Investigations on the Usefulness of CEACAMs as Potential Imaging Targets for Molecular Imaging Purposes

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Investigations on the Usefulness of CEACAMs as Potential Imaging Targets for Molecular Imaging Purposes

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Abstract

Members of the carcinoembryonic antigen cell adhesion molecules (CEACAMs) family are the prototype of tumour markers. Classically they are used as serum markers, however, CEACAMs could serve as targets for molecular imaging as well. In order to test the anti CEACAM monoclonal antibody T84.1 for imaging purposes, CEACAM expression was analysed using this antibody. Twelve human cancer cell lines from different entities were screened for their CEACAM expression using qPCR, Western Blot and FACS analysis. In addition, CEACAM expression was analyzed in primary tumour xenografts of these cells. Nine of 12 tumour cell lines expressed CEACAM mRNA and protein when grown in vitro. Pancreatic and colon cancer cell lines showed the highest expression levels with good correlation of mRNA and protein level. However, when grown in vivo, the CEACAM expression was generally downregulated except for the melanoma cell lines. As the CEACAM expression showed pronounced expression in FemX-1 primary tumours, this model system was used for further experiments. As the accessibility of the antibody after i.v. application is critical for its use in molecular imaging, the binding of the T84.1 monoclonal antibody was assessed after i.v. injection into SCID mice harbouring a FemX-1 primary tumour. When applied i.v., the CEACAM specific T84.1 antibody bound to tumour cells in the vicinity of blood vessels. This binding pattern was particularly pronounced in the periphery of the tumour xenograft, however, some antibody binding was also observed in the central areas of the tumours. Still, a general penetration of the tumour by i.v. application of the anti CEACAM antibody could not be achieved despite homogenous CEACAM expression of all melanoma cells when analysed in tissue sections. This lack of penetration is probably due to the increased interstitial fluid pressure in tumours caused by the absence of functional lymphatic vessels.

Introduction

Members of the carcinoembryonic antigen (CEACAM) family are transmembrane glycoproteins belonging to the immunoglobulin superfamily, which are involved in a variety of biological processes [1,2]. These include regulation of cell growth, differentiation, immune response, cellular recognition and cell adhesion [3,4,5,6]. In addition to their normal function, expression of several members of the CEACAM family was found to be upregulated in colorectal and lung cancer as well as in melanoma [7,8,9]. Due to their up-regulation in these entities, members of the CEACAM family have served as valuable clinical markers both in tissue sections and patients’ sera [10,11]. In particular, the classical serum marker CEACAM5 (CEA) is highly expressed in cancers including colorectal, gastric, pancreatic, and small cell lung cancer [12,13,14]. Because of their high expression level in colon cancer, serum CEA levels are routinely used to monitor the recurrence of colonic adenocarcinoma after surgery and some of the antibodies have been used in patient studies [15,16,17].

However, marker analysis of serum samples does not disclose the site of CEA production and therefore the site of the (primary) tumour remains unresolved by serum analysis. To localise tumours, endoscopic as well as non-invasive imaging techniques like MRI are used, which, however, lack information about the specific proteins secreted by the tumour including CEA. To obtain information about the specific molecular composition of tumours, MRI techniques have to be combined with antibody based technologies resulting in molecular imaging techniques [18].

To discover the capabilities and limitations of molecular imaging we developed murine xenograft models for in vivo detection of CEACAMs. CEACAMs were chosen for targeting...
as they are often highly expressed in a variety of malignancies (see above). In order to broaden the specificity of our molecular probe, we used the T84.1 monoclonal antibody which is capable of recognizing several members of the CEACAM family including CEACAM 1, 5 and 6 [19].

This contribution describes the expression of T84.1 immunoreactivity in 12 different human cancer cell lines for CEACAM expression in vitro and when grown in immunodeficient mice in vivo as primary tumour in order to establish a xenograft model for CEACAM detection. With one of these models we additionally investigated the accessibility of CEACAMs to antibodies in the primary tumour after i.v. application of the anti pan-CEACAM antibody T84.1.

