Effects of EpCAM overexpression on human breast cancer cell lines

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Abstract

Background: Recently, EpCAM has attracted major interest as a target for antibody- and vaccine-based cancer immunotherapies. In breast cancer, the EpCAM antigen is overexpressed in 30-40% of all cases and this increased expression correlates with poor prognosis. The use of EpCAM-specific monoclonal antibodies is a promising treatment approach in these patients.

Methods: In order to explore molecular changes following EpCAM overexpression, we investigated changes of the transcriptome upon EpCAM gene expression in commercially available human breast cancer cell lines Hs578T and MDA-MB-231. To assess cell proliferation, a tetrazolium salt based assay was performed. A TCF/LEF Reporter Kit was used to measure the transcriptional activity of the Wnt/β-catenin pathway. To evaluate the accumulation of β-catenin in the nucleus, a subcellular fractionation assay was performed.

Results: For the first time we could show that expression profiling data of EpCAM transfected cell lines Hs578TEpCAM and MDA-MB-231EpCAM indicate an association of EpCAM overexpression with the downregulation of the Wnt signaling inhibitors SFRP1 and TCF7L2. Confirmation of increased Wnt signaling was provided by a TCF/LEF reporter kit and by the finding of the nuclear accumulation of β-catenin for MDA-MB-231EpCAM but not Hs578TEpCAM cells. In Hs578T cells, an increase of proliferation and chemosensitivity to Docetaxel was associated with EpCAM overexpression.

Conclusions: These data show a cell type dependent modification of Wnt signaling components after EpCAM overexpression in breast cancer cell lines, which results in marginal functional changes. Further investigations on the interaction of EpCAM with SFRP1 and TCF7L2 and on additional factors, which may be causal for changes upon EpCAM overexpression, will help to characterize unique molecular properties of EpCAM-positive breast cancer cells.

Background

EpCAM is frequently overexpressed in human invasive breast cancer [1]. We reported EpCAM overexpression to be an independent prognostic marker for poor overall survival in node-positive breast cancer [2,3]. An independent group confirmed this finding in node-negative patients too [4]. Moreover, the magnitude of EpCAM antigen expression semiquantitatively assessed by immunohistochemistry showed a dose-dependent relationship with survival. In this retrospective analysis the patient subgroup with breast carcinomas overexpressing both EpCAM and Her-2/neu had the worst prognosis [5]. Targeting EpCAM with a humanized monoclonal antibody (Adecatumumab) in a randomized multi-centre phase II trial for the treatment of metastatic breast cancer yielded an expression- and dose-dependent reduction in formation of new metastatic lesions [6]. Recently, a trifunctional anti-EpCAM antibody (Catumaxomab) has received approval by the European Commission for the treatment of patients with EpCAM positive tumours [7].

EpCAM has initially been identified as a cell adhesion molecule located within intercellular adherens junctions, where it modulates cadherin-mediated cell adhesion and promotes epithelial cell migration and proliferation. EpCAM expression is not only involved in epithelium

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formation and epithelial-mesenchymal transition during organ development and tissue repair but also contributes to epithelial cell transformation [8,9]. Regarding EpCAM target genes, overexpression of EpCAM was found to be associated with enhanced transcription and translation of the proto-oncogene c-myc and the cell cycle proteins cyclin A and E in human epithelial 293 cells as well as in murine NIH3T3 fibroblasts [10]. Furthermore, proteome analysis revealed the epidermal fatty acid binding protein E-FABP, a major target of c-myc, to be upregulated upon EpCAM expression in HEK293 cells. Enhanced E-FABP expression correlated with a mitogenic signal, participating in a multimeric nuclear complex together with FHL2, β-catenin and Lef-1 for the induction of target gene transcription in FaDu hypopharynx and HCT-8 colon carcinoma cells [11]. Very recently, the proteolytic shedding of the intracellular domain of EpCAM (EpICD) was shown to confer a mitogenic signal, participating in a multimeric nuclear complex together with FHL2, β-catenin and Lef-1 for the induction of target gene transcription in FaDu hypopharynx and HCT-8 colon carcinoma cells [11].

Methods

Cell lines and plasmids

All cell lines were obtained from American Type Culture Collection. Hs578T, MDA-MB-231 and MCF-7 cell lines were grown in phenol red free MEM medium (PAA) supplemented with 5 mM glutamine (Gibco) and 10% fetal bovine serum (FBS superior, Biochrom) at 37°C under 5% CO2. SKBr-3 cells were cultivated in McCoy’s (Gibco) substituted with 5 mM glutamine and 10% FBS. A mixture of proteolytic and collagenolytic enzymes (Accutase, PAA) was used for detachment of the adherent cells for passaging. All cell lines were cultivated at maximum to passage 25 (3 months) and the absence of mycoplas contamination was controlled by using Venorgem PCR based detection kit (Minerva Biolabs).

EpCAM coding sequence was cloned into the pIRE-Spuro3 vector (Clontech) to get a high CMV promoter driven protein expression. Stable EpCAM expressing cell lines as well as respective empty vector control lines were established by transfections using Nucleofector Kit V (Amaxa) followed by puromycin selection.

Western blotting

50 μg of protein heated in Laemmli sample buffer was loaded per lane, resolved by a 4-20% SDS-PAGE (Criterion, Biorad) and transferred onto a nitrocellulose membrane (0.2 μm pore size, Schleicher & Schuell) for western blotting at 150 mA for 2 hours. To control protein transfer, the membrane was incubated 5 min in a Ponceau S staining solution until protein lanes were visible. After washing steps with TBS-T, the membrane was blocked 1 hour with 10% non-fat dry milk diluted in TBS-T. Primary antibodies were incubated in 1% non-fat dry milk diluted in TBS-T at 4°C over night. Secondary HRP-labelled antibodies were incubated for 1 hour at room temperature. After incubation with ECL Western blot detection reagent (Amersham), the detection followed by exposing the membrane to an X-ray film (Agfa).

