A Transfected Sialyltransferase That Is Elevated in Breast Cancer and Localizes to the medial/trans-Golgi Apparatus Inhibits the Development of core-2–based O-Glycans

Caroline Whitehouse,* Joy Burchell,* Stephen Gschmeissner,† Inka Brockhausen,§ Kenneth O. Lloyd,‖ and Joyce Taylor-Papadimitriou*

*Epithelial Cell Biology Laboratory, †Electron Microscopy Unit, Imperial Cancer Research Fund, Lincoln’s Inn Fields, London WC2A 3PX, UK; ‡Department of Biochemistry, Hospital for Sick Children and University of Toronto, Toronto M5G 1X8, Canada; and ‖Sloan-Kettering Institute, New York 10021

Abstract. The α2,3 sialyltransferase, α2,3 SAT (O), catalyzes the transfer of sialic acid to Galβ1,3 N-acetyl-d-galactosamine (GalNAc) (core-1) in mucin type O-glycosylation, and thus terminates chain extension. A Core-2 branch can also be formed from core-1 by the core-2 β1,6 N-acetyl-d-glucosamine transferase (β1,6 GlcNAc T) that leads to chain extension. Increased levels of the α2,3 SAT (O) and decreased levels of the core-2 β1,6 GlcNAc T are seen in breast cancer cells and correlate with differences in the structure of the O-glycans synthesized (Brockhausen et al., 1995; Lloyd et al., 1996). Since in mucin type O-glycosylation sugars are added individually and sequentially in the Golgi apparatus, the position of the transferases, as well as their activity, can determine the final structure of the O-glycans synthesized. A cDNA coding for the human α2,3 SAT (O) tagged with an immunoreactive epitope from the myc gene has been used to map the position of the glycosyltransferase in nontumorigenic (MTSV1-7) and malignant (T47D) breast epithelial cell lines. Transfectants were analyzed for expression of the enzyme at the level of message and protein, as well as for enzymic activity. In T47D cells, which do not express core-2 β1,6 GlcNAc T, the increased activity of the sialyltransferase correlated with increased sialylation of core-1 O-glycans on the epithelial mucin MUC1. Furthermore, in MTSV1-7 cells, which do express core-2 β1,6 GlcNAc T, an increase in sialylated core-1 structures is accompanied by a reduction in the ratio of GlcNAc: GalNAc in the O-glycans attached to MUC1, implying a decrease in branching. Using quantitative immunoelectron microscopy, the sialyltransferase was mapped to the medial- and trans-Golgi cisternae, with some being present in the TGN. The data represent the first fine mapping of a sialyltransferase specifically active in O-glycosylation and demonstrate that the structure of O-glycans synthesized by a cell can be manipulated by transfecting with recombinant glycosyltransferases.

In eukaryotic cells, proteins are synthesized in the ER from extensions to different locations, either within the cell or to the plasma membrane. Along the exocytic pathway, proteins undergo various modifications including proteolytic cleavage, glycosylation, and sulfation. Within this pathway there are two main types of glycosylation, N-linked and mucin-type O-linked. The addition of N-linked glycans is initiated in the ER by the addition to asparagine of an oligosaccharide chain via an intermediate lipid carrier. This chain is then modified by trimming and the addition of sugars. In contrast, mucin-type O-linked glycosylation is thought to be initiated in the cis-Golgi cisternae where the first sugar, N-acetyl-a-galactosamine (GalNAc),1 is added to the hydroxyl groups of serine and threonine (Roth et al., 1994; Clausen and Bennett, 1996). The oligosaccharide chains are then built up by the sequential addition of individual sugars, each reaction being catalyzed by a specific enzyme or enzymes (Brockhausen, 1996). Thus, the final structure of the O-glycans is strongly influenced, not only by the level of activity of an enzyme but also by its position within the Golgi apparatus.

After the addition of the first GalNAc to threonine or

1. Abbreviations used in this paper: GalNAc, N-acetyl-a-galactosamine; GlcNAc, N-acetyl-d-glucosamine; GalT, β1-4 galactosyltransferase; HPAEC, high performance anion exchange chromatography; PNA, peanut agglutinin.
serine, chains are extended via various core structures, generally with polyolactosamine units, and the final structures (added to the same core protein) can be different in different tissues, depending on the profile of glycosyltransferases expressed (Brockhausen, 1996). Of great interest is the observation that, in carcinomas, the composition of the O-glycans added to the glycoproteins produced by the tumor cells may be altered compared with those expressed in normal cells. This has a profound effect on the structure of those glycoproteins that carry multiple O-linked glycans such as the epithelial mucins. This change in glycosylation pattern has been best documented in breast cancer where the MUC1 mucin (Gendler et al., 1990) has been shown to carry shorter and less complex O-glycans than the mucin produced by normal cells (Hanisch et al., 1989; Hull et al., 1989; Lloyd et al., 1996).

