CA19.9 antigen circulating in the serum of colon cancer patients: Where is it from?

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A B S T R A C T

CA19.9 antigen is a glycoprotein present in human serum and found elevated in various diseases. It is intensively studied since long time as a potential marker for managing cancers of the gastrointestinal tract, but its reliability is widely accepted only for pancreatic cancers. Here, we focused on the tetrasaccharide epitope (NeuAcα2-3Galβ1-3[Fucα1-4GlcNAc]) sialyl-Lewis a studying the biosynthesis, expression, and secretion in colon cancers and related cancer cell lines. We found that the β1,3 galactosyltransferase βGal-T5, responsible for sialyl-Lewis a synthesis, is dramatically reduced in colon adenocarcinomas, in terms of both transcript and enzyme activity levels. Moreover, no or very faint antigen is detectable in colon cancer homogenates, by dot-blot or enzyme immunoassay, while it is commonly evident in sera from different patients. In cancer cell lines synthesizing CA19.9, the amount of antigen secreted is proportional to that expressed on the cell surface, and depends on appreciable levels of β3Gal-T5, which appear much higher than those measured in colon cancer specimens. Since colon cancers appear unable to synthesize relevant amount of CA19.9, we suggest that the antigen circulating in the serum of colon cancer patients may have a different and more complex origin than expected so far.

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1. Introduction

Sialyl-Lewis a is a the tetrasaccharide epitope (NeuAcα2-3Galβ1-3[Fucα1-4]GlcNAc) characterizing the CA19.9 antigen (Magnani et al., 1983; Yue et al., 2011a) present in human serum and found elevated in various diseases (Mann et al., 2000) including cancers of the digestive tract, as pancreas (Ballehanina and Chamberlain, 2012), bile ducts (Kikkawa et al., 2012), stomach (Kim et al., 2011), and colon (Yamashita and Watanabe, 2009). At present, CA19.9 is recommended by medical societies and study groups for managing cancers of the pancreas but not of the colon (Duffy et al., 2003, 2007, 2010; Locker et al., 2006), but is still widely used in clinical practice, trials, and studies concerning colorectal cancer diagnosis, prognosis, surveillance, and response to therapy (de Haas et al., 2010; Bystrom et al., 2012; Lin et al., 2012). The existence of such controversial issues and the finding of high CA19.9 in the serum of patients suffering non-malignant diseases (van der Veek et al., 2011), open questions about the biology of such molecule and the rationale of the use as a reliable marker. Little is known about CA19.9 structure, synthesis and secretion in the different tissues, and nothing about the origin of that circulating in health individuals. Recently, it was found that multiple proteins carry the epitope in pancreatic cancer and none appears responsible for cancer up-regulation (Yue et al., 2011b). This may suggest that the synthesis of the carbohydrate epitope is limiting in determining the serum levels. In fact, Narimatsu et al. (1998) reported that FUT3 (Lewis gene) dosage affects CA19.9 positively, while FUT2 (Secretor type α1,2 fucosyltransferase, competing for the synthesis) negatively. In colon cancer, however, among specific glycosyltransferases required for epitope biosynthesis (fucosyltransferase Fuc-TIII, α2,3 sialyltransferases ST3GalIV or ST3GalIII, and β1,3 galactosyltransferase β3Gal-T5) none is up-regulated (Ito et al., 1997; Kudo et al., 1998; Misonou et al., 2009; Dall’Olio et al., 2012). In particular, β3Gal-T5 transcript is expressed in normal colon mucosa but dramatically down-regulated in adenocarcinomas (Salvini et al., 2001; Ishihiki et al., 2003; Caretti et al., 2012), confirming the observation that β1,3Gal-T activity is impaired in colon cancers (Seko et al., 1996; Misonou et al., 2009), where type 2 chain oligosaccharides (based on Galβ1-4GlcNAc sequence) predominate over type 1 chain (based on Galβ1-3GlcNAc sequence) in various glycoconjugates (Misonou et al., 2009). On this light, the

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1 The abbreviations used are: sLea, sialyl-Lewis a, NeuAcα2-3Galβ1-3[Fucα1-4]GlcNAc; sLex, sialyl-Lewis x, NeuAcα2-3Galβ1-4[Fucα1-3]GlcNAc; Gal-T, galactosyltransferase; Fuc-TIII, α1,3,1,4fucosyltransferase type III (product of FUT3 gene); ElA, enzyme immunoassay; CEA, carcinoembryonic antigen.
concept that CA19.9 is a cancer-associated antigen in human colon appears paradoxical and deserves more investigations.