Antibodies used for Western analysis were: C-10 (mouse monoclonal against amino acids 24-93 of human EpCAM, Santa Cruz Biotechnology), E144 (rabbit monoclonal antibody against the C-terminus of human EpCAM, Epitomics), β-Catenin (BD Biosciences), pan-actin (Ab-5 Clone, ACTN05, Neomarkers), GAPDH 6C5 (Santa Cruz Biotechnology), lamin A/C (2032, Cell Signalling) and anti-TCF7L2 (clone 6H5-3, Upstate Biotechnology).

Immunofluorescence

Cells were plated at different densities on non-coated or gelatine-coated glass coverslips and prepared for immunofluorescence after additional 48 h hours. Cells were
washed 3 times in PBS, extracted in 2% PFA with 0.3% Triton X-100 in PBS for 5 minutes and fixed in 2% PFA in PBS for 30 minutes. Primary antibodies (NCL-ESA, 1:100, mouse monoclonal anti-EpCAM, clone VU1D9, Novocastra and C-10, 1:50, Santa Cruz Biotechnology) were used with the respective Alexa 488 labelled secondary serum (Molecular probes). F-actin was visualized using Alexa 568 Phallodin (Molecular probes).

Confocal microscopy was performed with a micro lens-enhanced Nipkow disk-based confocal system UltraVIEW RS (Perkin Elmer, Wellesley MA, USA) mounted on an Olympus IX-70 inverse microscope (Olympus, Nagano, Japan). Images were acquired using the UltraVIEW RS software (Perkin Elmer).

Subcellular fractionation
To generate nuclear and cytoplasmic lysates NE-PER fractionation kit was used (Pierce 78833). A stepwise lysis of cells generated both functional cytoplasmic and nuclear protein extracts. Protocol was performed according manufacturer’s instructions.

Cell counting kit
Proliferation and cell viability analysis was performed by using Cell Counting Kit-8 (CCK-8, Dojindo Molecular Technology) according manufacturer’s instructions. Cells plated in 96-well plates (5 replica wells for each cell line). At intervals of 24 hours, CCK-8 solution was added directly to the cells (dilution 1:10) and the absorbance at 450 nm was measured after 4 hours of incubation in a microplate reader (Biorad). Cell numbers of non transfected cells have been quantified once in parallel with the respective empty vector control lines.

Chemotherapeutic sensitivity testing
Cells were seeded at equal amounts (1000 cells for MDA-MB-231, 5000 cells for Hs578T) into 96 well plates and allowed to adhere over night. Then, cells were treated for 24 hours with different doses of Docetaxel (Taxotere®, Sanofi Aventis) respectively Hs578T - 0, 0.005, 0.01, 0.1, 0.5 µg/mL; MDA-MB-231 - 0, 0.01, 0.1, 0.5, 1 µg/mL. CCK-8 solution was added to determine cell viability and proliferation. Absorbance was measured at 450 nm after 4 hours of incubation.

Matrigel invasion assay
The FluoroBlok BioCoat™ Tumour Invasion System (BD Biosciences) was used to assess the invasive potential of cell lines in vitro. Membranes (8 µm FluoroBlok™ PET membranes coated with a uniform layer of BD Matrigel™ matrix) were rehydrated and layers were seeded into the upper chamber insert. To facilitate migration and invasion a serum gradient was used. Cells were allowed to migrate for 22 hours. To quantify migration, cells in the lower chamber were stained with calcein (BD), and the fluorescence was measured at Cytofluor4000 fluorescence reader (MTX Lab Sytems). The amount of migrated cells in the EpCAM transfected lines was compared to the migration rates of their respective empty vector control lines.

Wnt signal activity determination
The Cignal™ TCF/LEF Reporter Kit (SABiosciences, MD) was used to determine Wnt signaling activity. A construct containing a promoter with TCF binding motifs in front of a Firefly luciferase gene was transfected into the cell lines. ß-catenn binding to the TCF motif resulted in the transcription and translation of the luciferase and in the emission of a bioluminescence signal which was detected on a CHAMELEON multi-technology platereader (Hidex). To normalize on transfection efficiency, a CMV promoter driven Renilla luciferase was co-transfected. The luciferase substrates were purchased from Promega (Dual-Luciferase® Reporter (DLR) Assay, E1910) and added automatically by a dual-injector system according to manufacturer’s protocol.

Each cell line was transfected in triplicates with the TCF construct (or negative/positive control construct containing manipulated TCF binding motifs) plus the Renilla control using SUREfect transfection reagent (SABiosciences) and assayed after 48 hours. Bioluminescence signal intensity was determined after normalization on transfection intensity according to following formula: Relative Response Ratio = (SignalReporter-SignalNegative Control)/(SignalPositive Control-SignalNegative Control)

RNA isolation and analysis
Total RNA was isolated by using TRI-reagent (Molecular Research Centre) alone or in combination with the RNeasy MinElute Cleanup Kit (Qiagen) following the manufacturer’s protocol. Concentration and purity of each sample were assessed by absorbance at 260 nm and by the 260/280 nm ratio, respectively. Phenol contaminations were controlled by the 260/270 nm ratio and the concentration was adjusted [19]. The integrity of the RNA was controlled on ethidium bromide-stained agarose-formaldehyde gels. For the samples used for Affimex Chip analysis, RNA quality was determined on an Agilent 2100 Bioanalyzer and by photometric analysis.

Array data analysis
GC Robust Multi-array Average (GCRMA) background adjustment, quantile normalization and median-polish summarization on Affymetrix HGU133 plus 2.0 microarray probe-level data were performed using R 2.7.1 and...
Bioconductor 2.2. (gcrma package: justGCRMA function). The log2 transformed absolute expression values of the treatment sample (EpCAM positive line) were subtracted from the absolute expression values of the control sample (control line) in a gene-wise manner. The resulting log2 transformed relative expression values were ranked according to the level of the values. The top 5% and lowest 5% of the probesets were selected for further analysis. Ingenuity® Pathway Analysis (IPA) software (Ingenuity Systems Inc., Redwood City, CA) was used to inter-connect GO classified categories of the obtained gene set in a context specific manner in order to establish a regulatory network model.

