplatin. Cell death was measured with Annexin V/7-AAD staining in TOP-GFP<sup>lo</sup> cells. Significance is indicated (*p<0.05, **p<0.01, ***p<0.001)
between stem cells and differentiated progeny is observed during intestinal tumorigenesis. APC mutations in stem cells was suggested to effectively lead to intestinal tumors, while more differentiated cells would not fully transform. However, differentiated cells can be the tumor initiating cells when the NFKB pathway is co-activated, leading to enhanced nuclear $\beta$-catenin activity. In other words dedifferentiation can occur under the right conditions and result in tumor-initiating capacities. Tumors are surrounded by stromal cells, including myofibroblasts, which we have previously shown to install a CSC-phenotype in differentiated tumor cells, restoring their clonogenic potential. The dominant factor in MFCM was shown to be hepatocyte growth factor (HGF). Next to HGF, osteopontin and stromal-derived factor 1a can revert differentiated colorectal cancer cells into metastatic CSCs. In addition, it was shown that the microenvironment can even enhance the metastatic potential of non-metastatic progenitors and as such also form metastatic CSCs. Similar observations of plasticity have been made in other tissues. For instance, in mammary epithelium, stochastic conversion of differentiated cells occurs at low levels during homeostatic conditions. Moreover, oncogenic mutations increase the conversion of differentiated cells into stem cells, which then can act as tumor-initiating cells. Similarly, activation of the RAS signaling pathway, by NF1 deletion, and inhibition of p53 tumor suppression in terminally differentiated neurons and astrocytes can also induce stem cell markers and tumorigenicity. Finally, in breast cancer induction of epithelial-mesenchymal transition (EMT) by overexpression of TWIST or SNAIL or exposing cells to TGFβ strongly enhances the CSCs phenotype while, mesenchymal stem cells and cancer stem cells regulate stemness by a network of CXCL6 and IL6. Importantly, transitions between more differentiated cancer cells and CSCs may also represent a mere stochastic process as suggested by the group of Lander. When differentiated breast cancer cells were sorted, these cells were able to form stem cells. Interestingly, these differentiated are not tumorigenic in vivo, but become tumorigenic when injected with irradiated carrier cells, which gives them the time to revert to CSCs in vivo. In line with these observations, we have shown that differentiated glioblastoma cells can re-acquire stem cell traits under the influence of endothelial like cells (unpublished observations).

Therefore, these lines of evidence suggest that plasticity in the form of a bidirectional relationship between stem cells and their differentiated progeny is present and is increased in the presence of oncogenic mutations, as is the case within a tumor. Besides the fact that CSCs are tumorigenic it is now well accepted that CSCs are also resistant to tumor therapy. By measuring mitochondrial priming in CSCs and differentiated cells we have shown that mitochondria of CSCs are less primed
MFCM induces stemness and therapy resistance to death, which is due to an elevated anti-apoptotic machinery. Targeting the anti-apoptotic molecule BCLXL directly, resulted in cell death of CSCs, while sublethal doses of the BH3 mimetics allowed for mitochondrial priming of CSCs, which then sensitizes CSCs towards various chemotherapies. Likewise to our previous study, here we used sublethal doses of ABT-737 that did not induce cell death but sensitized chemotherapy resistant cells towards oxaliplatin. This combination treatment resulted in more than just an additional effect of ABT-737 on oxaliplatin. It is worth to mention that there is increasing evidence that also CSCs from other cancers may depend on BCLXL. In a recent study it is shown that chemotherapy resistant lung CSCs require BCLXL for resistance and treatment with ABT-737 eliminated these CSCs in vitro and in vivo. This indicates that our findings are not only restricted to colorectal cancer, but can also be important for other cancers. Therapy resistance of CSCs and plasticity has important implications when treating cancer. Therapy aimed at CSCs might not be effective, since CSC-pools will readily be restored by reversion of the differentiated progeny. Therefore, to kill all tumor cells and overcome relapses caused by remaining or reverted CSCs, more study needs to be done on regulation of plasticity of CSCs and their tumor therapy resistance mechanisms. Nevertheless, targeting microenvironmental factors or the anti-apoptotic features induced by the microenvironment could prove useful new avenues for therapy.

Materials and Methods

Cell culture and reagents
Colon spheroid cultures were derived from colorectal cancer patients and maintained as previously described in stem cell medium (advanced DMEM/F12 (Gibco) supplemented with N2 Supplement (Gibco), 6 mg/ml glucose, 5 mM HEPES, 2 mM l-glutamine, 4 μg/ml heparin, epidermal growth factor (50 ng/ml) and basic fibroblast growth factor (10 ng/ml). After lentiviral transduction with TCF/LEF reporter driving expression of GFP (TOP-GFP) cells were single-cell plated in 96-well ultra-low adhesion plates (Corning) with FACSaria (BD Biosciences). This TOP-GFP vector was a gift from Dr. Laurie Ailles and was described previously.