The key glycosyltransferases involved in the changes seen in breast malignancies are the α2,3 sialyltransferase (EC 2.4.99.4), which adds sialic acid to Galβ1,3 GalNAc (core-1), and the β1,6 N-acetyl-d-glucosamine (GlcNAc) transferase (EC 2.4.1.102), which forms core-2 from core-1 and is crucial for chain branching (Fig. 1) (Kuhns et al., 1993). These enzymes use the same substrate, and their effect is to terminate or initiate chain branching leading to extension, respectively. We have recently observed that the activity of the chain terminating enzyme, α2,3 sialyltransferase, is increased 8–10-fold in some breast cancer cell lines, while core-2 β1,6 GlcNAc T activity is either lost or reduced (Brockhausen et al., 1995). The expression of the MUC1 mucin is upregulated in breast cancers and the difference in glycosylation pattern causes the cancer-associated mucin to be antigenically distinct from the normal mucin. MUC1-based immunogens are therefore prime candidates for cancer vaccines and several formulations are being tested in the clinic.

The position of the α2,3 sialyltransferase in the Golgi apparatus will determine to some degree whether it can compete directly with core-2 β1,6 GlcNAc T for the Galβ1,3 GalNAc substrate. It is therefore important to precisely map its distribution within the Golgi apparatus. cDNAs coding for the α2,3 sialyltransferase have been isolated from porcine submaxillary glands (Gillespie et al., 1992), murine brain (Lee et al., 1993), and more recently from human placenta (Chang et al., 1995). It is therefore now possible to directly locate the transferase by transfecting a cDNA tagged with a sequence encoding an immunoreactive epitope (Nilsson et al., 1993). We have tagged the cDNA coding for the human α2,3 SAT (O) with an epitope from the mtsv1 gene, and the construct has been used to transduce or transfect two mammary epithelial cell lines, both of which express the MUC1 mucin. The T47D cell line was derived from a metastatic breast carcinoma (Keydar et al., 1979), does not express core-2 β1,6 GlcNAc T, even at the level of mRNA (Brockhausen et al., 1995), and produces MUC1 carrying short O-glycans (Lloyd et al., 1996). The MTSV1-7 cell line was derived from normal human milk epithelial cells (Bartek et al., 1991) and shows many characteristics of normal cells (Shearer et al., 1992), including the ability to add core-2–based O-glycans to the MUC1 mucin (Lloyd et al., 1996). In the MTSV1-7 cell line, the transfected α2,3 SAT (O) has been localized to the medial- and trans-Golgi cisternae, with some enzyme being detected in the TGN. We have also demonstrated that transfection of a cell with the α2,3 SAT (O) results in increased sialylation of the surface protein, the MUC1 mucin, thus allowing the manipulation of O-linked glycosylation. In the MTSV1-7 transfectants, the increase in sialylated core-1 is accompanied by a decrease in the GlcNAc content of the O-glycans attached to MUC1, suggesting a reduction in the synthesis of core-2–based structures.

Materials and Methods

Cell Culture

The cell line MTSV1-7 was grown in DME supplemented with 10% FCS (GIBCO BRL, Gaithersburg, MD), 10 μg/ml insulin (Sigma Chemical Co., St. Louis, MO), 5 μg/ml hydrocortisone (Sigma Chemical Co.), and 0.3 μg/ml glutamine. The retroviral infectants of MTSV1-7 were maintained in the same media with the addition of 2 μg/ml of puromycin (Sigma Chemical Co.). T47D and AM12 cells were grown in DME supplemented with 10% FCS (GIBCO BRL) and 0.3 μg/ml glutamine. The T47D transfectants were grown in the same medium as the parental line but with the addition of 500 μg/ml of G418 (GIBCO BRL).

Development of the α2,3 Sialyltransferase mtsv1 Constructs

The cDNA encoding human α2,3 SAT (O) was cloned into the HindIII/XbaI site of pBluescript (Stratagene, La Jolla, CA). This was digested with MroI and XbaI, and two annealing oligos encoding the myc epitope were inserted. The sequence of these oligos was as follows: 5′-CCGGATCTCCAAAGGGGAGACCAGAACAACTCTACAGACAGAGACATGTTGAT-3′ and 5′-CTAGATCACCAGCTCTTTACAGATCTGTCAGGATCCATTGGAT-3′. Bases encoding the myc epitope recognized by the 9E10 mAb (Evan et al., 1985) are underlined.

Figure 1. Alternative pathways for O-linked glycosylation of MUC1 involving either chain branching via core-2 β1,6 GlcNAc transferase or chain termination by addition of sialic acid via α2,3 sialyltransferase.
The product was digested with HindIII/NotI and cloned into the pcDNAIneo expression vector (Invitrogen, San Diego, CA). This neo3STMYC construct was sequenced, and expression of the myc epitope tag was checked by transient expression in COS cells before being used for stable transfection of T47D cells.