To address this issue we measured the levels of β3Gal-T5 transcript and β1,3Gal-T activity in biotic samples pairs from colon adenocarcinomas and surrounding normal mucosa, and determined the amount of sLea antigen expressed in the tissues. We also evaluated in cultured cell lines expressing the antigen their ability to secrete it in the medium. We then compared the obtained data with the amount of antigen detected in control and patient sera.

2. Materials and methods

2.1. Cell cultures

COLO-205, HCT-15, CACO-2, HT-29 (from human colon adenocarcinomas), and MKN-45 (from human gastric cancer) cells were cultured as previously described (Valli et al., 1998; Salvini et al., 2001). MKN-45 cells expressing Fuc-TIII, named MKN-45-FT, and human pancreatic adenocarcinoma cells BxPC3 and Panc-1, were obtained and cultured as reported (Mare and Trinchera, 2004). CACO-2 cells differentiated in culture were obtained according to the procedure described (Isshiki et al., 2003).

2.2. Preparation of human colon samples and cultured cells for analytical procedures

Human biopptic specimens were collected at surgery, immediately frozen in dry ice and placed in liquid nitrogen until used, as previously reported (Salvini et al., 2001; Caretti et al., 2012). They originated from different parts of the large intestine, including the rectum (samples 3 and 6). Cancers were staged from B2 to C2 (Dukes') and graded from moderately to poorly differentiated. The health mucosa surrounding the cancer lesion in the same surgical resection was referred to as the normal mucosa. For RNA extraction, 1–2 mm³ of frozen material was cut, placed in 0.3 ml of lysis buffer (Ambion RNAqueous minikit). In vitro, homogenated with a rotary homogenizer, and processed for total RNA extraction and DNA analysis according to the manufacturer's recommendations. For biochemical analysis, 2–3 mm³ of frozen material was cut, carefully rinsed twice in 0.5 ml of ice-cold phosphate buffered saline, placed in 0.5 ml of 0.1 M Tris/HCl buffer, pH 7.5, and homogenated with a rotary homogenizer. An aliquot of the homogenate was diluted to different protein concentrations and used for dot-blot, another aliquot was made 0.5 mg/ml Triton X-100 and used as the enzyme source for β1,3Gal-T in vitro assay, and a third aliquot was lysated for ELISA according to a published procedure (Baeckström et al., 1991). Cultured cells were harvested and freshly processed for flow cytometry as reported (Bardon et al., 1999; Salvini et al., 2001). For dot-blot, ELA, and enzyme assay, cell pellets were treated as described for biotic samples but vortexing instead of using the rotary homogenizer.

2.3. RNA analysis

For transcript quantification, competitive RT-PCR was performed essentially as previously reported (Trinchera et al., 2011; Caretti et al., 2012). First strand cDNA was prepared for samples and controls in the presence, or omitting, the reverse transcriptase, respectively, and reactions incubated under reported conditions. It was amplified (25 μl reaction volume) in the presence of 10 fg of competitor for 35 cycles (β3Gal-T5), or 10 pg of competitor for 25 cycles (β-actin), under reported conditions. No amplification was detected when the control reactions were used as template. Human β-actin and β3Gal-T5 competitors, and oligonucleotide primers, were those already described (Salvini et al., 2001). Fuc-TIII transcript was quantitated exactly as reported (Trinchera et al., 2011).

2.4. Enzyme assay

β1,3Gal-T activity was determined in the reported reaction mixture (Salvini et al., 2001), using 0.5 M GlcNAc as acceptor, in the presence of cell or tissue homogenates at various protein concentrations: 0.5–4.0 mg/ml for cell lines and clones, 0.5–2.0 mg/ml for normal colon mucosa, and 5.0–10 mg/ml for colon cancers. Incubations were done at 37 °C for 120 (cancers) or 60 min (all others). At the end of incubation, reaction products were assayed by Dowex chromatography and characterized according to previously reported protocols (Valli et al., 1998). In all cases the reaction product was found to be a disaccharide sensitive to β1,3galactosidase, as expected. In fact, GlcNAc is not used as acceptor by β1,4galactosyltransferases under the reported assay conditions (Valli et al., 1998; Bardon et al., 1999). One unit of β1,3Gal-T activity corresponds to one nanomole of transferred Gal per mg of protein homogenate per hour.

