Extracellular sialyltransferase st6gal1 in breast tumor cell growth and invasiveness

Nitai C. Hait, Aparna Maiti, Rongrong Wu, Valerie L. Andersen, Chang-Chieh Hsu, Yun Wu, Digantkumar G. Chapla, Kazuaki Takabe, Michael E. Rusiniak, Wiam Bshara, Jianmin Zhang, Kelley W. Moremen, Joseph T. Y. Lau

INTRODUCTION

The sialyltransferase ST6GAL1, mediating the α2,6-sialylation of N-glycans, is prominently associated with human cancers, including ovarian, prostate, pancreatic, colon, lung, colorectal, pancreatic, gastric, breast cancers, and acute myeloid leukemia [1–3]. Elevated cancer ST6GAL1 is often correlated with high tumor grade, metastasis, and poorer patient prognosis [3–7]. Furthermore, ST6GAL1-mediated α2,6-linked sialylation has been implicated in activating PI3K/AKT, epidermal growth factor receptor (EGFR) in some cancers, and cell growth and proliferation [3, 8]. In addition, ST6GAL1-mediated sialylation modulates cell surface receptor function, including integrin, death receptor, TNFR1, and the promotion of epithelial to mesenchymal transition (EMT) and resistance to chemo- and radio- treatments [1, 3, 9, 10]. These and related studies have driven the idea that ST6GAL1 is a cancer-promoting factor, including breast cancer [11]. However, a few reports correlated better long-term patient survival among individuals with the most aggressive cancers and the highest cancer ST6GAL1 [12, 13]. This conundrum brings to question whether cancer ST6GAL1 overexpression is beneficial or ultimately detrimental to long-term patient outcomes and highlights the incomplete mechanistic understanding of how ST6GAL1 is involved in cancer progression. Canonically, functioning ST6GAL1 resides within the intracellular ER-Golgi secretory complex, where sialylation of nascent cell surface and secreted components occurs cell-autonomously. It is within the cell-autonomous context that ST6GAL1 involvement in cancer progression has been interpreted.

Aggressive breast cancer is a highly heterogeneous disease caused by a variety of distinct cell-intrinsic genetic alterations in mammary epithelial cells, leading to vastly heterogeneous disease manifestation in individual patients and predominantly affecting patient prognosis and treatment options [14]. Cancer cell-extrinsic mechanisms, which are poorly understood, are also believed to contribute to disease progression and the heterogeneous genetic mutations with diverse presentations. Elevated N-glycan signatures are present in the disseminated cancer cells in breast cancer...
patients [15], and heterogeneity in α2,6-linked sialic acids have been suggested to potentiate the invasion of aggressive breast cancer cells [11].

In addition to the canonical intracellular ER-Golgi locale, catalytically active ST6GAL1 is also present in the extracellular spaces and systemic circulation. The physiologic roles of these extracellular glycosyltransferases have received scant attention, but we and others have identified extracellular ST6GAL1 as a potent modifier of hematopoiesis, inflammatory cell production, B cell differentiation, and proliferation, and in the sialylation of the anti-inflammatory IgG [16–22]. Furthermore, ST6GAL1 is released into the extracellular milieu from cancer cells and healthy non-involved sources such as the liver [18, 23]. We hypothesize that ST6GAL1 action is the combined contribution of intracellular cell-native ST6GAL1 acting cell-autonomously and the extracellular ST6GAL1 acting non-cell-autonomously.

We confirm that cancer ST6GAL1 is variably expressed in triple-negative breast cancer (TNBC) clinical specimens that confer heterogeneous presence of α2,6-sialic acids on aggressive breast tumors. Breast cancer cell-native ST6GAL1 is associated with metastatic progression of aggressive breast cancer, based on online KM-plotter analysis, but we also observed that high expression of cancer ST6GAL1 mRNAs is associated with a favorable relapse-free survival (RFS) in TNBC patients. Cancer cell-native ST6GAL1 is preferentially released in association with exosome-like particles and smaller exomer-like particles. Exosome-like particles from ST6GAL1-high cancer cells elicit increased cell proliferation and invasiveness in cancer cells with low native ST6GAL1 expression. Our data further suggest the presence of undefined cofactors within the cancer exosomes, and these cofactors potentiate the action of extracellular ST6GAL1. Extracellular ST6GAL1 and the undefined exosome cofactors can compensate cell-intrinsic ST6GAL1 expression and potentiate aggressive cancer cell growth and proliferation, cancer cell invasion, EMT, CSCs (cancer stem cells) transcription cofactors can compensate cell-intrinsic ST6GAL1 expression and confer heterogeneous presence of α2,6-sialic acids on aggres- sive breast cancer, based on online KM-plotter analysis, but we also observed that high expression of cancer ST6GAL1 mRNAs is associated with a favorable relapse-free survival (RFS) in TNBC patients. Cancer cell-native ST6GAL1 is preferentially released in association with exosome-like particles and smaller exomer-like particles. Exosome-like particles from ST6GAL1-high cancer cells elicit increased cell proliferation and invasiveness in cancer cells with low native ST6GAL1 expression. Our data further suggest the presence of undefined cofactors within the cancer exosomes, and these cofactors potentiate the action of extracellular ST6GAL1. Extracellular ST6GAL1 and the undefined exosome cofactors can compensate cell-intrinsic ST6GAL1 expression and potentiate aggressive cancer cell growth and proliferation, cancer cell invasion, EMT, CSCs (cancer stem cells) transcription factors, and serum-withdrawal stress-mediated cancer cell apoptosis. Our data highlights a previously unappreciated, non-cell-autonomous mechanism linking extracellular ST6GAL1 in cancer pathobiology.