Real-time PCR
Single stranded cDNA was synthesized from 1 μg total RNA using random primers and SuperScript II Reverse Transcriptase (Invitrogen) according manufacturer’s instructions. Real-time analysis was performed using 20 ng single stranded cDNA as well as specific primers and SensiMixPlus containing SYBR® Green (Quantace) on a Rotor Gene 6000 cycler (Corbett Research) using following conditions: 95°C 300 sec; 40 cycles: 95°C 10 sec, 60°C 15 sec, 72°C 10 sec (acquiring on Sybr Green). Primer specificity was checked by melting curve analysis. For each primer pair (Table 1) the PCR amplicon length was verified once by gel electrophoresis and the sequence was analysed. All applications of different cDNA input (n = 3, n = 4) were performed in replicates. The TATA box-binding protein (TBP) was used as endogenous control for normalization [20,21].

Relative expression ratios (R) of target genes were calculated based on the normalized Ct deviation of the EpCAM expressing cells versus the control cells according the mathematical model described by M. Pfaffl: ratio = (2^ΔCt_target(control-EpCAM))/(2^ΔCt_TBP(control-EpCAM)). The Ct value is defined by the cycle at which the threshold is crossed, and ΔCt is the crossing point difference between sample and control [22].

The relative expression software tool REST 2008 (by M. Pfaffl and Corbett Research) was used for statistical analysis. Two groups (controls and EpCAM-positive cells) were compared based on their mean crossing point deviation for significance by a randomization test. The resulting hypothesis test value $p(H1)$ is an indicator of probability that the difference between sample and control group is significant [23,24].

Results
Only few commercially available immortalized human breast cancer cell lines lack EpCAM expression. Hs578T and MDA-MB-231 cells were found to express none or only very little EpCAM mRNA and protein in comparison to established and well characterized breast cancer cell lines such as MCF-7 or SK-BR-3 [25].

Both Hs578T and MDA-MB-231 cells were stably transfected with the EpCAM cDNA containing construct pIRESpuro3_EpCAM. The resulting cell lines were designated Hs578T_EpCAM and MDA-MB-231 EpCAM; the cell lines transfected with the empty control vector pIRESpuro3 were named Hs578Tcontrol and MDA-MB-231 control, respectively. Parental MDA-MB-231 cells were originally derived from a malignant pleural effusion. MDA-MB-231 cells are highly proliferative, invasive in vitro and tumourigenic in nude mice [26]. Hs578T cells were derived from an invasive ductal carcinoma, show low invasive potential and are non-tumourigenic [27,28]. Of note, both cell lines lack E-cadherin expression (due to promoter methylation), which is a common feature among carcinoma cell lines with low or absent EpCAM expression.

EpCAM protein undergoes posttranslational modification
Using SDS-PAGE, EpCAM protein was detectable as a double band in both transfected breast cancer cell lines as well as in MCF-7 control lines under denaturing conditions. After treating cell lysates with the glycosidase PNGase F, both of these EpCAM protein bands were shifted to a lower molecular weight but were still detectable as a doublet, indicating that N-linked glycosilation is not responsible for these two different species of the
EpCAM protein (Figure 1). However, this posttranslational modification might be of functional importance.

Localization of EpCAM determined by confocal microscopy

Confocal microscopy analysis revealed that localization of EpCAM antigen was strongly influenced by cell density in monolayer cell cultures. Highly confluent monolayers showed a predominantly membranous EpCAM staining, while single cells lacking contact with neighbouring cells exhibited much weaker membrane but stronger cytosolic staining. Cell membrane areas at intercellular contact regions stained strongest for EpCAM protein (Figure 2). Since physical interactions of EpCAM and E-cadherin have been proposed [9], confocal microscopy analysis demonstrated that E-cadherin expression at the cell surface was not necessary to recruit EpCAM to the cell membrane in confluent Hs578T EpCAM and MDA-MB-231 EpCAM cells.

The results obtained by confocal microscopy and subcellular fractionation in the transfected EpCAM-positive cell lines were comparable to the endogenously EpCAM-expressing MCF-7 and SK-BR-3 cell lines. For this reason, Hs578T EpCAM and MDA-MB-231 EpCAM cells provided us with an excellent tool to investigate EpCAM-mediated changes on cellular and molecular levels.

Hs578T EpCAM cells show enhanced proliferation and chemosensitivity

EpCAM overexpression has been reported to be associated with a strongly invasive and aggressive tumour phenotype in breast cancer [2,3,17]. However, by using a Matrigel™ coated Boyden chamber assay to compare the in vitro migration/invasion properties of EpCAM-expressing cell lines with their EpCAM-negative empty vector counterparts, no difference in cell migration and invasiveness could be found even by optimizing experimental conditions and prolonging migration time (Figure 3).