Materials and Methods

Cell lines

The human prostate cancer cell lines LNCAP [20] and PC3 [21] (both established from metastatic adenocarcinomas) were obtained from the German Collection of Microorganisms and Cell Culture (DSMZ, Germany). The human breast cancer cell lines T47D [22] and MCF7 [22] (both established from pleural effusions) were obtained from European Cell Culture Collection (Porton Down, Wiltshire, UK). The human melanoma cell lines MEWO [23] and FemX-1 [24] (both established from metastatic melanoma lymph nodes) were kindly provided by the Klinik für Dermatologie, Universitätsklinikum Hamburg-Eppendorf, Germany. The human colon cancer cell line HT29 [25] (established from a primary carcinoma of the colon) was obtained from Cell Lines Service (Germany). The human colon cancer cell lines Caco2 and SW480 [22] (both established from primary adenocarcinomas of the colon) were obtained from European Cell Culture Collection (Porton Down, Wiltshire, UK). The human small cell lung cancer cell line OH-1 [26] (established from pleural effusion) was kindly provided by Prof. Uwe Zangemeister-Wittke, University of Bonn, Department of Pharmacology. The human pancreatic cancer cell line 5061 [27] (established from an advanced pancreatic adenocarcinoma) was kindly provided by the Klinik und Poliklinik für Allgemein-, Viszeral- und Thoraxchirurgie, Universitätsklinikum Hamburg-Eppendorf, Germany. The human pancreatic cell line 5072 (established from a advanced pancreatic adenocarcinoma from a 71-year-old Caucasian woman) was kindly provided by the Klinik und Poliklinik für Allgemein-, Viszeral- und Thoraxchirurgie, Universitätsklinikum Hamburg-Eppendorf, Germany. Histopathological examination of the surgical specimen confirmed a low-differentiated adenocarcinoma of the pancreas, which was staged pT3, pN2, G3, M0, R0. Written informed consent of the patient for the removal of tissue samples for investigational purposes was obtained prior to surgery. The study was approved by the ethical committee of the Medical Council of Hamburg (Arztekammer), Germany.

The cell lines LNCAP, PC3, T47D, MCF7, MEWO, FemX-1, HT29, Caco2, SW480, OH-1 were cultured in vitro under standard cell culture conditions (37°C, 100% relative humidity, 5% CO2) in RPMI medium (Gibco/Life Technologies, Paisley, Scotland) supplemented with 10% heat inactivated fetal bovine serum (FBS, Gibco), 2 mM L-glutamine (Gibco), 100 U/ml penicillin and 100 μg/ml streptomycin (Gibco). The cell lines 5061 and 5072 were cultured in complete (TUM) RPMI 1640 medium with Glutamax (Invitrogen, NY, USA) supplemented with 10% of fetal calf serum (FCS), 200 IU/ml of penicillin-streptomycin, 0.1 mg/ml gentamycin (Biochrom AG, Berlin, Germany), 50 μmol/ml of human transferrin (Sigma-Aldrich, Steinheim, Germany), 0.01 μg/ml of bovine insulin (Sigma-Aldrich, Steinheim, Germany), 0.01 μg/ml of recombinant human epidermal growth factor (Pepro Tech, London, UK), and 0.01 μg/ml of human basic fibroblast growth factor (Pepro Tech, London, UK). Before reaching confluence, cells were routinely harvested for passaging using 0.05% trypsin-0.02% EDTA (Gibco).

Real-time PCR

To quantify CEACAM mRNA amount in relation to GAPDH mRNA amount of the human tumour cell lines, real-time PCR was carried out. In brief, total RNA from tumour cells was isolated using RNeasy Mini Kit (Qiagen, Hilden, Germany) according to the manufacturer’s instructions. The RNA was eluted in 50 μl RNase free water. The RNA-concentration was measured and the quality was checked on a NanoDrop® ND-1000 Spectrophotometer (Peqlab, Erlangen, Germany). The cDNA synthesis was performed in a Biometra thermal cycler (Biometra, Göttingen, Germany) in a total volume of 20 μl for each sample and followed the manufacturer’s instruction for the First Strand Transcriptor cDNA Synthesis Kit (Roche Diagnostics, Mannheim, Germany). Two parallel cDNA approaches were used for reverse transcription separately, with anchored-oligo(dT)18 and random hexamer primer. Two μl of total RNA were used for each cDNA approach and were pooled afterwards. Real-time polymerase chain reaction was performed in a 96 well format with the LightCycler® 480 System (Roche Diagnostics GmbH, Mannheim, Germany). For the real-time PCR, the LightCycler Fast Start DNA MasterPLUS SYBRGreen 1 Kit (Roche Diagnostics GmbH, Mannheim, Germany) was used. Two μl of cDNA was used as a template for the PCR reaction and incubated in a total reaction volume of 10 μl, containing 1×SYBR Green 1 Master mix including Taq DNA polymerase, Taq PCR buffer, a dNTP mixture and 1 mmol/l MgCl2, 10 pmol specific CEACAM or GAPDH primers. Forward CEACAM primer (TGTTGAAAGACAGGACAC), reverse CEACAM primer (CAGCCTGGGACTCACCGG), forward GAPDH primer (AAA TTGGACCGCAGCCTCCC), and reverse GAPDH primer (CCAGGCCGCAATACGACAAAT) were synthesized by MWG-BIOTECH AG (Ebersberg, Germany). The PCR conditions were initially 5 min 95°C, followed by 40 cycles of 10 s 95°C, 10 s 62°C (CEACAM) or 72°C (GAPDH) and 20 s 72°C, respectively. The primer pair detects in parallel CEACAM 1, 5 and 6.