**MATERIALS AND METHODS**

**Cells and reagents**

MCF7, MDA-MB-231 (MDA-231), BT-474, T-47D, ZR-75-1, E0771, and BT-549 cells were obtained from ATCC, Virginia, USA. LM-2,4 cells were kindly provided by Dr. John Ebos, Roswell Park Comprehensive Cancer Center (RPCCNY, USA). 4T1 parental and metastatic clone 4T1,2 cells were donated by Prof. Cheryl L. Jorcyk, Boise State University (ID, USA). Breast cancer metastatic clone MDA-231/BR was obtained from Dr. Patricia S. Steeg, National Cancer Institute, Bethesda, MD, USA. MCF7 cells were cultured in phenol red-free improved minimum essential medium supplemented with 0.25% glucose and 10% fetal bovine serum (FBS). BT-474, BT-549, 4T1, 4T1,2, and E0771 cells were grown in RPMI medium with 10% FBS. T-47D, ZR-75-1, MDA-231, LM-2,4, and MDA-231/BR cells were grown in DMEM medium with 10% FBS. All cell culture media were supplemented with 100 units/ml penicillin and 100 µg/ml streptomycin (24, 25). All experiments were performed during the logarithmic growth phase, and cell lines were passaged for no more than 6 weeks. All cell lines used here are certified in our laboratory to be mycoplasma-free. Cells were plated 24 h before treatment. Recombinant rat secretory ST6GAL1 enzyme (rST6GAL1) [24] was produced by Dr. Kelley W. Moremen, University of Georgia, Georgia, USA. CMP-sialic acid (CMP-Sia) (MilliporeSigma, Massachusetts, USA), DAPI (BioLegend, CA, USA; 0.5 µg/ml), Crystal Violet (MilliporeSigma, Massachusetts, USA), Matrigel (Biosciences, CA, USA), and Cell Counting Kit-8 (CCK-8), WST-8 was from Dojindo Molecular Technologies, MD, USA. shRNA targeting mouse ST6GAL1 (shST6GAL1 #1), established and validated in Lau Laboratory; shST6GAL1 #2, Sigma, cat # TRCN000018819; sequence CCGCGCAGGATGTTAATGTTCACATCGGAGATGTTATCATGATGTTT; shST6GAL1 #3, Sigma, cat # TRCN000018821, sequence CCGCGCAGGATGTTAATGTTCACATCGGAGATGTTATCATGATGTTT; human ST6GAL1 (shST6GAL1 #1, Sigma, cat # TRCN0000035432, sequence CCGCGCTGTGTGCTACTATACAAGCTTGAAGAGATGTTATCATGATGTTT; shST6GAL1 #2, Sigma, cat # TRCN000035429, sequence CCGCGCAGGATGTTAATGTTCACATCGGAGATGTTATCATGATGTTT; and shControl (pLKO.1) were used to knockdown ST6GAL1. Lipofectamine 3000 transfection reagent (Invitrogen, MA, USA) was used for the transfection of RNA plasmids. Successful knockdown was confirmed by immunoblotting using anti-ST6GAL1 goat polyclonal antibody (R&D Systems, MN, USA; cat. # AF5924) and qPCR analysis.