To assess cell proliferation, a tetrazolium salt based assay (CCK-8) was performed. The same methodology was used by other groups for the functional analysis of EpCAM function in EpCAM overexpressing HEK293 cancer cells, where EpCAM positive cells exhibited enhanced proliferative and metabolic activity compared to cells with low or no antigen expression [10,11,29]. We additionally compared proliferation by thymidine incorporation with the results of the CCK-8 test, and found the CCK-8 assay to be a reasonable alternative to classical radioactive methods for these cell lines. In our experiments, Hs578T EpCAM cells showed a shortened doubling time as compared to non transfected cells and empty vector controls. This effect has been observed 48 to 72 hours after cell seeding (Figure 4) and was reproduced in different transfected cell clones.
**Figure 2** Microscopical analysis of EpCAM localization. Confocal microscopy of generated and endogenously EpCAM overexpressing cell lines as well as vector controls revealed an increase of EpCAM membraneous staining with cell density. Cells with a cytosolic distribution could be observed in cultures of low confluence. Actin staining with phalloidin revealed slight changes of the actin cytoskeleton formation upon EpCAM expression.
However, this growth-promoting effect could not be detected by comparing MDA-MB-231 EpCAM cells to the respective empty vector control cells, suggesting that the in vitro growth-promoting effect of the EpCAM antigen is not universal but depends on individual cell features. In line with these results, Hs578T<sup>EpCAM</sup> but not MDA-MB-231<sup>EpCAM</sup> cells showed enhanced chemosensitivity to Docetaxel treatment compared to their empty vector counterparts (Figure 5).

Figure 3 Migration/invasion assay. (A) Schematic presentation of an Fluoroblock BioCoat<sup>TM</sup> invasion chamber (BD) composed of a Matrigel<sup>TM</sup> coated transwell insert containing an 8 μm pore size membrane. The Matrigel matrix locks the pores of the membrane and therefore blocks non invasive cells from migrating. Invasive cells can degrade this basement membrane reconstitution and migrate through the pores. (B) Cells have been seeded at high density into the upper chamber. Migration was stimulated by a serum gradient. After 22 hours the amount of migrated cells has been measured with the Cytofluor<sup>4000</sup> fluorescence reader (MTX Lab Systems) from the bottom after cell labelling with calcein (BD), a staining dye for vital cells only. The FluoroBlok<sup>TM</sup> membrane efficiently blocked the transmission of light into the upper chamber, therefore the measured signal derived from migrated cells only. Although all cells migrated efficiently, no difference could be detected by comparing Hs578T<sup>EpCAM</sup> and MDA-MB-231<sup>EpCAM</sup> cells with their respective empty vector control lines (set as 100%).

Figure 4 Cell proliferation assay. In a tetrazolium salt based assay, the proliferative behavior of cell lines with stable EpCAM expression in comparison to empty vector control and untransfected cells has been measured. EpCAM transfected Hs578T cells show enhanced doubling frequency than non-transfected cells and empty vector controls. This effect appears two days after cell seeding. No similar effect could be detected in the stable transfected MDA-MB-231<sup>EpCAM</sup> cell line. Absorbance was normalized to the 24 hours values.
Hs578T\textsuperscript{EpCAM} and MDA-MB-231\textsuperscript{EpCAM} cells show expression changes of Wnt pathway components

In order to investigate changes of the transcriptome on EpCAM gene overexpression in human breast cancer cells, global gene expression analysis using the human genome U133 Plus 2.0 chip (Affymetrix) was performed. Differential gene expression analysis was performed by comparing expression levels from Hs578T\textsuperscript{EpCAM} cells (GEO accession numbers: GSE25743, GSM632607, GSM632608) with the respective empty vector control. The top and lowest 5% of probe sets were used for further analysis by Ingenuity® Pathway Analysis (IPA) software (Ingenuity Systems Inc., Redwood City, CA).

Interestingly, alterations of the Wnt signaling pathway components were found in both EpCAM-transfected cell lines. In more detail, we found the secreted frizzled related protein 1 (SFRP1) - a Wnt ligand competitor and therefore signaling inhibitor - and the HMG box containing transcription factor 7 like 2 (TCF7L2), a Wnt-responsive transcription factor which can also act as repressor of signaling, to be significantly downregulated after EpCAM overexpression. Additionally, the mRNA level for the immunoglobulin transcription factor 2 (ITF-2) was diminished. Only recently, ITF-2 has been identified as a Wnt downstream target [30]. A schematic representation of SFRP1, TCF7L2 and ITF-2 proteins is illustrated in Figure 6.

Changes on the mRNA level detected by expression arrays were confirmed by real-time PCR. TATA box-binding protein (TBP) mRNA expression was not significantly different between control and EpCAM-positive cell lines and served as optimal housekeeping gene. To determine the expression differences of SFRP1, TCF7L2 and ITF-2 in EpCAM positive cells we calculated normalized mean expression levels (Table 2). SFRP1-specific mRNA was consistently downregulated up to 35.8-fold in Hs578T\textsuperscript{EpCAM} and up to 11-fold in MDA-MB-231\textsuperscript{EpCAM} cell lines in comparison to the respective controls. TCF7L2, was downregulated in MDA-MB-231\textsuperscript{EpCAM} cells with a 3.6-fold change while in Hs578T\textsuperscript{EpCAM} cells the TCF7L2 mRNA level was only marginally decreased. In line with these results, protein levels of TCF7L2 were downregulated in MDA-MB-231\textsuperscript{EpCAM} cells (Figure 7). ITF-2 mRNA was downregulated in MDA-MB-231\textsuperscript{EpCAM} cells until the limit of detection (>1500-fold, ΔΔCt value > 10). In Hs578T\textsuperscript{EpCAM} cells, ITF-2 expression was decreased only 3.5 times.

In order to evaluate the obtained regulations in a group-wise comparison we tested the expression differences between EpCAM positive cell lines and the respective controls for significance with a Pair Wise Fixed Reallocation Randomization test by using the REST® relative expression software tool. The perceived differences between EpCAM expressing cells and controls were of statistic significance for the downregulation of SFRP1 in both MDA-MB-231\textsuperscript{EpCAM} and Hs578T\textsuperscript{EpCAM} cell lines and for TCF7L2 and ITF-2 in MDA-MB-231\textsuperscript{EpCAM} cells (p(H1) < 0.05).