For production of an amphotropic c2,3 sialytransferase-expressing retrovirus, the HindIII/XbaI fragment was excised from pBluescript, blunted ended with Klenow, and subcloned into the StuII site of the pBabe puro vector. The amphotropic packaging cell line AM12 was transfected with this construct using calcium phosphate-mediated transfection. Briefly, AM12 cells were grown to 70% confluency in 10-cm dishes and refed with fresh medium 1 h before transfection. 15 μg of pBabe puro-3STMYC DNA and 25 μg of carrier salmon sperm DNA were coprecipitated with calcium phosphate at pH 7. After 6 h at 37°C, dishes were rinsed five times with serum-free medium and then refed with fresh medium. 48 h after transfection, cells were split 1:10 into growth medium containing 2 μg/ml puromycin for selection. Medium was changed every 3–4 d for 4 wk until selection was complete. The c2,3 sialytransferase-retrovirus producer cell line was grown to 70% confluency, and the spent medium was replaced with half the volume of fresh medium. 3 d later, the virus-containing medium was removed, filtered, quick frozen in dry ice, and stored at −70°C.

**Transfection and Transduction of Cell Lines**

T47D cell line was transducted directly with the neo3STMYC construct using the method of calcium phosphate transfection described above with the following alterations. The DNA precipitate was left on the cells overnight after being washed with serum-free medium. The cells were then washed three times with PBS before refedding with fresh medium. 2 d later, the plates were split 1:10 into medium containing 500 μg/ml G418, and selection was carried out until individual clones were isolated and expanded. Selected clones were referred to as T47D 3STMYC. Cells were also transfected with the vector pcDNAIneo and a clone, T47D neo, was isolated.

For transduction of MTSV-1-7 cells with the c2,3 sialytransferase retrovirus, the cells were grown to 70% confluency in 10-cm dishes and infected with 3 ml of viral stock (with or without dilution) with 8 μg/ml polybrene. Infection proceeded for 3 h before virus was replaced with fresh medium. 2 d later, the plates were split 1:10 into medium containing 2 μg/ml puromycin, and selection continued until individual colonies could be identified and expanded. Selected colonies were referred to as T47D 3STMYC. Cells were also infected with a retrovirus derived from the pBabe puro vector, and a clone MTSV-1-7 pBpuro was isolated.

**Northern Analysis of mRNA from Cell Lines**

Total cellular RNA from the cell lines was isolated according to the method of Chomczynski and Sacchi (1987). 25 μg of RNA from each cell line was denatured in 1× MOPS buffer, 0.66 M formaldehyde, and 50% (vol/vol) formamide, and subsequently size fractionated on a 1.3% agarose-formaldehyde gel. The RNA was transferred and immobilized onto Hybond-N membrane (Amersham Intl., Little Chalfont, UK). The membrane was hybridized with a 1.2-kb HindIII/XbaI cDNA fragment from the c2,3 SAT (O) plasmid according to the method of Church and Gilbert (1984) and washed to highest stringency as described previously (Lloyd et al., 1996). Briefly, cells were metabolically labeled with 100 μCi/ml [3H]glucosamine-hydrochloride (Amersham Intl.) and MUC1 immunoprecipitated with CT1, an antibody to the cytoplasmic tail of MUC1 (Pemberton et al., 1992). The carbohydrate side chains were released by alkaline borohydride treatment. Samples containing 10,000 cpm were then separated on a Carbo Pak PA1 column (Dionex Corp., Sunnyvale, CA) using a gradient of 0.2 M NaOH to 0.2 M NaOH–0.25 M sodium acetate at 1.0 ml/min over 30 min (Lloyd and Savage, 1991). Collected radioactive fractions were neutralized with 1 M HCl before counting. For hexosamine analysis, the immunoprecipitate was eluted in 2% SDS and hydrolyzed in 2 N trifluoroacetic acid at 100°C for 3 h, and 5,000 cpm samples were analyzed by HPAEC on a CarboPak PA1 column by isocratic elution with 0.01 M NaOH at 1.0 ml/min.

**FACS® Analysis**

**Reactivity of Peanut Lectin with Live Cells.** Cells were incubated with or without neuraminidase and analyzed by FACS® can for Anachis hypogaea peanut agglutinin (PNA) (Sigma Chemical Co.) lectin binding as described previously (Burchell and Taylor-Papadimitriou, 1993).

**Detection of the Tagged Sialytransferase in Permeabilized Cells.** Cells were stained with 9E10 mAb after incubating the cells with 0.3% saponin for 20 min. Subsequent steps were as previously described (Burchell and Taylor-Papadimitriou, 1993) with the exception that 0.1% saponin was included in all incubations and washes, which were carried out at room temperature. The reactivity of an mAb (LE61) to keratin 18 (Lane, 1982) was included as a positive control for staining the permeabilized cells.