**Immunoblotting**

Cell lysates were used for Western blot analysis as previously described [25]. Briefly, cells were washed twice with the ice-cold 1X PBS and lysed by probe-sonication in the lysis buffer (20 mM Tris (pH 7.4), 20% glycerol, 150 mM NaCl, 0.5% NP-40, 5 mM sodium orthovanadate, 40 mM β-glycerophosphate, 15 mM NaF, 0.5 mM 4-deoxy-D-glucose, and protease inhibitor cocktail (Sigma)). Unbroken cells were measured by centrifugation at 700 x g for 10 min at 4°C. Protein concentrations were determined by the Bradford method (Bio-Rad, CA, USA). A total of 25 µg of protein was loaded per lane and separated by 10% SDS-PAGE and transferred to nitrocellulose membranes. Membranes were blocked with 5% blocking solution in 1X TBST buffer, and then membranes were incubated at room temperature and followed by incubation (1 h at room temperature) with the rabbit or mouse, or goat secondary antibodies (1: 5,000 in TBST with 1% blocking solution). Membranes were washed thoroughly at room temperature and developed with enhanced chemiluminescence (ECL western blotting substrate (#32106; Pierce/Thermo Fisher Scientific, MA, USA). Western blot images were obtained by exposing the membranes to X-ray films (Thermo Fisher Scientific, MA, USA). The following primary antibodies were used for western blot analysis: β-tubulin, Cytochrome c, Caspase 3 cleaved protein, PARP, N-cadherin, E-cadherin. Snap are from Cell Signaling Technology, MA, USA; TGSTG101, CD63, and GM130 are from Santa Cruz Biotechnology, CA, USA. In addition, the following secondary antibodies were used for Western blot analysis: peroxidase-conjugated affghage goat anti-rabbit IgG (#111-035-045) and goat anti-mouse IgG (#115-035-062) (both from Jackson ImmunoResearch Laboratories, Inc, PA, USA), and anti-GOat IgG HRP-conjugated antibody (#HAF109, R&D Systems, Inc, MN, USA).

**In vitro sialyltransferase assay**

Sialyltransferase assays using cell lysates were carried out as described previously [18]. Briefly, 1.4 µl of cell lysates were mixed with Acceptor A (Galβ1,4GlcNAc-O-Bn; Toronto Research Chemicals Inc, Ontario, Canada) for ST6GAL1 activity or Acceptor B (Galβ1,3GlcNAc-O-Bn; Toronto Research Chemicals Inc, Ontario, Canada) for ST6GAL3 enzyme activity, the substrate CMP-Siaβ1,3Galβ1,3GlcNAc-O-Suc (25 mM sodium cacodylate pH 6.5, 0.3% Triton X-100). The enzymatic reaction was performed at 37°C for 1 h. After the enzymatic reaction, the enzyme product (Siaβ1,3Galβ1,3GlcNAc-O-Suc) was separated by using a Waters Sep-Pak® 18 column cartridge. Liquid scintillation counting of the methanol flow-through counts radioactivity from reacted product exclusively. Enzyme activity was calculated as fmol product/min/mg of cell extract protein.

**Real-time PCR**

Quantitative real-time PCR (qPCR) analysis was performed as described before [25]. Briefly, total RNA was prepared with Trizol (Life Technologies, CA, USA). DNA contamination was removed from RNA with DNase treatment (Qiagen, MD, USA). RNA (2 µg) was reverse transcribed with the high-capacity cDNA Archive Kit (Life Technologies). cDNAs were diluted 10-fold (target genes) and 100-fold (house-keeping gene Glyceraldehyde 3-phosphate dehydrogenase (GAPDH), and amplified with SYBR Green qPCR on CFX96 cytos (Bio-Rad). Gene expression levels were calculated by the 2-ΔΔCT method [26], and for some experiments, 2-ΔΔCT method [27] were normalized to GAPDH expression. Following real-time PCR, primers are listed below in this manuscript. Human-ST6GAL1-F: 5′-CGAAGATCAGTGAGTGAAACAG-3′; human-ST6GAL1-R: 5′-CCAACTCTAGCGACAGATCG-3′; mouse-ST6GAL1-F: 5′-CTGCGTCCTACGGACGACAGGATCTTCTGGTCGAGGT-3′; mouse-ST6GAL1-R: 5′-TCCACTCTCCAGGAGAGGGA-3′; mouse-GAPDH-F: 5′-CTGACCCACTTCTGGACGAGGAGGGA-3′; mouse-GAPDH-R: 5′-ATGTTCACTTGGAGACGCCCG-3′; mouse-GAPDH-F: 5′-TCACCATCCTCGGAGAGGGA-3′; mouse-GAPDH-R: 5′-TCGAGTGTTCACTTGGAGACGCCCG-3′; and mouse-GAPDH-F: 5′-TCACCATCCTCGGAGAGGGA-3′; mouse-GAPDH-R: 5′-TCGAGTGTTCACTTGGAGACGCCCG-3′.
mouse-GAPDH-R: 5′-GACCTGTGCAATCTTGTGAGC-3′; human-Twist-1-F: 5′-CCTCAGAGGTGTGGCTAAT-3′, human-Twist-1-R: 5′-CCCGATTTATTTTCCAAAGTTT-3′; human-Slug-R: 5′-CAAGCTGACACACCATACGTG-3′; Slc2a1-F: 5′-CTAGGATCTTGGTTGGT-3′, human-ALDH1-F: 5′-AACTCTCTCTACTGCTCTCCAG-3′, human-ALDH1-R: 5′-GTCACCTCTTACAGGTCTTCC-3′; human-c-Myc-F: 5′-CACGGAAAATTCCTGGGTCACG-3′, human-c-Myc-R: 5′-GCAAGGAGAAGCCTGATGTCG-3′, human-Nanog-F: 5′-CCCTGCTGCTGCGTGAAGAAA-3′, human-Nanog-R: 5′-AGG GCT GTC CTT GAG AAG CAG C-3′; human-Oct4-F: 5′-CCCTGAGAACAGAAGATGTA-3′, human-Oct4-R: 5′-CCGGAGCTTCAACCAGTTCT-3′; human-E-Cadherin-F: 5′-CTGAGGATCCAGCGATTTGAGC-3′, human-E-Cadherin-R: 5′-GAGCCGGCCGGCAGGAAGA-3′, human-Nan-Cadherin-R: 5′-TCAAGCCGCTTGAGAAGCTT-3′, human-Nan-Cadherin-F: 5′-ATGGCACTATCTGGTAAGCTC-3′; human-Snail-F: 5′-GCCATTTAACATAAATGGAAGGG-3′, human-Snail-R: 5′-GAATATTGTTGAGACACATCG-3′; human-CD44-F: 5′-AGAAGATGTGTCGCGGGAGAA-3′, human-CD44-R: 5′-AAATGACCATATTCTT GAGA-3′, human-Vimentin-F: 5′-AGTCCACTGATTCCGGAGAC-3′, human-Vimentin-R: 5′-CATTCCAGGATGTTGCCGTC-3′, human-Sox2-F: 5′-GAGC TTGGACGGAAGTTCGTG-3′, human-Sox2-R: 5′-GCAGAGACCGCTCCGTA-3′.