An EpCAM-associated downregulation of inhibitory and repressor molecule expression might contribute to the activation or enhancement of Wnt signaling in breast cancer and therefore further corroborate the ongogenic potential of the EpCAM tumour antigen.
Nuclear accumulation of \(\beta\)-catenin in MDA-MB-231\textsuperscript{EpCAM} cells

Fractionation analysis of cell lysates revealed the appearance of EpCAM protein in the soluble cytosolic, the nuclear and the insoluble membraneous fractions in the transfected Hs578TEpCAM and MDA-MB-231\textsuperscript{EpCAM} cell lines (Figure 8). This confirmed the distribution pattern obtained by our group and others in breast cancer-derived as well as in mouse fibroblast cell lines [17,31]. In MDA-MB-231\textsuperscript{control} and in HS578T\textsuperscript{EpCAM} as well as in Hs578T\textsuperscript{control} cells, similar amounts of \(\beta\)-catenin were found in the cytosolic and in the nuclear protein fraction (Figure 8). In MDA-MB-231\textsuperscript{EpCAM} cells, a nuclear accumulation of \(\beta\)-catenin was accompanied consistently by a decrease in \(\beta\)-catenin levels in the cytosol, whereas in Hs578T cells the expression of EpCAM had no significant impact on \(\beta\)-catenin distribution.

As cells were cultivated only to 70-80% density, a majority of the EpCAM protein (detected by C-10 antibody) was present in the cytosol. EpCAM was additionally detected with a C-terminal directed antibody (E144). The presence of EpCAM in the nuclear fraction suggests localization in the perinuclear compartment.
Higher activity of Wnt pathway signaling in MDA-MB-231EpCAM cells

To further confirm these data, the Cignal™ TCF/LEF Reporter Kit (SABiosciences, MD) was used to measure the transcriptional activity of a β-catenin-responsive luciferase reporter. After 48 hours of incubation luciferase activities were evaluated, normalized to the transfection controls and the signaling intensities of EpCAM positive cells were then compared with the values obtained from the corresponding control cell lines (Figure 9).

Wnt activity in Hs578TEpCAM cells was only slightly higher than in Hs578Tcontrol cells (3.7% +/- 1.4%) suggesting that down-regulation of the SFRP1 inhibitor

Table 2 Real-time PCR

|                | TBP     | SRFP1   | TCF7L2   | ITF-2   |
|----------------|---------|---------|----------|---------|
| MDA-MB-231control | 22.11 ± 1.90 | 31.42 ± 2.90 | 19.19 ± 1.45 | 23.47 ± 0.43 |
| MDA-MB-231EpCAM   | 21.81 ± 0.10 | 34.37 ± 1.80 | 20.75 ± 0.27 | 36.26 ± 5.50 |

comparison EpCAM expressing versus control

Regulation Factor

P(H1) 0.000 0.000 0.000

Result DOWN DOWN DOWN

Quantification of differential mRNA expression of the Wnt signaling components SFRP1, TCF7L2 and ITF-2 in the stable EpCAM expressing cells in comparison to the empty vector control lines. (SD = standard deviation, CV = coefficient of variation, REST P(H1) = hypothesis test p value, significant if < 0.05).

Figure 7 TCF7L2 protein expression. Western Blot showing lower amounts of TCF7L2 protein expression in MDA-MB-231EpCAM cells compared to their empty vector counterparts.
alone is not sufficient to achieve a strong activation of Wnt signaling. In contrast, MDA-MB-231 EpCAM cells showed a 19.5% +/- 7.0% higher activity of Wnt pathway signaling compared to empty vector control cells.

Discussion

Our results show the effect of constitutive EpCAM expression in previously EpCAM antigen negative or low expressing parental human breast cancer cell lines Hs578T and MDA-MB-231. Of note, most other commercially available human breast cancer cell lines are characterized by a high level of EpCAM protein expression, thus low or even no EpCAM expression is a rare occurring property. Hs578T cells are used as an example for a low-tumourigenic EpCAM-negative cell line, whereas MDA-MB-231 is used as an example for a more tumourigenic one. Given that only these two breast cancer cell lines could be used for gain-of-function studies, and the resulting phenotypes were not consistent, further investigation will be needed to elucidate a potential in vivo relevance of the presented findings. However, the expression of EpCAM affected cancer-related signaling molecules in both cell lines and seemed to contribute in triggering complex biological processes, which account for an aggressive tumoural behaviour.

In the transfected, constitutively EpCAM-expressing human breast cancer cell lines described here, we observed a reduced expression of Wnt signaling associated components after EpCAM overexpression. We found the downregulation of mRNA levels for SFRP1 in both Hs578T EpCAM and MDA-MB-231 EpCAM cells, and TCF7L2 in MDA-MB-231 EpCAM only. These are two important negative regulators of Wnt signaling [32]. The results on proliferation and Wnt signaling activation between cell lines were not consistent. As such, Wnt signaling was activated significantly in MDA-MB-231 EpCAM but not in Hs578T EpCAM. In contrast to these data, proliferation and chemosensitivity was increased in Hs578T EpCAM but not in MDA-MB-231 EpCAM.

Canonical Wnt signaling hinges on four complexes: the Wnt-frizzled receptor complex in the membrane, the β-catenin destruction complex, the nuclear TCF/LEF/β-catenin transcriptional activator complex and the nuclear TCF/LEF transcriptional repressor complex [33]. SFRP1 acts as a repressor, it binds directly to Wnt ligands and competes with their ability to activate the
frizzled receptors [34]. SFRP1 siRNA treatment in the immortalized, non-transformed human mammary epithelial cell line MCF-10A caused a modest but reproducible increase in luciferase activity in a TCF/LEF activation assay. This activity was further enhanced by addition of Wnt-3A conditioned media [34]. As expression of SFRP1 in breast cancer counteracts Wnt signaling, we proposed that its reduction leads to enhanced Wnt signaling in breast cancer cells.