Protein extraction and Western blotting

Total protein extracts from cell lines were obtained by lysing the cells in cold radioimmunoprecipitation assay (RIPA) buffer (50 mM Tris, 2 mM EDTA, 1% NP-40, 0.1% SDS, and 150 mM NaCl) in the presence of protein inhibitor cocktail set I (Calbiochem, La Jolla, USA). After centrifugation to remove cell debris, protein concentrations of the supernatants were measured using the BCA method [28]. 40 μg protein per lane were resuspended in loading buffer (0.5 M Tris-HCl, pH 6.8, glycerol, 10% SDS, 0.5% bromophen blue, mercaptoethanol) and then subjected to sodium dodecyl sulfate–polyacrylamide gel electrophoresis (8% gels). Proteins were subsequently blotted onto a nitrocellulose membrane (Hybond-ECL®, Amersham Biosciences, Freiburg, Germany) following conventional protocols. Finally, blots were blocked in 4% bovine serum albumin/Tris-buffered saline–0.1% Tween 20 (TBS-T) for 30 min at room temperature. Membranes were incubated with primary antibody T84.1 (1 μg/ml) and anti-beta-Actin (0.5 μg/ml, Abcam, Cambridge, UK) over night at 4°C, washed with TBS-T and incubated with a 1:200 diluted polyclonal goat-anti-mouse antibody (DakoCytomation, Carpinteria, USA) conjugated with horseradish peroxidase for 60 min at room temperature. The
bound immune complexes were visualised using ECL Western Blotting Substrate (Pierce, Rockford, USA) and the ChemiDoc XRS System (Bio-Rad, Munich, Germany).

Flow Cytometry
Cultured cells were detached with Cell Dissociation Buffer ( Gibco™, Carlsbad, USA) and incubated on ice for 30 min with T84.1 primary antibody at a concentration of 1 μg/ml. The corresponding marine isotype control was IgG1 (DakoCytomation, Carpenteria, USA). The cells were washed and the primary antibody was detected with allopheocyanin-conjugated goat-anti-mouse antibody (Becton Dickinson Biosciences, Heidelberg, Germany). Flow cytometry was performed using a FACs CALIBUR flow cytometer (Becton Dickinson, Heidelberg, Germany). Mouse antobody was detected with allophycocyanin-conjugated goat-anti-mouse antibody (Becton Dickinson Biosciences, Heidelberg, Germany). The cells were washed and the primary antibody was detected with allopheocyanin-conjugated goat-anti-mouse antibody (Becton Dickinson Biosciences, Heidelberg, Germany). Flow cytometry was performed using a FACs CALIBUR flow cytometer (Becton Dickinson, Heidelberg, Germany). Data were analyzed using Win MDI 2.9 software.