**Isolation of extracellular vesicles (EVs) by ultracentrifugation**

As previously described, EVs were isolated from the cancer cell-conditioned medium with minor modifications [23, 26]. Briefly, confluent breast cancer cells were cultured in serum-free media for 48 h. The supernatants were collected and centrifuged at 500 x g for 20 min at 4 °C to discard the cells. The supernatants were further centrifuged at 10,000 x g for 30 min at 4 °C to discard cell debris and apopotic bodies. Finally, the supernatants were ultra-centrifuged at 100,000 x g for 90 min at 4 °C to precipitate the EVs. The EVs were suspended with 20 volumes of ice-cold buffer (25 mM HEPES, pH 7.0 in 0.1% Tween 20) and washed by centrifuging again at 100,000 x g for 90 min at 4 °C. The washed pellet (1/10th volume of original supernatant) was designated as exosome-like particles. The protein concentration of the nanoparticles was measured and analyzed by Western blot analysis with EV-specific marker antibodies (positive markers: TSG-101 and CD63; negative markers: GM130) and analyzed by NIH ImageJ software.

**Nanoparticle tracking analysis (NTA)**

As previously described, [29] exosome-like particles were analyzed by NTA to measure the size, size distribution, and concentration of exosomes using a NanoSight LM10 system (NanoSight, Malvern Instruments, Worcestershire, UK) and Transmission Electron Microscopy (TEM).

**Transmission electron microscopy (TEM)**

TEM analysis of the exosome-like particles was carried out as previously described [23]. The exosome-like particles fixed with 2.5% glutaraldehyde in 0.1 M sodium cacodylate buffer. Samples were incubated on grids and negatively stained with 2% uranyl acetate. Imaging was performed on a Philips/FEI T-12 transmission electron microscope and on grids and negatively stained with 2% uranyl acetate. Imaging was performed on a Philips/FEI T-12 transmission electron microscope and on grids and negatively stained with 2% uranyl acetate. Imaging was performed on a Philips/FEI T-12 transmission electron microscope and on grids and negatively stained with 2% uranyl acetate. Imaging was performed on a Philips/FEI T-12 transmission electron microscope and on grids and negatively stained with 2% uranyl acetate.

**3D invasion assay**

Cancer cell spheroid assays were performed in a 3D cell culture setting with modifications [31–34]. Briefly, 20,000 cells were cultured in DMEM F12 medium (Gibco-Thermo Fisher Scientific, NY, USA) supplemented with 20 ng/ml each of human EGF (Pepro Tech, NJ, USA), human FGF, and 1× B27 (Gibco-Thermo Fisher Scientific) in ultra-low attachment 24-wells plates (VWR) for 24 h in the presence of 1–2 µg/ml rST6G, CMP-Sia, or equivalent vehicle controls. In addition, for some experiments, cells were grown in the presence of 1–2 µg self- or source cell exosome-like particles with or without rST6G, as indicated in the figure/legends. Then, cells were mixed thoroughly with the BD Matrigel (Biosciences, CA, USA) in the DMEM F12-Growth Factors medium and plated on ultra-low attachment plates. Finally, tumor cell spheroids were treated with the rST6G or exosome-like particles, or equivalent vehicle control, as indicated in the figure/legends for another three days. 3D spheroids in 5 randomly selected fields were imaged at ×10 magnification using a Nikon Eclipse Ti-U Inverted Contrast Phase Microscope (Nikon Instruments Inc., NY, USA). The aggressive tumor cell invasion phenotypes were analyzed using ImageJ software (National Institute of Health, MD, USA). 3D spheroids invasion was quantified as the average number of invadopodia/protrusions per spheroid (n = 5–10). In addition, for some experiments, the spheroids invasion was quantified as the average length (µm) of invadopodia/protrusions per spheroid (n = 5–10).