Myofibroblast 18Co cells were purchased from the American Type Culture Collection and maintained in DMEM medium supplemented with 10% FCS and 1% glutamine. To obtain MFCM, 1.5 × 10^6 18Co cells were seeded in 75-cm² flasks. The next day, cells were washed five times with PBS and incubated for 24 h with 10 ml of CSC medium without EGF and bFGF. MFCM was then collected, cleared by centrifugation and supplemented with EGF (50 ng/ml) and bFGF (10 ng/ml) prior to addition to the cells.
RNA extraction, cDNA synthesis and RT-PCR

Extraction of RNA from cells was performed with Trizol reagent (Invitrogen) in accordance with the manufacturer’s protocol. After quantification of RNA using NanoDrop ND-2000 (Thermo Scientific) 2μg of RNA was used to synthesis cDNA using Super-Script III in accordance with the manufacturer’s protocol (Invitrogen). RT-PCR was performed with LC480 SYBR green (Roche) in accordance with the manufacturer’s instructions on a LC480. The following primers were used: 18S sense: 5'-AGACAACAA-GCTCCGTGAAGA-3', 18S antisense: 5'-CAGAAGTGACGCAGCCCTCTCTA-3'; Mucin 2 sense: 5'-CGAACCACGCCCACAACGT-3', Mucin 2 anti-sense: 5'-GACCACGG-GCCCGTTAAGCA -3', Ck20 sense: 5'- TGTCTCTGCAAATTGATAATGCT -3', and Ck20 anti-sense: 5'- AGACGTATTCCTCTCTACTCTCACTCTATA -3'.

Immunoblotting and antibodies

Spheroid cultures were grown in MFCM or control medium and 24 h later cultures were lysed in 1 × RIPA lysis and extraction buffer (Thermo Fisher Scientific) supplemented with complete protease inhibitor (Roche). Subsequently, lysates were cleared by centrifugation (14000 rpm, 10 min, 4°C). After quantification of protein concentration using BCA protein assay (Thermo Scientific), 20μg extracted protein was separated on 12% precast gels (Bio-Rad) and transferred to Hybond-P membranes (Amersham). Membranes were blocked with 5% milk in phosphate-buffered saline solution containing 0.2% TWEEN (PBS-T) for 1h. The membranes were incubated with anti-BCLXL (clone S18, 1:500, Santa-Cruz) in 2.5% milk/PBS-T overnight at 4°C and subsequently washed three times with PBS-T. Membranes were then incubated with anti-mouse IgG horseradish peroxidase conjugates for 1h, washed three times and detection of bound antibody was performed with ECLplus reagents (Amersham). Western blots were analyzed by LAS4000. For loading control, blots were incubated with anti-ERK1/2 (1:10000, Cell Signaling).

Limiting-dilution assay

Spheroid cultures were dissociated and CSCs (10% TOP-GFP<sup>hi</sup>) or differentiated (10%TOP-GFP<sup>lo</sup>) cells were FACS deposited using the FACSaria (BD Biosciences) in a limiting dilution fashion at 1, 2, 4, 8, 16, 24, 32, 40, and 64 cells per well in ultra-low 96-well plates (Corning). CSCs were sorted in control medium, while differentiated cells were sorted in either control medium or MFCM. Clonal frequency was evaluated after two weeks and calculated using the Extreme Limiting Dilution Analysis ‘limdil’ function as described.10

136
**FACS staining**
Colon spheroid cultures were grown in control medium or MFCM. After 24 h cells were dissociated and stained with AC133/CD133-APC antibody (1:25, Miltenyi Biotec) or anti-LGR5-biotin antibody (4D11F8, 1:100, BD Biosciences) in PBS containing 1% bovine serum albumin (PBS-B) for 30 min at 4°C. Subsequently, cells were washed with PBS-B and resuspended in PBS-B. For LGR5 staining, cells were incubated with APC conjugated streptavidin (1:500, E biosciences) and washed twice with PBS-B. Dead cells were excluded with 7-AAD (BD Biosciences). Stainings were analyzed on a FASCanto (BD Biosciences).

**Cell death assay**
TOP-GFP transduced spheroid culture Co100 was seeded as single cells on an adherent cell culture 12-well plate (Greiner) overnight. The next day, adherent cells were refreshed with control medium or MFCM and 24 h later cells were left untreated or treated with 100 nM ABT-737 (Selleck Chemicals) and/or 50 μM oxaliplatin. After 24 h treatment cells were resuspended to single cells using trypsin-EDTA. 50,000 cells were washed with PBS and stained with Annexin V-APC (BD biosciences) and 7-AAD (BD biosciences) for 15 min at RT, followed by flow cytometry performed with FASCanto (BD biosciences). Cell death (AnnexinV+) was measured in CSCs by gating on TOP-GFP<sup>hi</sup> cells and in differentiated tumor cells by gating on TOP-GFP<sup>lo</sup> cells.

**Acknowledgements**
We thank the members of the laboratory for useful discussion. In addition, we thank Kate Cameron, Berend Hooibrink and Toni van Capel for assistance with fluorescence-activated cell sorting experiments. We also thank Laurie Ailles (University of Toronto) for providing Tcf/Lef reporter construct. JPM is sponsored by grants from the Netherlands Organization for Scientific Research (NWO; Gravitation-Cancer Genomics Center The Netherlands Zwaartekracht), from the Dutch Cancer Society (UVA2009-4416 and UVA2012-5735), from MLDS (FP13-07) and Alpe d'Huzes/KWF (CONNECTION).

**Conflict of interest**
The authors declare no conflict of interest.
References


MFCM induces stemness and therapy resistance