Immunocytochemistry
For immunocytochemistry, cells cultured in eight-well chamber slides (Becton Dickinson, Heidelberg, Germany) were washed with PBS, fixed in ice-cold acetone for 2 min, air dried, and blocked with 10% rabbit sera in blocking reagent (TBS). Slides then were incubated with T84.1 or IgG1 control antibody at a dilution of 0.6 μg/ml over night (4 °C) followed by a biotinylated rabbit anti mouse antibody (DakoCytomation) at a dilution of 1:200 for 30 min. After careful washes in TBS, an incubation with a avidin-alkaline phosphatase complex (ABC kit, Vectastain, Vector, Burlingame, CA) for 30 min followed and thereafter, additional washes in TBS were performed. Alkaline phosphatase activity was visualised using Naphthol-AS-bi-phosphate as a substrate and New Fuchsin was used for simultaneous coupling. Slides were counterstained with Mayer’s hemalum diluted 1:1 in distilled water for ten seconds, blued under running tap water and mounted with Aquatex® (Merck KGaA, Darmstadt, Germany). The intensity (plus signs) and extent (percentage) of the positive areas of five histological sections were determined by visual inspection of three independent observers. The plus sign indicates the intensity of the staining. It ranges from “+++” to “−”. The percentage indicates how many cells are labelled.

Immunohistochemistry of tumour cell lines xenotransplanted in SCID mice
Sections of all primary tumors were drawn from our in house tumor bank. They had all been fixed with 4% formalin and embedded in paraffin wax according to standard procedures. For immunohistochemistry, 5 μm thick sections were cut, dewaxed and microwaved in 10 mM citrate buffer (pH 6.0) at 500 W five times for 2 min and then cooled for 20 min. After washing, non-specific binding was blocked by incubating the sections in 10% normal rabbit serum (Dakocytomation, Hamburg, Germany) for 30 min at room temperature. The following steps were as described above.

Ethics Statement
The methodology for carrying out the experiment was consistent with the UKCCCR guidelines for the welfare of animals in cancer research [29]. The experiment was supervised by the institutional animal welfare officer and approved by the local licensing authority (Behörde für Soziales, Familie, Gesundheit und Verbraucherschutz; Amt für Gesundheit und Verbraucherschutz, Hamburg, Germany, project no. 92/09).

In vivo experiments
To investigate in vivo CEACAM binding sites, specific pathogen-free BALB/c SCID (scid/scid) mice were used. The mice were 8–16 weeks old and weighed 20–30 g at the beginning of our experiments. They were housed in filter top cages and provided with sterile water and food ad libitum. For injection, FemX-1 melanoma cells were harvested by trypsinisation and viable cells (5×10⁶) were suspended in 1 ml of cell culture medium. An aliquot of 200 μl of this suspension was injected subcutaneously between the scapulae of each SCID CD mouse. Twelve mice bearing melanomas were included in the experiment when the tumour had reached maximal growth or started to ulcerate.

Iodination of antibodies was done by Iodobeads (Pierce, Rockford, USA). Specifically, 5 μl sodium [¹²⁵I]Iodine (3.7 MBq/μl, Amersharm, UK) was added to 1 mg T84.1 or control IgG1 in 1 ml PBS in the presence of one Iodobead and the reaction was left for 15 min. Free [¹²⁵I]Iodine was removed by exclusion chromatography through a PD-10 column (GE Healthcare, Buckinghamshire, UK). The protein concentration was determined by the BCA method and the specific activity of the [¹²⁵I]-T84.1 and [¹²⁵I]-IgG1 per μg protein was measured by gamma counting (Perkin Elmer, Waltham, USA).

Ten μg of [¹²⁵I]-labelled antibodies were injected via the tail vein into FemX-1 melanoma bearing mice. Six mice received [¹²⁵I]-IgG1 and six mice [¹²⁵I]-T84.1 antibody. After 0.5, 2.5, 6, 17.5, and 22.5 hours blood samples were taken by tail vein to measure antibody blood half life. Afterwards mice were sacrificed, intracardially perfused with NaCl and tumour and organs were removed and weighed. Radioactivity of organs, tumours and blood were determined in the gamma counter. Statistical analyses (Two-way ANOVA) were performed using GraphPad Prism 5 software (GraphPad software, USA).

Evans Blue was injected i.v. at a dose of 25 μl (50 mg/ml PBS) per FemX-1 s.c. bearing mouse 4 hours before scarification. Anaesthetised mice were intracardially perfused with 1% BSA in NaCl to remove free dye and afterwards fixed with 4% PFA. Tumours were removed and embedded in 5% agarose gel, cut in 200 μm thick slices with a vibratome (VT1000E, Zeiss, Wetzlar, Germany) and scanned with 2400 dpi on a scanner (Epson, Long Beach, USA).

Results

CEACAM expression in vitro
The CEACAM expression levels of twelve human cancer cell lines derived from different tumour entities are summarized in figure 1 and table 1.