**Immunofluorescence Staining**

Cells were grown on glass coverslips, washed with PBS, and fixed with 4% paraformaldehyde for 15 min. Cells were permeabilized with 0.1% Triton for 5 min, and then nonspecific binding was blocked with 10% FCS/PBS for 30 min. The cells were then incubated with the 9E10 mAb to the myc epitope (10 μg/ml) or TEx-1, a rabbit antiserum which recognizes mannosidase II (diluted 1:50) (Slusarewicz, 1994), and binding was detected with FITC-conjugated goat anti–mouse (diluted 1:40; Dako) or rhodamine-labeled swine anti-rabbit (diluted 1:40; Dako) secondary antibodies.

**Immunoelectron Microscopy**

Cells were fixed for 1 h at room temperature in 0.1% glutaraldehyde/4% paraformaldehyde before being scraped, spun, and stored overnight in 2% paraformaldehyde at 4°C. Double labeling was performed as described previously (Slot et al., 1991). Antibody details are as follows: the grids were incubated overnight at 4°C with 9E10 supernatant, diluted 1:5, followed, after washing, for 30 min with goat anti–mouse Ig conjugated to 10-nm gold particles. Galactosyltransferase rabbit polyclonal antibody (Watzele et al., 1991) was diluted 1:50 in 1% BSA, and the grids were incubated for 30 min at room temperature before washing followed by a 30-min incubation with protein A coupled to 5-nm gold. The grids were embedded in 1.8% methyl cellulose/0.4% uranyl acetate before examination using a Zeiss 10C (Oberkochen, Germany) or JEOL 1010 (JEOL USA, Peabody, MA) electron microscope.

**Immunogold Quantitation**

Golgi apparatus profiles were selected at random, photographed, and printed at a final magnification of 75,000. The compartments of the Golgi apparatus were defined as described previously (Nilsson et al., 1993). Briefly, the trans ciservae were defined as the last continuous cisternae that la-
beled for Gal-T, and the TGN comprises the tubuloreticular network adjacent to the trans side of the Golgi apparatus stack. The boundary of the Golgi cisternae and TGN (defined as the interface between the outermost membranes of the tubular network and the adjacent cytoplasm) (Rabouille et al., 1995) was drawn on each micrograph. Gold particles over the TGN and the Golgi cisternae were directly counted and included gold over budding vesicles but not structures such as vacuolar endosomes, which can be found in the TGN and were excluded.

To assess the polarized distribution of gold particles within the Golgi apparatus, for every Golgi apparatus analyzed, the position of each gold particle was calculated as a fraction of the distance across the Golgi apparatus as follows: \(d_1 = \text{distance from cis face}, \quad d_2 = \text{distance from trans face}, \quad \text{and} \quad \frac{d_1}{d_1 + d_2} = \text{the position within the Golgi apparatus.}\n
Individual gold particles were assigned to one of ten equal fractions, and the sum of gold particles of each size in each fraction was expressed as a percentage frequency of the total number of each size of gold (Rabouille, C., and T. Nilsson, personal communication).

Comparison of Sequences Coding for the Endogenous and Transfected \(\alpha_{2,3} SAT(O)\)

Bases 1–278 of the \(\alpha_{2,3} SAT(O)\) DNA were PCR amplified from T47D and MTSV1-7 genomic DNA prepared as described previously (D’Souza et al., 1993), using the following oligonucleotides: 5′GAATTCGAA-TTCGACGTTCCGAAGATG and 5′AAGCTTAAAGCTTAAAGCG-CTGTGGCC. The PCR product was cloned into the HindIII/EcoRI sites of pBluescript and sequenced using the ABI PRISM automated sequencing method (Perkin-Elmer Corp., Norwalk, CT), for comparison with the \(\alpha_{2,3} SAT–expressing\) plasmid.

Results

Development of Stable Cell Lines Expressing \(\alpha_{2,3}\) Sialyltransferase

The cDNA encoding the human \(\alpha_{2,3}\) sialyltransferase tagged with a 30-bp stretch of DNA encoding the 9E10 epitope of the myc protein was cloned into a mammalian expression vector (pcDNAI neo). At the COOH terminus of this type II transmembrane protein, the 9E10 epitope is far away from the membrane-spanning domain that has been shown to be involved in the localization of glycosyltransferases to their correct compartment within the Golgi (Mac-hamer, 1993). The construct was transfected into the breast carcinoma cell line, T47D, and two stable cell lines (designated T47D 3ST 2 and 3) were selected in the presence of gentamycin. The MUC1 mucin produced by the parental T47D cells carries short O-glycans that do not contain core-2 structures (Lloyd et al., 1996).

The cell line MTSV1-7 has many characteristics of normal mammary epithelial cells. It is nontumorigenic, forms organized colonies in collagen gels (Berdichevsky and Taylor-Papadimitriou, 1991; Shearer et al., 1992; Lu et al., 1995), and glycosylates MUC1 in a manner similar to that of normal mammary epithelial cells in adding core-2–based O-glycans (Lloyd et al., 1996). MTSV1-7 has proven to be difficult to transfect, and a retrovirus expressing the myctagged \(\alpha_{2,3}\) sialyltransferase was constructed (see Materials and Methods) and used to transduce MTSV1-7. Five stable cell lines (designated MTSV1-7 3ST 3.10, 3.11, 3.14, 4.1, and 5.2) were selected in the presence of puromycin.