2.5. sLeα detection

Detection of sLeα by immunofluorescence and flow cytometry was performed as previously reported (Salvini et al., 2001). For dot-blot, aliquots from homogenates, sera, bile, or culture media were applied to the membrane by vacuum aspiration. Serial dilutions of samples were performed in preliminary experiments to set the optimal protein concentrations and amounts needed for detection. Blotting membranes were washed, blocked, stained with primary and peroxidase-labeled secondary antibodies, and visualized by enhanced chemiluminescence as reported for western-blot (Caretti et al., 2012). Monoclonal anti-sLea (from hybridoma 1116-NS-19-9) antibody was prepared as reported (Bardon et al., 1999). Quantification of CA19.9 in serum and bile samples, in culture media, or in tissue and cell lysates was performed by ELISA with an automated analytical system (Cobas Core II, Roche Diagnostics) equipped with dedicated reagents, according to the instructions of the manufacturer.

3. Results

3.1. β1,3Gal-T activity, β3Gal-T5 transcript levels, and sLeα expression in normal colon mucosa and adenocarcinomas.

To evaluate the actual ability of colon tissues to synthesize large amounts of sLeα, we determined β3Gal-T5 transcript and activity in 9 sample pairs from patient biopsies, each representing colon adenocarcinomas and surrounding normal mucosa. Serum levels of CA19.9 were 115, 84.6, and 19.2 U/ml in patients 9, 8, and 4, respectively (normal range <37 U/ml), and not available in the others. Fuc-TIII transcript was found to be heterogeneously expressed in all samples (not shown), as expected (Ito et al., 1997; Salvini et al., 2001; Trinchera et al., 2011).

The amount of β3Gal-T5 transcript expressed in normal colon mucosa is high but variable, ranging from 3 to 20 fg/pg of β-actin, and the enzyme activity ranges from 22 to 120 U. In cancer samples, β1,3Gal-T activity closely follows transcript down-regulation (Fig. 1, panel A, note the different scales used). The calculated reduction is 36.2-folds on average, ranging from a minimum of 6.2-folds in sample 6 to a maximum of 68-folds in sample 9. Interestingly, in a single adenocarcinoma case (sample 5) where the transcript level is relatively maintained (0.9 fg/pg of β-actin) the enzyme activity reaches the value of 13.2 U, while it is lower than 4 U in all other samples. This data indicate that β3Gal-T5 down-regulation in colon cancers leads to extremely low values of enzyme activity.
To directly evaluate the synthesis of sLea in the tissues, sample homogenates were analyzed through immunostaining of dot-blot. Surprisingly, under conditions used for detection in cell lines, sLea is undetectable in both normal colon mucosa and adenocarcinomas (Fig. 1, panels B and C). Using much more protein in the blots, sLea is detected in many normal mucosa specimens and faintly in few cancers, but remains undetectable in others. To better assess the actual amount of CA19.9 expressed in colon tissue with respect to cell lines, we measured CA19.9 by EIA in tissue and cell lysates. Such assay linearly detects CA19.9 in COLO-205 and BxPC3 lysates starting from 5.0 and 10.0 μg/ml of cell protein, respectively, while none is detected using up to 40.0 μg/ml of protein lysate from HCT-15, a cell line known to lack sLea (Fig. 2, lower panel). Using 20.0 μg/ml of protein from colon tissue lysates, CA19.9 is detected at very low levels in one adenocarcinoma and one normal mucosa sample, and remains undistinguishable from the background (value detected in HCT-15 cells) in all other samples (Fig. 2, upper panel). These findings demonstrate that no accumulation of sLea occurs in colon adenocarcinomas.