**Immunofluorescence**

Cells are grown onto 13 mm sterile glass slides overnight with 50% confluences in a 5% FBS-containing medium. For some experiments, the cells were fixed with 4% paraformaldehyde for 10 min at 37 °C; before paraformaldehyde-fixing, cells were washed twice with serum-free medium and treated with exosome-like particles ± rST6G or equivalent vehicle control, as indicated in the figure/legends. First, slides were permeabilized with 0.1% Triton X-100 (Sigma) for 5 min at room temperature (RT). Next, slides were washed with 1× PBS and blocked with 1% BSA (Sigma) for one hour at RT. After brief washing with PBS-T (0.1% Tween 20), slides were incubated with Direct FITC-conjugated NanoSight (NanoSight, Malvern, UK) and Transmission Electron Microscopy (TEM).}

**Cell proliferation assay**

Cell proliferation assays were carried out as described [30]. Briefly, the day before treatment, cells were cultured in full-serum medium on the 96-well tissue culture plate. The next day, cells were washed twice in a serum-free medium, and treatments were performed, as indicated in the figures/legends. Cell growth was also measured by adding WST-8 reagent and incubating at 37 °C for 1 h; absorbance was measured at 450 nm with background subtraction.

**Bioinformatics analysis**

Transcriptome data was acquired from The Cancer Genome Atlas (TCGA) breast cancer cohort through cBioportal [38]. ST6GAL1 mRNA expression levels were compared between breast cancer (TCGA, n = 1078) and normal healthy breast tissues (n = 52). ST6GAL1 mRNA expression data were analyzed by cBioPortal [38].
Elevated ST6GAL1 expression is associated with activating gene networks that promote a metastatic phenotype. A boxplot shows a high expression score of the ST6GAL1 gene in normal adjacent breast tissues (n = 117) vs. breast cancer patients of TCGA breast cancer cohort (n = 979) (A). Student’s t-test, p = 0.031. Boxplots of the ST6GAL1 high expression score by immunohistochemistry (IHC) determined subtype in the TCGA breast cancer (B) and Nottingham pathological grades are shown for the METABRIC cohort (C). All boxplots are Tukey type, and the boxes depict medians and inter-quartile ranges. One-way ANOVA and Tukey’s tests were used to calculate p-values. Box plots of the intratumor heterogeneity [57, 58] in the TCGA breast cancer cohort by low and high ST6GAL1 expression scores (D). Mann–Whitney U test and Kruskal–Wallis tests were used to calculate the p-value. Median cut-off was used to divide two groups. IHC of ST6GAL1; representative (n = 3). E TNBC tissue section IHC with anti-ST6GAL1 antibodies are shown (red arrow higher and green arrow lower ST6GAL1 expression, respectively). Gene Set Enrichment Analysis (GSEA) of high ST6GAL1 in the TCGA breast cancer cohort revealed enrichment in the Hedgehog (F), EMT (G), and Hypoxia (H) pathways. The normalized enrichment score (NES) and false discovery rate (FDR) values for TCGA-BRCA cohort are: Hedgehog, NES = 1.467799; FDR = 0.161496; EMT, NES = 1.458644; FDR = 0.160286; and Hypoxia, NES = 1.622797, FDR = 0.085881. FDR of 0.25 was used as the statistical significance of GSEA. A median cut-off was used to divide two groups (high vs. low ST6GAL1). I The Kaplan–Meier estimates the probability of relapse-free survival (RFS) by mRNA of TNBC patients. Analysis was provided by using the online KM-plotter [60].

Gene Set Enrichment Analysis (GSEA) of hallmark gene sets for metastasis
Gene Set Enrichment Analysis (GSEA) was performed using software provided by the Broad Institute (http://software.broadinstitute.org/gsea/index.jsp) [41]. A collection of annotated gene sets for use with GSEA software can be found in the Molecular Signatures Database (MSigDB) (http://software.broadinstitute.org/gsea/msigdb). The result of the GSEA is the normalized enrichment score (NES), which reflects the degree to which a gene set is overrepresented in the top or bottom of a ranked list of genes. Statistical analyses were performed using R software (http://www.r-project.org). Statistical significance was defined by a false discovery rate (FDR) of less than 25%, as GSEA software recommended.