Expression and Activity of the \(\alpha_{2,3}\) Sialyltransferase in the Transfectants

Northern blot analysis revealed that all the selected cell lines were expressing the transfected DNA (Fig. 2 a) and, moreover, the expression level varied with the clone, with the MTSV1-7 transfectants expressing more than the T47D clones. The difference in molecular weight between the message in T47D clones and the MTSV1-7 is due to the \(\alpha_{2,3}\) sialyltransferase RNA in the MTSV1-7 cells being expressed within the context of the retrovirus vector. Fig. 2 b shows endogenous expression of \(\alpha_{2,3} SAT(O)\) in T47D and MTSV1-7 parental cell lines, where the Northern blot was exposed for a longer period of time.
Western blotting with the 9E10 antibody showed a specific band in the region of 46–48 kD, which corresponds to the myc-tagged α2,3 SAT (O). Fig. 3a shows the blots for the highest expressing MTSV1-7 and T47D transfecteds, indicating higher expression of the enzyme in the MTSV1-7 clone. Expression of the protein was also confirmed by FACS® analysis of permeabilized cells. Fig. 3b shows the 3STMYC clone of T47D, the MTSV1-7 3STMYC, and the T47D neo and MTSV1-7 puro cell lines. As seen on the Western blots, the expression of α2,3 SAT (O) by the T47D clones was considerably weaker than by the MTSV1-7 clones. However, the positive shift in 9E10 staining observed in the T47D transfecteds was consistent and significant as the expression of keratin 18 completely overlaps in the T47D neo and T47D transfecteds (Fig. 3b). Functionality of the expressed enzyme was demonstrated by showing an increase in the specific activity of the enzyme in the transfected clones. Table I shows the activity of the enzyme in cell extracts of the clones in comparison with cells transfected or transduced with vector alone. In the T47D clones, α2,3 sialyltransferase activity was elevated about sevenfold compared with T47D neo, whereas the MTSV1-7 clones had increased activity ranging from 19–30-fold in comparison with MTSV1-7 puro or wild-type cells. The levels of activity of the enzyme correlated with the levels of expression, (Figs. 2 and 3; Table I). (The levels of enzyme activity of the T47D series of cell lines and the MTSV1-7 series are not comparable since they were done at different times and do not reflect the increased level of activity present in the T47D cells [Brockhausen et al., 1995]). The overexpression of α2,3 sialyltransferase had no obvious effect on the morphology of the Golgi apparatus as shown by electron micrographs from wild-type MTSV1-7 cells (Fig. 4a) and one of the MTSV1-7 3ST clones (Fig. 4b).

Figure 3. Expression of α2,3 SAT (O) protein. (a) Western blot analysis of myc-tagged α2,3 SAT (O) expression in T47D and MTSV1-7 3STMYC transfecteds, carried out as described in Materials and Methods. The expected band was detected by the anti-myc antibody (9E10) in the transfecteds but not in the parental cell lines. M, markers. (b) Permeabilized cells were stained with antibodies 9E10 (to the myc epitope) or LE61 (to keratin 18) and subjected to FACS® analysis as described in Materials and Methods. (Left) Staining of T47D transfecteds. (Right) Staining of MTSV1-7 transfecteds. (Thin continuous lines) T47D neo or MTSV1-7 puro stained with 9E10; (thick continuous lines) T47D neo or MTSV1-7 puro stained with LE61; (dashed lines) T47D 3STMYC or MTSV1-7 3STMYC stained with 9E10; (dotted lines) T47D 3STMYC or MTSV1-7 3STMYC stained with LE61.

Table I. Sialyltransferase Activity in Mammary Epithelial Cell Lines Transfected with cDNA Encoding α2,3 Sialyltransferase, α2,3 SAT (O), or with Vector Only

| Cell line   | Activity (nmol/h/mg) |
|-------------|----------------------|
| MTSV1-7 puro| 2.2                  |
| MTSV1-7 3STMYC 3.10 | 64.3                       |
| MTSV1-7 3STMYC 3.11 | 42.2                       |
| MTSV1-7 3STMYC 3.14 | 45.0                       |
| MTSV1-7 3STMYC 4.1 | 67.3                       |
| MTSV1-7 3STMYC 5.2 | 55.7                       |
| T47D neo    | 2.4                  |
| T47D 3STMYC 2 | 13.9                       |
| T47D 3STMYC 3 | 17.6                       |