3.2. Expression and secretion of sLea in cultured cell lines

As a model for studying sLea secretion, we evaluated in cultured cell lines the relationship between sLea secreted in the culture medium and expressed on the cell surface, since cancer cell lines secrete mucin-type glycoproteins carrying the epitope (Baeckström et al., 1991, 1995). To this purpose, a set of cancer cell lines from the digestive tract expressing different amount of antigens were selected in preliminary experiments. They were plated at the same density and cultured for 48 h. The amount of sLea present on the surface was then quantitated by immunofluorescence and flow cytometry, while that accumulated in the culture medium was detected by dot-blot immunostaining and quantitated by EIA (Fig. 3). Among colon adenocarcinoma cell lines, COLO-205 cells, expressing very much sLea on the surface, secrete large amount of antigen in the medium, while CACO-2 and HT-29, expressing low amount of sLea on the surface, secrete minimal amount of antigen, as detected by both dot-blot immunostaining and EIA. Moreover, all cell lines lacking expression on the surface, as HCT-15, Panc-1, and MKN-45, do not secrete detectable antigen at all. In addition, CACO-2 cells differentiated in culture and expressing about three times more β1,3Gal-T activity than undifferentiated cells, also express more sLea on the surface and secrete more sLea in the medium. On the other side, BxPC3 cells (from pancreas) expressing intermediate amount of sLea on the surface, and MKN-45-FT (from stomach), a recombinant clone forced to express sLea on the surface through over-expression of Fuc-TIII, secrete a moderate amount of antigen in the medium as detected by dot-blot immunostaining, but EIA detects much more antigen in the culture medium of BxPC3. Altogether these data indicate that secretion of CA19.9 antigen in cancer cell lines does not occur in the absence of concurrent expression on the surface, and only cells expressing β3Gal-T5 are able to secrete relevant amounts of antigen. Interestingly, there is a very good correlation between the amounts of antigen expressed on the surface and secreted in the culture medium of colon cancer.
cells (Fig. 3C), while those from pancreas or stomach seem to have different behavior (Fig. 3C).

Noteworthy, \( \beta_3 \text{Gal-T} \) activity measured in CACO-2 and HT-29 cells is 15.2 and 11.4 U, respectively. Such values, able to direct only minimal secretion of antigen, are still 7- to 22-folds larger than those measured in cancers patients 8 and 9 of Fig. 1A, whose serum CA19.9 is indeed above the cut-off level.

### 3.3. Detection of sLea in the serum

Unfortunately, we lack serum samples from the same patients where the biopsies are from. As an alternative, we checked samples from individuals whose serum levels of CA19.9 cover a wide range. Fig. 4 shows that CA19.9 is very easily detected by dot-blot immunostaining, even blotting minimal volumes of sera having low antigen amounts measured by EIA. This indicates that the blotting procedure is highly sensitive and that at least some immunogenic properties of CA19.9 present in tissues, cells, and sera are similar. On the other side, the quantitative relationship between EIA and blotting detection in cancer patient serum and in the bile is very poor, as found in the culture medium of BxPC3 cells. Such discrepancy probably accounts for the structural heterogeneity of the antigen, already proved to occur among different biological sources but even within the same one (Baekström et al., 1991, 1994, 1995; Yue et al., 2011a, 2011b). At this regard, it is interesting to note that CA19.9 circulating in the serum of cancer patients or present in the bile is better detected by EIA than by blotting, while that found in colon biopsies or colon-derived cell membranes and culture media is not.

### 4. Discussion

We found that the synthesis and secretion of CA19.9 antigen in colon cancer presents several problems not supposed so far. In fact we found that \( \beta_3 \text{Gal-T} \), the enzyme responsible for one of the glycosylation steps necessary for epitope synthesis, is so strongly down-regulated in colon cancers that drops down to values far below those needed to synthesize the antigen in cell lines. Furthermore, the antigen is not detectable in the cancer biopsies, or more difficult to detect than in the normal mucosa. Since in colon cancer cell lines the amount of antigen secreted in the culture medium is proportional to that expressed on the cell surface, our data open the question if colon cancers are able to secrete large amounts of sLea.

In previous studies, sLea was found in some colon cancers analyzed by histochemistry (Ito et al., 1997; Kudo et al., 1998) or sporadically distributed in some parts of the tumors (Tabuchi et al., 1988), but western blot and histochemical data were not overlapping (Kudo et al., 1998). On the light of our present data, we suggest that such discrepancies reflect the degree of blood contamination in the tissues, or depend on binding to the cell membrane of the antigen carried by the blood. In fact, we found that patient serum is very easily stained by anti-sLea antibody, so that 0.01 ml of serum seem to contain more sLea than some tens or even hundreds of milligrams of colon cancer tissue. Moreover, studies performed on colon cancer at surgery (Tabuchi et al., 1988; Nakagoe et al., 2000) showed that CEA and sLex, but not CA19.9, are more elevated in the blood from vessels draining the tumor region than in peripheral blood, with spikes coincident with surgery manipulation. Since CEA (Maxwell, 1999) and sLex (Trinchera et al., 2011) are actually synthesized by colon cancer cells, these in vivo results corroborate the data that we obtained in cultured cell lines. It would be very interesting in the future to validate such data in vivo comparing antigen levels in cancer biopsies and serum samples from the same patient. At this stage, we speculate that the metabolic origin of circulating CA19.9 should be searched in other tissues actually able to synthesize and secrete relevant amounts of antigen. Normal intestinal mucosa is a potential candidate, due to the high expression levels of glycosyltransferases as \( \beta_3 \text{Gal-T} \) and