Statistical analysis
Quantitative data are presented as mean ± SE of three independent replicates. We have used a one-way ANOVA test for datasets containing multiple group comparisons and Tukey’s post hoc test for the family-wise error rate comparison. Association between variables (e.g., gene expression and mutation load scores) was determined using the Mann–Whitney U test. Kaplan–Meier method with log-rank test was used to compare survival curves between groups. All statistical tests were performed using the GraphPad Prism 8 software. P < 0.05 was considered statistically significant (*P < 0.05, **P < 0.01, and ***P < 0.001).

RESULTS
ST6GAL1 is heterogeneously expressed and elevated in aggressive human breast cancers
TCGA, one of two gene expression databases with large breast cancer patient cohorts, was interrogated. Even though ST6GAL1 transcript abundance was diminished overall in breast cancer patients compared to matched controls (Fig. 1A), TNBC aggressive cancers have elevated ST6GAL1 mRNA compared to HER2+ and ER+ subtypes (Fig. 1B). In addition, METABARIC cohort analysis validated that ST6GAL1 expression was significantly elevated in TNBC and HER2+ compared with the ER+ subtype (Supplemental Fig. S1A).

Importantly, elevated ST6GAL1 mRNA was significantly associated with a higher tumor grade in breast cancer patients (Fig. 1C), even though ST6GAL1 expression does not appear correlated with the aggressive stage of breast cancer (Supplemental Fig. S1B, C). Importantly, elevated ST6GAL1 mRNA is associated with intratumor heterogeneity in breast cancer patients (Fig. 1D). When TNBC clinical specimens were examined by immunohistochemistry (IHC) using anti-ST6GAL1, we observed heterogeneous ST6GAL1 protein expression (Fig. 1E), similar to published results in other clinical cases [18, 42].

Overall, analysis using TCGA (Fig. 1F–H) and METABRIC cohorts (Supplemental Fig. S1D–F) revealed that elevated ST6GAL1 is
associated with enrichment in gene networks associated with cancer stemness (Hedgehog), epithelial-mesenchymal transition (EMT), and hypoxia pathways. The data agree with the idea that ST6GAL1 is linked to the aggressive progression of breast cancers. Paradoxically in TNBC patients, an apparently opposing trend was observed, that higher ST6GAL1 expression is also associated with better relapse-free survival (RFS) (Fig. 1).

**ST6GAL1 is heterogeneously expressed in breast tumor cells**

ST6GAL1 is a catalytically active enzyme that constructs the α2,6-sialic acid moiety on N-linked glycans of glycoproteins. To gain further insight into how ST6GAL1 expression is regulated that ultimately affects N-glycan sialylation in cancer, we examined an array of breast tumor cell lines (ER+ and TNBC) for ST6GAL1 expression on multiple levels of mRNA, protein, enzymatic activity, and by the abundance of the cognate enzymatic product, the α2,6-linked sialic acids on cell surface glycans (Fig. 2 panel A, B, C, D, E, respectively). Published data already suggested that various tumor cell lines, including breast cancer cell lines, have variable levels of ST6GAL1 protein [9, 11, 18].

Here, we examined an array of breast tumor cell lines (ER+ and TNBC). Two molecular forms of ST6GAL1 are generally observed, a larger 50 kDa and a smaller 42 kDa forms presumptively representing, respectively, the full-length and the smaller soluble catalytic domain is known to be proteolytically released from its N-terminal cytosolic and transmembrane anchor [18]. Among the panel of breast cancer cells examined, the human cells had the larger 50 kDa form exclusively (Fig. 2B). The observed electrophoretic mobility of the largest ST6GAL1 form in the three mouse cell lines examined (4T1, 4T1.2, and E0771) appeared somewhat smaller than the corresponding largest ST6GAL1 from the human cells. Presently, the molecular basis for this inter-species difference in electrophoretic mobility is not known, and presumably might be due in part to glycosylation differences on the ST6GAL1 protein.

Figure 2 also illustrates that the levels of ST6GAL1 mRNA and protein were also highly variable within the cell line panel. Enzymatic assays were performed to measure the transfer of sialic acid from CMP-sialic acid to form Sia α2,6 joined to Gal (β1,4)GlcNac-R (Fig. 2C). While there is general concordance between high ST6GAL1 protein with high ST6GAL1 catalytic activity amongst the cell lines examined, some notable differences were observed. Among the human cell panel, BT-474 had moderate ST6GAL1 protein presence but low detectable enzymatic activity. The mouse 4T1 and its metastatic variant 4T1.2 had roughly equivalent ST6GAL1 protein levels, but 4T1.2 had drastically less enzymatic activity than 4T1. The most striking discordance between protein and activity levels was in E0771, where high enzymatic activity levels but modest ST6GAL1 protein were observed. Moreover, MDA-231/BR also had the highest measured enzymatic activity but modest ST6GAL1 protein (compare Fig. 2A and C).