Enzyme assays were carried out as described previously (Brockhausen et al., 1995). The absolute values for α2,3 sialyltransferase activity in the MTSV1-7 and T47D series of cell lines are not directly comparable as the assays on the two series were performed at different times.
and 2 (Fig. 5, a and b). We have previously shown that peak 1 contains neutral oligosaccharides consisting mainly of Galβ1-3 GalNAc-ol with a small amount of GalNAc-ol, whereas peak 2 contains monosialylated Galβ1-3 GalNAc-ol (Lloyd et al., 1996). Strikingly, the α2,3 SAT (O) transfected T47D clones show an increase in the proportion of the monosialylated peak (Fig. 5, c and d, peak 2; Table II), and a corresponding reduction of the neutral peak (Fig. 5, c and d, peak 1; Table II). These data indicate that activity of the α2,3 sialyltransferase in the transfected cell lines has resulted in the increased sialylation of the Galβ1,3GalNAc substrate. Further confirmation of the increase in sialylated core-1 comes from FACS® analysis using PNA lectin, which binds to the unsialylated disaccharide (Gillespie et al., 1993). Fig. 6 shows that, while the parental T47D (a) and T47D neo transfectant (b) bind the lectin, in the transfected clones PNA binding is totally absent (c and d). However, after removal of sialic acid with neuraminidase treatment, both the transfectants and the untransfected cells show equivalent binding of peanut lectin.

Figure 4. EM analysis of (a) MTSV1-7, (b) MTSV1-7 3STMYC, (c) T47D, and (d) T47D 3STMYC, showing that the structural morphology of the Golgi apparatus has not been obviously altered by overexpression of the α2,3 sialyltransferase. Both transfected and untransfected cell lines show well-stacked Golgi consisting of several cisternae.
As for the T47D series, the binding of PNA seen in the parental MTSV1-7 cell line and the puromycin transfectant (Fig. 6, e and f) was not evident in the MTSV1-7 transfectants (Fig. 6, g and h), but it could again be induced by neuraminidase treatment. Thus, the sialyltransferase was also active in the MTSV1-7 transfectants in increasing the level of sialylated core-1. To test whether the \( \alpha \)2,3 SAT (O) enzyme was affecting side chains that contain GlcNAc (Fig. 1), the radiolabeled MUC1 precipitates from the MTSV1-7 puro and MTSV1-7 3ST clones were analyzed for hexosamine content. Fig. 7 a shows the elution profiles of the puromycin clone, and Fig. 7 b shows the corresponding profile of one of the sialyltransferase transfectants. It can be clearly seen that, in the \( \alpha \)2,3 SAT (O) transfectant, there is a marked increase in the proportion of counts eluting in the first peak, which corresponds to galactosamine, compared to the second peak, which corresponds to glucosamine. This reduction in GlcNAc content clearly demonstrates a loss of chain branching and/or extension of the side chains on the MUC1 expressed by the MTSV1-7 transfectants.

**Intracellular Localization of the \( \alpha \)2,3 Sialyltransferase**

**Light Microscopy.** Immunofluorescence microscopy with the 9E10 mAb that recognizes the myc-tagged \( \alpha \)2,3 SAT (O) of the MTSV1-7 3ST and T47D 3ST cell lines gave the expected pattern of staining. The \( \alpha \)2,3 sialyltransferase gave a perinuclear staining, often located to one side of the nucleus, characteristic of Golgi apparatus staining (Fig. 8 a). No staining was observed elsewhere in the cell and staining of unfixed cells showed no positive surface staining (data not shown). To confirm the localization in the Golgi apparatus, the cells were double labeled using a polyclonal antibody to mannosidase II, a resident of the Golgi medial cisternae, and 9E10. Fig. 8 b shows that these enzymes are coexpressed, as shown by the yellow staining resulting from the overlapping of fluorescein and rhodamine second antibodies, thus confirming localization to the Golgi apparatus of the recombinantly expressed \( \alpha \)2,3 sialyltransferase.

**Immunoelectron Microscopy.** Thin-sections of pellets of the transfected MTSV1-7 and T47D cells were labeled with the myc antibody followed by goat anti–mouse coupled to gold particles. From visual observations of the immunoelectron micrographs of both cell types, the gold labeling appeared to be localized over the Golgi stack and was polarized to one side (Fig. 9, a and b). Since the structure of the Golgi apparatus was more clearly defined in MTSV1-7 cells and the expression of the transfected gene

**Table II. Relative Distribution of Radiolabeled Oligosaccharides of MUC1 from T47D, T47D neo, and T47D 3STMYC Cell Lines**

| Cell line       | Percentage of total counts applied |
|-----------------|-----------------------------------|
| T47D            | T47D neo | T47D 3STMYC 2 | T47D 3STMYC 3 |
| Peak 1 (neutral)| 48       | 42           | 23           | 17           |
| Peak 2 (mono sialo)| 46     | 50           | 61           | 73           |
| Peak 3 (di sialo)| 6       | 8            | 15           | 10           |
was used in double-labeling experiments (Fig. 9, c and d).