The highest expression of CEACAM mRNA was detected in the pancreatic tumour cell lines 5061 and 5072, the colon cancer cell lines HT29 and Caco2, and the prostate cancer cell line LNCAP. Low levels could be found in the breast cancer cell lines T47D and MCF7 and in the melanoma cell line FemX-1. Almost no CEACAM expression could be detected in the lung cancer cell line OH-1, the melanoma cell line MEWO, and the colon cancer line SW480.

Presence of the CEACAM proteins was analyzed by Western Blot of cell lysates (figure 2, table 1). All cell lines showed distinct protein bands except for the cell line T47D, which showed only a very faint band. Several bands were detected in the blots, presumably because of the different glycosylation isoforms that exists of the CEACAM proteins. No obvious correlation between mRNA and protein levels was observed.

The amount of membrane bound CEACAM proteins were furthermore determined by FACS analysis using mAb T84.1 (figure 3). As summarized in table 1, almost all cell lines showed the same staining pattern, as seen in the immunocytochemistry.
To visualize the CEACAM expression pattern under cell culture conditions we stained the cells, grown on chamber slides, with the panCEACAM T84.1. As shown in figure 4, the pancreatic (5061, 5072), colon (Caco2, HT29, SW480), and melanoma (FemX-1, MEWO) cancer cell lines clearly bound mAb T84.1. Lower or no binding could be detected by prostate (LNCAP, PC3) and breast (MCF7, T47D) cancer cell lines.

CEACAM expression in vivo
Only FemX-1 and LNCAP primary xenograft tumours showed strong mAb T84.1 binding by more than 50% of all cells (figure 4 and table 1), while all other primary tumours did not show substantial T84.1 binding.

CEACAM binding detection in vivo
To analyse whether the CEACAM binding sites detected in tissue sections were also accessible in vivo, [125I]-labelled T84.1 mAb and non-specific [125I]-labelled control IgG were injected into the tail vein of FemX-1 melanoma bearing mice. A significant two-fold higher enrichment of the [125I]-T84.1 mAb in the FemX-1 melanoma was detected compared to the [125I]-IgG1 control (figure 5) (Two-way ANOVA, P<0.001). There was no significant difference concerning blood half life or tissue distribution between specific [125I]-T84.1 mAb and control [125I]-IgG (figure 6).

In a parallel experiment the distribution of the injected mAbs 24 hours after application were analysed in histological sections regarding tissue distribution of the antibody (figure 7). No detectable levels of antibody could be found in control animals injected with control IgG1 mAb. In contrast, T84.1 binding could be detected at the margin of the tumour and in the central area of the FemX-1 tumour around blood vessels.

To study the mechanism of T84.1 penetration of the tumour tissue we investigated the tumour vessel permeability with an alternative technique. We injected the albumin-binding dye Evans Blue i.v. in FemX-1 melanoma bearing mice and investigated the dye distribution in vibratome sections of the tumour. Evans Blue-Albumin complex was present only around blood vessels, at the margin of the tumour, and in the transition zone between vital tumour tissue and central necrosis and thus was similar to the T84.1 binding (figure 8).

Discussion
Identifying malignant tumours is one of the most important aims of molecular imaging. In order to bring molecular imaging techniques into clinical application, the technical approach has to be validated in suitable animal models first [30]. We therefore systematically investigated the CEACAM expression of twelve human cancer cell lines and their primary tumour xenografts in immunodeficient mice for their suitability to detect the tumour using an anti CEACAM antibody as a pre-requisite for imaging studies.

As human cancer cell lines often serve as the first choice for detecting human specific antigens, we analyzed CEACAM mRNA expression of human malignant cells. The mRNA of the malignant cell lines 5061, 5072 (pancreas), PC3, LNCAP (prostate), FemX-1, MEWO (melanoma), MCF7, T47D (breast), HT29 (colon) and OH-1 (small cell lung cancer) were relatively quantified by real time PCR, using GAPDH for normalization. Cell lines 5061, 5072, LNCAP and HT29 showed high expression levels of CEACAM mRNA.

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Table 1. Summary of in vitro and in vivo CEACAM expression of all cell lines by different methods.