In MDA-MB-231EpCAM cells mRNAs for both SFRP1 and TCF7L2 were down-regulated compared to the corresponding empty vector line and Wnt signaling was about 20% more active. Of note, Low SFRP1 expression in human breast cancer has been reported by several groups independently [35,36]. Both low SFRP1 and TCF7L2 levels have been described in a subset of breast carcinomas derived from infiltrating ductal carcinoma [32]. However, reduction of SFRP1 expression only, was not sufficient to enhance Wnt signaling in Hs578TEpCAM cells compared to Hs578Tcontrol cells. Probably, the reduction of both SFRP1 and TCF7L2 is required to more potently enhance Wnt signaling in breast cancer cell lines, and a contribution of additional factors cannot be excluded.

The transcription factor TCF7L2 belongs to the TCF/LEF (T-cell factor/lymphoid enhancer factor) protein family and is expressed in many isoformal variants which can coexist in one cell. The short isoforms lack the C-terminal protein binding sites (CtBP) which convert the long variants of TCF7L2 into a repressor [37]. It is therefore conceivable that downregulation of a repressive variant of the TCF7L2 protein in MDA-MB-231EpCAM cells might favour Wnt signaling mediated through other activating TCF/LEF proteins.

Although downregulation of tumour suppressors is a common mechanism during cancer progression, activation of Wnt signaling via the suppression of repressors observed in our breast cancer cells differs from the mechanism discovered for colon cancer, where Wnt pathway activation occurs through loss-of-function mutations of negative pathway components e.g. adenomatous polyposis coli (APC) or gain-of-function mutations in genes activating Wnt signaling (i.e. β-catenin) [38]. Our findings in human breast cancer lines underscore the complexity of Wnt pathway in human cancer [33].

EpCAM and β-catenin proteins were detectable in nuclear lysates of both MDA-MB-231EpCAM and Hs578TEpCAM, but the nuclear staining for EpCAM was negative in confocal microscopy. As the antibody used for microscopical analysis was directed against the extracellular domain, a detection of the shedded cytoplasmic fragment EpICD was not possible. Regarding the results obtained for full-length EpCAM in the fractionation experiment, we suppose that EpCAM has a perinuclear location and cofractionates with nuclear proteins. However, further experiments are required to establish a relationship to the perinuclear space and the endoplasmatic reticulum.

The activation of TCF/LEF by β-catenin is a crucial step in the direct activation of Wnt target genes. In many studies the expression of key target genes such as cyclin D and c-myc is used as a read-out for Wnt signaling activity [10,39]. Upregulation of cyclin D and c-myc has been found to be strongly associated with breast cancer progression [38-40]. However, although expressed at moderate levels in parental cell lines, no significant upregulation of both genes could be detected in Hs578TEpCAM and MDA-MB-231EpCAM cells (data not shown). In
line with our observation, it has been reported that in the absence of distinct TCF7L2 variants the regulation of common TCF/LEF target genes such as cyclin D1, c-myc, MMP7 and c-jun is not detectable in renal cell carcinoma and colorectal cancer cell lines [41]. As each cell type is able to express multiple isoforms of TCF7L2 the combination of long repressors and short activating variants of TCF7L2 adds even more complexity to the signaling events, thus target gene expression can be affected in many different ways [37,42]. Furthermore, the type of trigger and feedback loops within the Wnt pathway have been shown to shape the gene expression pattern and thus the cellular response in normal and transformed epithelial cells [43]. Of note, Western blot analysis of MDA-MB-231 EpCAM cells showed reduced levels of both short and long TCF7L2 variants.

The role of ITF-2, which is strongly down-regulated in MDA-MB-231 EpCAM cells, remains unclear. ITF-2 is a basic-helix-loop-helix (bHLH) transcription factor. Its interaction with the inhibitors of DNA binding Id-1 and Id-2 proteins has been described. In complex with Id proteins, ITF-2 can modulate the mammary epithelial cell phenotype and malignant transformation [44]. Zhai et al showed an interference between ITF-2 and Wnt signaling [45]. Furthermore, the appearance of two isoformal variants, ITF-2A and ITF-2B, differing substantially in domain architecture and function, has been reported [30].

Finally, an increase in invasion and migration after EpCAM overexpression was not observed. Such complex processes are governed by multiple molecular changes. The addition of a single gene might not be sufficient and essential interaction partners might be absent in our analysed cell lines. As such, EpCAM has been described to be a transcriptional target of p53 [46]. A loss of wild-type p53 during breast cancer progression does probably not result in a sole increase of EpCAM expression but also in the regulation of other important target genes.

Conclusions
The effects of constitutive expression of the EpCAM oncogene are dependent on the properties of parental cell lines. Further, the functional alterations observed in vitro were marginal, potentially due to the lack of accessory molecules which might be necessary for EpCAM to exert its full oncogenic potential. In MDA-MB-231 EpCAM cells a nuclear accumulation of β-catenin and a significant upregulation of Wnt reporter assay activity were observed. In Hs578T EpCAM cells, overexpression was accompanied by a considerable decreased expression of the Wnt pathway inhibitor SFRP1, by an increased proliferation rate and enhanced sensitivity to Docetaxel. Certainly, future investigations are warranted to define the functional relationship between increase of EpCAM expression, Wnt signaling and oncogenic features.

List of abbreviations
bHLH: (basic-helix-loop-helix), CAM: (cellular adhesion molecule), c-myc: (v-myc myelocytomatosis viral oncogene homolog); E-FABP: (epidermal fatty acid binding protein); EpCAM: (epithelial cell adhesion molecule), EPLD: (EpCAM intracellular domain); FHL2: (four and a half LIM-only protein 2); SFRP: (secreted frizzled related protein); GAPDH: (glyceraldehyde-3-phosphate dehydrogenase); IRES: (internal ribosome entry site); LEF: (lymphoid enhancer binding factor); SDS-PAGE: (sodium dodecylsulfate polyacrylamide gel electrophoresis), Wnt pathway (wingless-type MMTV integration site family signaling pathway)

Acknowledgements
This work was supported by the doctoral college in Molecular Cell Biology and Oncology Innsbruck (MCBO) and the FWF Grant number P19552.