The relative distribution of the two enzymes was then determined by counting each size of gold particle across the Golgi stacks, using a method developed by T. Nilsson and C. Rabouille (personal communication). The distance between the beginning of the cis cisternae and the end of the trans compartment was divided into 10 equal fractions and converted to real distances (nm) using the magnification factor of the micrograph. The number of gold particles of each size, in each of the 10 fractions, was counted, converted to a percentage frequency, and plotted against distance as shown in Fig. 10. This method provides a more detailed outline of the distribution of the GalT and α2,3
SAT (O) across the Golgi apparatus, rather than in each cisternal profile, as the stack often consisted of more than three cisternae, which could not therefore be simplified to cis, medial, or trans.

The number of each size of gold particles in the TGN was also estimated, and the distribution in the Golgi apparatus and TGN is shown in Table III. Quantitation of the 5-nm gold particles showed that the GalT was localized late in the Golgi stack, corresponding to the trans cisternae, and in the TGN (Table III and Fig. 10) as previously reported (Rabouille et al., 1995). The larger (10-nm) gold particles identifying the α2,3 SAT (O) show that this enzyme is coexpressed in the trans cisternae but is equally well expressed earlier in the stack in an area probably corresponding to the medial cisternae. 26% of the α2,3 sialyltransferase was also found in the TGN.

Sequencing the cytoplasmic tail, transmembrane domain, and part of the stalk region of endogenous α2,3 sialyltransferase, prepared by PCR from T47D and MTSV1-7 (see Materials and Methods), showed that the sequence was identical to the transfected enzyme (data not shown). Thus, the transfected enzyme contains the same sequence that determines the residency of the endogenous α2,3 sialyltransferase in the Golgi apparatus.

Figure 8. Immunofluorescence microscopy of an MTSV1-7 3STMYC cell line. (a) MTSV1-7 3STMYC cells were fixed, permeabilized, and labeled with 9E10 for myc-tagged α2,3 SAT (O) (green). (b) MTSV1-7 3STMYC cells show double labeling of the same cells with an antibody against a resident Golgi marker, mannosidase II, such that α2,3 SAT (O) myc (green) and mannosidase II (red) show overlapping distribution (yellow) to a compact, juxtanuclear reticulum. Bar, 20 μm.
The α,2,3 sialyltransferase studied here is particularly relevant to the study of breast cancer since the activity of the enzyme is elevated in breast cancer cell lines (Brockhausen et al., 1995). Using in situ hybridization, we have also recently noted elevation of expression of the mRNA coding for the enzyme in some primary breast cancers, with the less differentiated tumors showing a higher expression (Burchell, J., and R. Poulsom, manuscript in preparation). The data presented here show that increasing the expression of the α,2,3 sialyltransferase in breast epithelial cell lines results in increased sialylation of the O-glycans added to the MUC1 glycoprotein, manifest as an increase in sialylated core-1 structures. Moreover, the enzyme may compete with the core-2 β1,6 GlcNAc transferase (when it is expressed) for the common core-1 substrate with the result that the GlcNAc content of the O-glycans is reduced (Fig. 7). Although the core-2 β1,6 GlcNAc T is absent in some breast cancer cell lines, in others such as MCF-7, mRNA and enzyme activity have been demonstrated (Brockhausen et al., 1995). Our results indicate that the increase in the α,2,3 SAT (O) activity seen in the breast cancers could still inhibit chain extension and influence the composition of the O-glycans added to a tumor-associated antigen, even when the core-2 β1,6 GlcNAc T is present and active. This in turn could affect the behavioral properties of the tumor cell. As a result of the complexity of the structure of the core-2-based oligosaccharides on MUC1 produced by MTSV1-7 (Lloyd et al., 1996), a detailed structural analysis of the side chains is necessary to prove unequivocally that the decrease in GlcNAc content is due to a reduction in core-2 branching rather than a reduction in chain extension from galactose in core-1.

Since in O-glycosylation sugars are added individually and sequentially, the position of an enzyme in the Golgi apparatus relative to other enzymes active in the synthesis of O-glycans will influence the final structure. The conclusion from the studies reported here is that the α,2,3 sialyltransferase is located in the medial and trans cisternae of the Golgi apparatus with some of this enzyme also being present in the TGN. To localize the protein by immunoelectron microscopy, the cDNA was tagged with an immunoreactive epitope (a sequence from the myc gene), and cells were transfected with the tagged gene. The inclusion of the myc epitope (10 amino acids) at the extreme COOH terminus (which in this case is in the lumen of the Golgi compartment) has previously been shown to have no effect on the localization of glycosyltransferases (Rabouille et al., 1995; Nilsson et al., 1993; Munro and Pelham, 1986). In our studies, the tag clearly had no obvious effect on the folding of the protein as the α,2,3 SAT (O) was highly active, not only in cell extracts (Table I) but also in the intact transfected cells, as shown by the increased sialylation of the MUC1 glycoprotein (Figs. 5, 6, and 7). Thus, anomalous localization because of misfolding is unlikely. In addition, the sequences that have been shown to be involved in the localization of glycosyltransferases were identical in the endogenous and transfected genes, making it likely that the coded proteins were directed to the same compartment of the Golgi apparatus. The location of the sialyltransferase to within more than one compartment of the Golgi apparatus supports the view that the Golgi cisternae are characterized by their different mixtures of these enzymes, not by discrete types (Nilsson et al., 1993; Rabouille et al., 1995).