![Fig. 3. Expression on the surface and secretion in the culture medium of sLea synthesized in different cell lines. Different cell lines were seeded at the same density, cultured for 48 h, harvested, and analyzed together with the culture media as detailed under Section 2. (A) sLea antigen expressed on the cell surface (full bars) was determined by immunostaining and flow cytometry performed on a fresh aliquot of the cell pellet. \( \beta_3 \text{Gal-T} \) activity (empty bars) was determined by in vitro assay as in Fig. 1. Results are the mean ± standard deviation for three determinations. (B) sLea secreted in the culture media was detected by immunostaining of dot-blot samples obtained with 50 µl of culture media, or quantified by EIA. Numbers are the mean for three determinations; standard deviations were less than 15%. (C) Correlation between sLea measured on the surface (by flow cytometry) and CA19.9 secreted in the culture media (quantitated by EIA) of colon cancer cell lines (HCT-15, CACO-2, HT-29, differentiated CACO-2 and COLO-205). Input data were from panel B. Correlation coefficient is 0.998 as determined by linear regression.](image)

![Fig. 4. Detection of CA19.9 in patient serum and in the bile. The amounts of serum or bile indicated were brought to the same volume, blotted, and stained with anti-sLea antibody as in Fig. 1. EIA quantification in the samples is shown at the top of the panel. Numbers are the mean for three determinations; standard deviations were less than 15%. “A-F” are the sample numbers indicated in the following lines: A is from a health control, B, E, and F from cancer patients, C from a chronic alcoholic, and D from a chronic C virus hepatitis patient. G is a bile sample, collected at cholecystectomy, that was diluted ten-times before blotting or assaying.](image)
Fuc-TIII. However, it does not accumulate large quantities of anti-
gen under basal conditions, probably because several additional
factors, unknown at present, control CA19.9 synthesis and secre-
tion in vivo, and any of them may be triggered by the cancer
lesion. The biliary tract is also a candidate, because an amazing
huge amounts of CA19.9 (millions of U/ml) is present in the bile,
as reported (Baeckström et al., 1994) and confirmed in our experi-
ments. However, a mechanism allowing CA19.9 elevation without
jaundice needs to be envisaged in this case. Anyway, our data are
compatible with the observation that appreciable levels of CA19.9
are commonly detected in healthy individuals (the source of which
remains totally unknown), and elevated levels, including extremely
high levels, occur in several benign conditions affecting the hepa-
tobiliary system (Mann et al., 2000; Akođan et al., 2001; van der Veek et al., 2011).

CA19.9 circulating in cancer patients was originally reported as an
high molecular weight mucin (size above 5.0 × 10^6 Da) (Magnani et al., 1983), perhaps difficult to immobilize on blotting mem-
banes. It is also predictable that mucinics bear several epitopes all recognized by EIA but not by dot-blot immunostain-
ning. We speculate that CA19.9 sources providing low-moderate
dot-blot staining but high titer by EIA, as the culture medium of
pancreatic BxPC3 cells, the bile, and the blood from some cancer
patients (E and F in Fig. 4), do contain a high molecular weight
form of the antigen, while the others do not. They include the cul-
ture medium from colon cancer cell lines and cell lysates from colon
cell lines and biopsies, where dot-blot immunostaining correlates
with EIA. Accordingly, the main molecules carrying CA19.9 in the
culture medium of Colo-205 cells were identified as CD-43 gly-
coforms (Baeckström et al., 1995), 150–300 kDa in size, expected
to easily bind blotting membranes. The molecular size heterogene-
ity of CA19.9 (Yue et al., 2011a) may be helpful in the future in
the attempt to identify the actual source of circulating antigen, an
information that we believe necessary to answer several clinical
questions.

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