Cell surface α2,6-Sia structures were interrogated using SNA, the lectin from *Sambucus nigra* (Fig. 2D and E). In general, the observed cell-surface SNA reactivity aligned with the level of cellular ST6GAL1 mRNA and protein expression. Most striking, however, is the variable and heterogenous SNA reactivity amongst cells of the same strain. This is most pronounced in the lung-metastatic variant of MDA-231 cells, LM-2-4, but present in all cell lines examined, where only a subset of cells was SNA+ (Fig. 2D, E). ST6GAL1 mRNA also did not correlate entirely with the observed expression on the protein level (Fig. 2A). Notable are ZR75-1 and LM-2-4; both lines had intermediate mRNA levels but little to no detectable ST6GAL1 protein or enzymatic activity. These observations are consistent with the published data of human lymphoblastoma cell lines [18]. Furthermore, in agreement with the human breast cancer clinical cases, we observed that human breast tumor cells have very low levels (normalized 2^Δ-ΔCT) of ST6GAL1 transcripts at the steady-state culture conditions. Assuming that human and mouse primers for the...
Fig. 2  Functional ST6GAL1 is variably expressed in breast cancer cells. A mRNA levels of ST6GAL1 were determined from ER+ and TNBC human and mouse breast cancer cells, as indicated by quantitative real-time PCR (qPCR) and normalized to GAPDH (2−ΔΔCt). N = 3, data are means ± s.e. B Total cell lysates from separate cell cultures (A) were used for Western blot analysis with antibodies against ST6GAL1. β-tubulin was used as a specific marker for cytosol and to show equal loading and transfer. C As indicated, an equal amount of proteins from cell lysates (B) was used for the ST6GAL1 enzyme assay, as previously mentioned [65]. ST6GAL1 activity is presented as fmol/min/mg protein; enzyme assays were performed in triplicates. Data are mean ± s.e. D Human and mouse representative ER+ and TNBC cell lines, as indicated, were immunostained with FITC- (green) labeled SNA lectin and DAPI (blue) for the nuclei before fixing. During fluorescent microscopy, exposure time and weighting for both DAPI and FITC fluorescence were kept consistent between samples. Representative images (N = 4) were shown on a scale bar of 50 µm. E Images were processed in ImageJ; background subtraction and MFI per cell calculations were carried out using the same parameters for each condition, and mean fluorescence intensity (MFI) per cell was shown with mean ± s.d. for four fields of view.

ST6GAL1 are similarly efficient; mouse TNBC 4T1 and breast cancer bone metastatic 4T1.2 cells have higher ST6GAL1 transcripts than ER+ E0771 cells (Fig. 2A, right panel).

Overall, the data describe highly variable expression of ST6GAL1 among breast cancer cell lines but also discordance between measurable enzymatic activity, ST6GAL1 protein, and mRNA levels among cells lines might be attributed to a poorly understood and largely unexplored complex differential kinetics of gene expression, protein translation, and longevity of the ST6GAL1 protein within the cells. As much of the intracellular ST6GAL1 is continually released into the systemic space, the levels of intracellular protein or measurable enzymatic activity might be highly dynamic and variable among different cells. Additional unknown factors, such as modifiers of intracellular ST6GAL1 mRNA and/or protein longevity, might also contribute.

Transient knockdown of ST6GAL1 reduced breast tumor cell growth and invasiveness

During malignant progression, invasive cancer cells proliferate and penetrate the basement membrane by forming invadopodia structures that are enriched with actin filaments, adhesion proteins, and proteinases for tumor cell motility and degradation of the extracellular matrix (ECM) [43].

To assess how ST6GAL1 expression might affect invasive cancer growth, a 3D spheroid culture method that closely mimics the in vivo environment, including a hypoxic tumor core, was used [33]. shRNA targeting was used to achieve significant knockdowns in several mouse and human breast cancer cell lines. The results for 4T1.2, E0771, and BT-549 are summarized. Cell growth and invasive characteristics were monitored in the shRNA knockdown cells (shST6GAL1) compared to cells transfected with random non-ST6GAL1 mRNA (shControl). Figure 3 panel A shows that shRNA knockdown of 4T1.2 cells resulted in an almost complete reduction of ST6GAL1 mRNA and protein levels accompanied by dramatically reduced cell growth and almost complete attenuation invadopodia formation (Fig. 3 panel B, C, respectively). Supplemental Fig. S2 panel A-E shows the results for E0771 cells. Three distinct sets of guide shRNAs (shST6GAL1#1, shST6GAL1#2, and shST6GAL1#3) targeting different sites of the ST6GAL1 mRNA sequence were used. All were effective in decreasing ST6GAL1 mRNA expression (Supplemental Fig. S2 panel A, C). shST6GAL1#1 was evaluated to be highly effective in attenuation of cell proliferation; shST6GAL1#2 and #3 had attenuated cell growth, although they were not as effective as #1 (Supplemental Fig. S2 panel B, E, respectively). ST6GAL1 protein levels were significantly decreased in #2 and #3 (Supplemental Fig. S2 panel D). The invasion was not measured in E0771 cells since they do not form invasive 3D spheroids in our cell culture. Figure 3D–F panels show BT-549 cells transfected with shST6GAL1#1 and shST6GAL1#2 guides or random control sequence (shControl). Both #1 and #2 gRNAs effectively diminished ST6GAL1 mRNA and protein levels, compared to shControl, and concomitantly reduced cell growth and invadopodia formation (Fig. 3D, E, and F, respectively). For the most part, the decrease of ST6GAL1 mRNA in the knockdown cells generally parallels decreased intracellular ST6GAL1 protein levels. However, as mentioned earlier, there is a complex differential kinetics of gene expression, protein translation, and longevity of the ST6GAL1 protein within the cells that are poorly understood and poorly studied. Therefore, intracellular ST6GAL1 may be a transient pool that does not truly reflect the effectiveness of the knockdown.