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Authors’ contributions
JMG established the stably transfected cell lines, carried out migration assays and TCF/LEF reporter assays and contributed substantially in the experimental design. FL and DF carried out Western blots and cell fractionation assays. AM carried out chemosensitivity assay. OAW performed array data analysis. MZ performed PCR analysis. MH carried out immunofluorescence imaging. SK participated in the design of the study. GS and GG conceived the study, and participated in its design and coordination and drafted the manuscript. All authors read and approved the final manuscript.

Competing interests
The authors declare that they have no competing interests.

Received: 5 January 2010 Accepted: 31 January 2011
Published: 31 January 2011

References
1. Tandon AK, Clark GM, Channess GC, McGuire WL: Association of the 323/ A3 surface glycoprotein with tumor characteristics and behavior in human breast cancer. Cancer Res 1990, 50:3317-3321.
2. Gastl G, Spizzo G, Obrist P, Dunser M, Mikuz G: Ep-CAM overexpression in breast cancer as a predictor of survival. Lancet 2000, 356:1981-1982.
3. Spizzo G, Went P, Dimhofer S, Obrist P, Simon R, Spichin H, Maurer R, Metzger U, Von Castelberg B, Bart R, et al: High Ep-CAM Expression is Associated with Poor Prognosis in Node-positive Breast Cancer. Breast Cancer Res Treat 2004, 86:207-213.
4. Schmidt M, Hasenclever D, Schaeffer M, Boehm D, Cotarelo C, Steiner E, Lebrecht A, Siggelkow W, Winkel W, Schiffer-Petry I, et al: Prognostic effect of epithelial cell adhesion molecule overexpression in untreated node-negative breast cancer. Clin Cancer Res 2008, 14:5849-5855.
5. Spizzo G, Obrist P, Ensinger C, Theurel L, Dunser M, Ramoni A, Gunsilius E, Eibl G, Mikuz G, Gastl G: Prognostic significance of Ep-CAM AND Her-2/ neu overexpression in invasive breast cancer. Int J Cancer 2002, 98:883-888.
6. Schmidt M, Scheulen ME, Dittrich C, Obrist P, Marschner N, Dirix L, Schmidt M, Ruttinger D, Schuler M, Reinhardt C, et al: An open-label, randomized phase II study of adecatumumab, a fully human anti-EpCAM antibody, as monotherapy in patients with metastatic breast cancer. Ann Oncol 2009.
28. Thompson EW, Prins FA, Bakker HA, Fleuren GJ, Warnaar SO, Litvinov SV: The structural analysis of adhesions mediated by Ep-CAM. Exp Cell Res 1999, 246:108-121.

29. Litvinov SV, Balzar M, Winter MJ, Bakker HA, Braude-de Bruijn IH, Prins F, Fleuren GJ, Warnaar SO: Epithelial cell adhesion molecule (Ep-CAM) modulates cell-cell interactions mediated by classic cadherins. J Cell Biol 1997, 139:1337-1348.

30. Munz M, Kieu C, Mack B, Schmitz B, Zeidler R, Gires O: The tumour-associated antigen EpCAM upregulates the fatty acid binding protein FABP. Cancer Lett 2005, 225:151-157.

31. Baueerle PA, Gires O: EpCAM (CD326) finding its role in cancer. Br J Cancer 2007.

32. Maetzel D, Denzel S, Mack B, Canis M, Went P, Benk M, Kieu C, Papior P, Baueerle PA, Munz M, et al: Nuclear signalling by tumour-associated antigen EpCAM. Nat Cell Biol 2009.

33. Spizzo G, Chaudry MA, Sales K, Ruf P, Lindhofer H, Winslet MC: EpCAM is overexpressed in breast cancer and is a potential target for breast cancer gene therapy. Cancer Res 2004, 64:5818-5824.

34. Trops M, McLaughlin PH, de Leij LM, Harsmenn MC: Epithelial cell adhesion molecule: more than a carcinoma marker and adhesion molecule. Am J Pathol 2007, 171:386-395.

35. Stuling TM, Amberger A: Exposing contaminating phenol in nucleic acid preparations. Biotechniques 1994, 16:402-404.

36. Bieche I, Olivi M, Champmore MH, Vidaud D, Lidereau R, Vidaud M. Novel approach to quantitative polymerase chain reaction using real-time detection: application to the detection of gene amplification in breast cancer. Int J Cancer 1997, 83:661-666.

37. Bieche I, Onody P, Laurendeau I, Olivi M, Vidaud D, Lidereau R, Vidaud M: Real-time reverse transcription-PCR assay for future management of ERBB2-based clinical applications. Clin Cancer Res 1999, 5:1148-1156.

38. Pfaffl MW: A new mathematical model for relative quantification in real-time RT-PCR. Natl Acad Sci USA 2001, 105:9469-9475.

39. Pfaffl MW, Horgan GW, Dempfle L: Relative expression software tool (REST) for group-wise comparison and statistical analysis of relative expression results in real-time PCR. Nucleic Acids Res 2002, 30:e36.

40. Ditzic A, Mohr A, Mayer K, Burger J, Mayer HH, Pfaffl MW: Effects of mycoplasmic acid (MCA) treatment on expression of Fc receptor (FcIRn) and polymeric immunoglobulin receptor (pIgR) mRNA in adult sheep tissues. Croat Med J 2004, 45:130-135.

41. Prang N, Preitner S, Brischke W, Goster P, Woppeil A, Muller J, Steiger C, Peters M, Baueerle PA, da Silva AJ: Cellular and complement-dependent cytotoxicity of Ep-CAM-specific monoclonal antibody MT201 against breast cancer cell lines. Br J Cancer 2005, 92:342-349.