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<tr>
<th>cell culture</th>
<th>QPCR</th>
<th>WB</th>
<th>FACS positive</th>
<th>immunocytochemistry</th>
<th>xenografted tumor</th>
<th>immunohistochemistry</th>
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<tr>
<td>S061</td>
<td>+++</td>
<td>+</td>
<td>99%</td>
<td>+++ (95%)</td>
<td>+/+ (45%)</td>
<td></td>
</tr>
<tr>
<td>S072</td>
<td>++</td>
<td>+</td>
<td>77%</td>
<td>+++ (90%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>LNCAP</td>
<td>++</td>
<td>+</td>
<td>93%</td>
<td>++ (100%)</td>
<td>+/+ (60%)</td>
<td></td>
</tr>
<tr>
<td>PC3</td>
<td>+</td>
<td>+</td>
<td>23%</td>
<td>++ (10%)</td>
<td>(+) (20%)</td>
<td></td>
</tr>
<tr>
<td>T47D</td>
<td>+</td>
<td>(+)</td>
<td>5%</td>
<td>+ (1%)</td>
<td>+/- (20%)</td>
<td></td>
</tr>
<tr>
<td>MCF7</td>
<td>+</td>
<td>(+)</td>
<td>23%</td>
<td>+++ (5%)</td>
<td>++/++ (20%)</td>
<td></td>
</tr>
<tr>
<td>MEWO</td>
<td>(+)</td>
<td>+</td>
<td>74%</td>
<td>+++ (85%)</td>
<td>+</td>
<td></td>
</tr>
<tr>
<td>FemX-1</td>
<td>+</td>
<td>+</td>
<td>95%</td>
<td>+++ (100%)</td>
<td>++/+ (90%)</td>
<td></td>
</tr>
<tr>
<td>HT29</td>
<td>++</td>
<td>+</td>
<td>86%</td>
<td>+++ (90%)</td>
<td>++ (45%)</td>
<td></td>
</tr>
<tr>
<td>Caco2</td>
<td>+</td>
<td>+</td>
<td>16%</td>
<td>+++ (90%)</td>
<td>+</td>
<td></td>
</tr>
<tr>
<td>SW480</td>
<td>–</td>
<td>+</td>
<td>31%</td>
<td>+ (5%)</td>
<td>(+) (30%)</td>
<td></td>
</tr>
<tr>
<td>OH-1</td>
<td>(+)</td>
<td>+</td>
<td>80%</td>
<td>++ (80%)</td>
<td>(+) (100%)</td>
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Note that the highest CEACAM expression in vitro and in vivo showed the melanoma cell line FemX-1. The intensity (plus signs) and extent (percent) of the positive areas of 5 histological sections were determined by visual inspection of 3 independent observers.
doi:10.1371/journal.pone.0028030.t001
expression in a panel of human malignant cell lines. This was followed by the detection of their CEACAM protein expression in Western Blots. We found that pancreatic and colon (except of SW480) cancer cell lines have the highest expression levels of CEACAM with good correlation between mRNA amount and protein level in Western Blots. Similarly high protein levels were detected in melanoma cells, but their mRNA level was generally lower than that observed in the pancreatic and colon cancer cell lines.

This discrepancy between mRNA and protein expression levels has already been observed e.g. in *Saccharomyces cerevisiae* for the PUP2, and by mammals for circadian Period2 gene [31,32]. RNA stability and/or translational efficiency between the cancer cell lines and different CEACAM family members could be a reason for the finding, that low mRNA level were associated with high protein levels.

A further discrepancy in CEACAM expression was observed between *in vitro* and *in vivo* CEACAM expression (figure 4). Except

![Figure 2. Western blot analysis of the CEACAM protein expression pattern. CEACAM family members as detected by pan CEACAM specific antibody T84.1 were present in all cancer cell lines except T47D. Beta-actin was used as a loading control. doi:10.1371/journal.pone.0028030.g002](image)

![Figure 3. CEACAM cell surface presence of tumour cells. CEACAM could positively be detected by FACS-analysis with mAb T84.1 by all cancer cell lines, except of T47D, MCF7 and PC3. Isotype controls are shown as dotted lines. doi:10.1371/journal.pone.0028030.g003](image)
melanomas, all malignant cell lines showed a downregulation of CEACAM expression in vivo as compared to in vitro. CEACAM family members are mainly expressed by epithelial and endothelial cells at the free surface of their apical pole [33,34]. In contrast to the three dimensional growth in vivo where tumour cells form a 3D conglomerate of cells with only a few free surfaces, almost all cultured cells have exposed free surfaces and therefore almost all cells display an apical cell pole enabling them to express CEACAM at this interface to the cell culture medium. This different growth behaviour results in a down regulation of CEACAM in the primary tumours compared to the cell culture growth.