There is a formal possibility that overexpression of a glycosyltransferase may alter the fine localization of the enzyme. This has been suggested from data localizing the β1,2...
N-acetyl glucosaminyltransferase (NAGT-1), where direct immunolocalization of the endogenous enzyme showed localization to the medial compartment of the Golgi apparatus (Dunphy et al., 1985), while the transfected tagged enzyme (fourfold increased expression) located to both the medial and trans compartments (Nilsson et al., 1993). It must be noted, however, that the sensitivity of the method used for quantitating the distribution throughout the Golgi apparatus can affect the outcome, and with an increase in the level of expression comes an increase in the sensitivity of detection. The data of Rabouille and colleagues argue against an effect of overexpression since the localization of a tagged N-linked α2,6 sialyltransferase was not altered when it was expressed at widely different levels (Rabouille et al., 1995). Even if overexpression of the enzyme does result in a broadening of the distribution in the Golgi apparatus, this would also be expected to occur when the endogenous enzyme is overexpressed, as it is in breast cancers by 8–10-fold. Thus, any overlap or competition with other glycosyltransferases such as α2,6 sialyltransferase in the trans compartment, the positioning of enzymes involved in mucin-type O-glycosylation has relied mainly on sucrose density gradient centrifugation (Chaney et al., 1989) or on following the glycosylation of marker proteins (Locke et al., 1992). The work presented here represents the first fine mapping of a sialyltransferase active specifically in O-glycosylation. Previous work by Locke and colleagues following the acquisition of oligosaccharides onto the M protein of Coronavirus and the effects of brefeldin A suggested that α2,3 SAT (O) is in an earlier compartment than the TGN (Locke et al., 1992; Lippincott-Schwartz et al., 1990). The results presented here support their interpretation of the data, as we place the α2,3 sialyltransferase in the trans and medial cisternae but also in the TGN. However, with the cloning of genes encoding the glycosyltransferases (Joziasse, 1992; Clausen and Bennett, 1996), it is becoming apparent that this group of enzymes is extremely complex and that there is a large family of sialyltransferases (Tsui, 1996) that may be resident in different Golgi cisternae. Thus, it becomes imperative to localize each specific enzyme directly, which can be achieved using tagged cDNA as described here, or by developing antibodies to the expressed recombinant enzymes.

There is some evidence to suggest that increased sialylation of glycoproteins may be involved in malignant progression, and this question can now be addressed more directly by manipulating the composition of the O-glycans synthesized by the tumor cells. In this context, it is significant that the product of the α2,3 SAT (O) enzyme, NeuAco2,3 Gal β1-3GalNAc, is the major ligand for sialoadhesin, a lectin expressed by macrophages (Kelm et al., 1994), and cells expressing MUC1 carrying this O-glycan are strongly bound by sialoadhesin (Crocker, P.R., personal communication). Whether increasing the level of this glycan affects the macrophage infiltrate in tumors is a specific question that can now be posed.

The change in glycosylation pattern seen in breast cancer also relates specifically to the studies on the MUC1 mucin as a potential target antigen in active specific immunotherapy of breast cancer. The extracellular domain of the MUC1 glycoprotein is made up largely of tandem repeats of 20 amino acids: 25–100 depending on the allele (Gendler et al., 1990), and each repeat contains potential glycosylation sites (Nishimori et al., 1994; Stadie et al., 1995). The antigenic profile of the mucin is therefore dramatically altered when the composition of the O-glycans added is changed from being core-2 based to the simpler, shorter, and more heavily sialylated glycans found on the tumor mucin. Indeed, both humoral and cellular responses to the MUC1 mucin have been observed in breast cancer patients, who show some specificity for the aberrantly glycosylated mucin expressed by the tumor cells (von Mensdorf-Pouilly et al., 1996; Magarian et al., 1993). Our findings show that it may be possible to produce the appropriate glycoform of the mucin in recombinant form by manipulating the glycosyltransferases in the producer cell. CHO cells, widely used for the production of recombinant proteins, do not in fact express the core-2 β1,6 GlcNAc T (Li et al., 1996) and have been reported to synthesize short O-glycans (Oheda et al., 1988). It may then be relatively simple to produce the MUC1 antigen in these cells with minimal manipulation. Furthermore, although the results presented here would support the hypothesis that the distributions of the core-2 β1,6 GlcNAc transferase and the α2,3 sialyltransferase show some overlap, it will be important to confirm this by detailed mapping of the position of the β1,6 GlcNAc transferase.

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