42. Zhang RD, Fiegl H, Price JE: Relative malignant potential of human breast carcinoma cell lines established from pleural effusions and a brain metastasis. Invasion Metastasis 1991, 11:204-215.

43. Hackett AJ, Smith HS, Springer EL, Owens RB, Nelson-Rees WA, Riggs JL, Gardiner MB: Two syngeneic cell lines from human breast tissue: the aneuploid mammary epithelial (Hs578T) and the diploid myoepithelial (Hs578BM) cell lines. J Natl Cancer Inst 1975, 58:1795-1806.

44. Thompson EW, Paik S, Brunner N, Sommers CL, Zaugnaita C, Clarke R, Shima T, Torri J, Donahue S, Lippman ME, et al: Association of increased basement membrane invasiveness with absence of estrogen receptor and expression of vimentin in human breast cancer cell lines. J Cell Physiol 1992, 150:534-544.

29. Zanna P, Trottola M, Vacca G, Bonasaia V, Palombo B, Guerra E, Rossi C, Lattanzio R, Pantelesi M, Alberti S: Trop-1 are conserved growth stimulatory molecules that mark early stages of tumor progression. Cancer 2007, 110:452-464.

30. Kolligs FT, Nieman MT, Winer L, Hu G, Van Mater D, Feng Y, Smith IM, Wu R, Zha Y, Cho KR, et al: ITF-2, a downstream target of the Wnt/TCF pathway, is activated in human cancers with beta-catenin defects and promotes neoplastic transformation. Cancer Cell 2002, 1:145-155.

31. Balzar M, Bakker HA, Braude-de Bruijn IH, Fleuren GJ, Warnaar SO, Litvinov SV: Cytoplasmic tail regulates the intercellular adhesion function of the epithelial cell adhesion molecule. Mol Cell Biol 1998, 18:6833-4834.

32. Shulewitz M, Soloviev I, Wu T, Koeppen H, Polakis P, Sakamaka C: Repressor roles for TCF-4 and Sfrp1 in Wnt signaling in breast cancer. Oncogene 2006, 25:4361-4369.

33. Hoppler S, Kavanagh CL: Wnt signaling: variety at the core. J Cell Sci 2007, 120:385-393.

34. Uren A, Reichman F, Aneet V, Taylor WG, Murasio K, Battoralo DP, Cumberledge S, Rubin JS: Secreted frizzled-related protein-1 binds directly to Wingless and is a biphasic modulator of Wnt signaling. J Biol Chem 2001, 276:4374-4382.

35. Ugozini F, Charlal-Jaffuret E, Bardou V, Geneix J, Adelia J, Labat-Molecule P, Renault-Lorca L, Longy M, Jacquierne J, Birnbaum D, et al: WNT pathway and mammary carcinogenesis: loss of expression of candidate tumor suppressor gene SFRP1 in most invasive carcinoma except of the medullary type. Oncogene 2001, 20:5810-5817.

36. Wong SC, Lo SF, Lee KC, Yam JW, Chan JK, Wendy Hsiao WL: Expression of frizzled-related protein and Wnt-signaling molecules in invasive human breast tumours. J Pathol 2002, 196:145-153.

37. Shina H, Igarashi M, Breault J, Ribeiro-Filho L, Pookot D, Uraiki S, Tissrima H, Deguchi M, Yamanaka M, Shirai M, et al: The human T-cell factor-4 gene splicing isoforms, Wnt signal pathway, and apoptosis in renal cell carcinoma. Clin Cancer Res 2003, 9:2121-2132.

38. Hattel S, Rowlands T, Herrimath M, Cowyn P: Beta-catenin and Tcf5 in mammmary development and cancer. J Mammary Gland Biol Neoplasia 2003, 8:145-158.

39. Rowlands TM, Pechkenina IV, Hattel S, Cowyn P: Beta-catenin and cyclin D1: connecting development to breast cancer. Cell Cycle 2004, 3:145-148.

40. Yang SZ, Kohno N, Yokoyama A, Kondo K, Hamada H, Hiwada K: Decreased E-cadherin augments beta-catenin nuclear localization: studies in breast cancer cell lines. Int J Oncol 2001, 18:541-548.

41. Tang W, Dodge M, Gundapaneni D, Michnoff C, Roth M, Lum L: A genome-wide RNAi screen for Wnt/beta-catenin pathway component identifies unexpected roles for TCF transcription factors in cancer. Proc Natl Acad Sci USA 2008, 105:9697-9702.

42. van Noort M, Clevers H: TCF transcription factors, mediators of Wnt-signaling in development and cancer. Dev Biol 2002, 244:1-8.

43. Sankpal NV, Willman MW, Fleming TP, Layfield JD, Gillanders WE: Beta-catenin and Wnt target genes of the Wnt/β-catenin pathway and statistical insights into Wnt target promoter regulation. FEBS J 2005, 272:1600-1615.

44. Parnell S, Lin CQ, Murata K, Itahana Y, Singh J, Krotkola A, Campisi J, Desprez PY: Id-1, IFT-2, and Id-2 comprise a network of helix-loop-helix proteins that regulate mammary epithelial cell proliferation, differentiation, and apoptosis. J Biol Chem 2001, 276:39213-39219.

45. Zha Y, Wu R, Schwartz DR, Dannah D, Reed H, Kolligs FT, Nieman MT, Fearon ER, Cho KR: Role of beta-catenin/TFI-cell regulator-gene regulated genes in ovarian endometrioid adenocarcinomas. Am J Pathol 2002, 160:1229-1238.

46. Sankpal NV, Willman MW, Fleming TP, Layfield JD, Gillanders WE: Transcriptional repression of epithelial cell adhesion molecule contributes to p53 control of breast cancer invasion. Cancer Res 2009, 69:753-757.

Pre-publication history
The pre-publication history for this paper can be accessed here: http://www.biomedcentral.com/1471-2407/11/45/Suppub