The observation that the melanoma cell line showed the highest CEACAM expression level might not be so unexpected, as periodic acid-Schiff (PAS) reaction positive loops and networks indicating micro vascular channels within melanoma have been described [35]. As these microvascular channels are lined by the melanoma cells themselves and are not covered by endothelial cells, this apical surface might provide additional space for CEACAM expression. A further reason for the difference in expression can be found in tumour-stroma interactions. Cancer cells in a tumour xenograft are exposed to mouse extracellular matrix and mouse cells like blood vessel endothelia, lymphocytes and macrophages and also to mouse hormones. All these tumor-stroma interactions can potentially alter the expression of genes in the human tumour cell in xenografts [36]. The few regions of higher CEACAM expression in vivo in the xenograft of cell line T47D could be explained by the fact, that hypoxia could upregulate CEACAM expression via HIF1α signaling. A HIFα response element is located in the promoter region of CEACAM [37].

Another factor to be investigated is the accessibility of the monoclonal antibody to the CEACAM binding sites after i.v.

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**Figure 4. CEACAM protein expression pattern in vivo and in vitro.** Cell lines LNCAP and FemX-1 showed in vitro and in vivo strong binding of mAb T84.1 by all cells. In contrast, cell lines 5061, MEWO, OH-1 (insert with higher magnification) HT29, Caco2, and particularly 5072 showed strong mAb T84.1 in vitro binding, but little or no binding in vivo. (red = T84.1 binding).

doi:10.1371/journal.pone.0028030.g004
application of the T84.1 antibody. Using labelled lectins as a probe, we have previously shown that vast differences in lectin binding site accessibility existed between the binding of the lectins to tissue sections and to the same tissue after i.v. injection of the same lectins [38]. We therefore investigated the presence of CEACAM binding sites in vivo using the CEACAM specific T84.1 mAb on FEMX-1 cells after its i.v. application. FEMX-1 cells were chosen as they readily grow in SCID mice and additionally show robust CEACAM expression in vitro and in vivo (figure 4). We could indeed show that the i.v. applied antibody reaches the target, but to a moderate extent only as its presence was limited to the direct vicinity of blood vessels (figure 5 and 7). Furthermore, melanoma is a particularly well suited malignancy to be investigated as CEACAM expression is positively correlated with metastasis formation [9]. Therefore more malignant cells express CEACAM-1 in melanoma, while in other tumour entities such as breast or prostate cancer CEACAM-1 expression is downregulated during malignant progression [39,40].

A known phenomenon of tumour xenografts is the uneven distribution of blood vessels within different areas of the tumour [41]. In our FemX-1 melanoma model the vascular density was more intense in the periphery than in the centre of the tumour xenograft (figure 8). The access of the T84.1 antibodies was limited to areas around blood vessels, resembling Kroghs’ cylinder which shows the limits of oxygen diffusion in tissues [42]. Even the Evans Blue-Albumin complex with a lower mass weight of 67 kDa could only be found in the same area, as the injected T84.1 antibody (figure 8). All transport processes beyond this cylinder are not based on diffusion but on convection. Convection, however, is severely altered in tumours because of the absence of functional lymphatic vessels within the tumour. This leads to an accumulation of interstitial fluid within tumours resulting in a high interstitial fluid pressure of tumours which could be the pathophysiological mechanism behind this finding of a limited access of the antibody to the melanoma cells, despite the presence of microvascular channels in melanomas [43,44].

The process of diffusion allows only the penetration of small molecules into the tumour, the size of the whole antibody molecule is obviously far too big to target cells behind the endothelial barrier as the target molecule CEACAM is clearly expressed in the tumour as shown in tissue sections. Therefore, the access of i.v. administrated anti CEACAM antibodies have not found acceptance in routine clinical practice [16,17]. In order to target a wider range of CEACAM positive tumour cells, smaller target-specific molecules like DARPinS or nanobodies against CEACAM will be tested to see...
if they could reach the in vivo CEACAM binding sites to a greater extent. However, we could show that the target CEACAM could be studied with our melanoma FEMX-1 model and used for molecular imaging.

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References