Breast cancer cells release ST6GAL1 in exosome-like vesicles

Cancer cells release heterogenic extracellular vesicular small particles such as exosome (50–120 nm) and exosomer (<50 nm) [23]. An enrichment of certain sialoglycoprotein and N-glycans [44], lectins [45], and neuraminidase [46] have been identified in exosomes, and adhesiveness of exosome to target cells may be mediated at least in part by exosome glycans and carbohydrate-binding receptors [47]. In addition, colon cancer exosomes encapsulating ST6GAL1 that can be delivered to target cells and involved in cell signaling have been reported recently [23]. Here, we examine whether breast cancer cells also encapsulate functional ST6GAL1 into extracellular vesicles (EVs).

Exosome-like vesicles were isolated from confluent monolayers of 4T1.2 breast cancer cells grown in a serum depleted medium for 48 h. The supernatants were centrifuged at 500 × g, 10,000 × g, and 100,000 × g, as previously described [23, 44]. 100,000 × g pellets were strongly enriched for EVs markers, CD63, Tsg101 but did not express the cis-Golgi matrix protein GM130 and mitochondiral localized protein Cytochrome c (Fig. 4A), as expected for an EV preparation without substantial contamination by cytosolic or mitochondrial protein [48]. 4T1.2 EVs carry ST6GAL1 as well as at least one other sialytransferase ST3GAL6. shST6GAL1#1 knockdown resulted in a ~4-fold decrease of ST6GAL1 in exosomes, but a negligible change in a2,3-sialytransferase activity, most likely ST3GAL6, measured as normalized to EV protein concentration (Fig. 4B). Representative size distributions measured by nanoparticle tracking analysis (NTA) and transmission electron microscopy (TEM) images of EVs from shControl and shST6GAL1 are also shown (Fig. 4C, D). Supplementary Figure S3A–F shows size distributions of exosomes released by ER+ (T47D, ZR-75-1, E0771) and TNBC (BT-549, MDA-231, and 4T1.2) cancer cells. In general, the EVs are a heterogeneous population in the range between 50 and 400 nm particle sizes. The maximum of the major peak ranged between 86 and 128 nm of EVs in the six breast cancer cell lines tested. Concentrations of EVs are also varied based on the cell of origin. A cell line-dependent difference in nanoparticle secretion is similar to what has been previously reported [49].

ST6GAL1 knockdown in 4T1.2 cells resulted in strikingly reduced ST6GAL1 in exosomes (see Fig. 4A). The morphology of the exosomes measured by TEM was similar between shRNA knockdown and shControl samples (Fig. 4C, D). There was a slight reduction of exosomes concentration measured by NTA and exosomal marker proteins enrichment (e.g., Tsg101 and CD63) in shST6GAL1 compared with shControl cells. Overall, our observations point to breast cancer cells packaging ST6GAL1 into exosome-like structures that are released into the extracellular milieu. The level of ST6GAL1 packaged and released within exosome-like particles varies depending on the breast cancer cells in our panel.
Cancer exosome-like particles potently amplify extrinsic ST6GAL1 to boost proliferation and invasiveness of breast cancer cells ex vivo

ST6GAL1 packaged within cancer exosomes can cell-surface sialylated and elicit signaling in recipient target cells [23]. Here, we demonstrate that extracellular ST6GAL1 elicits definitive phenotypic consequences in the recipient cells in the forms of enhanced cancer cell proliferation, growth, and invasive behavior.

Human TNBC breast cancer cells MDA-231, natively ST6GAL1-low and cell-surface SNA-dim, were used as targets. Adding recombinant ST6GAL1 (rST6G) but not self-exosomes to the culture medium enhanced MDA-231 cell growth; the addition of...
Natively expressing ST6GAL1 is involved in breast cancer cell growth and invasiveness. B Mouse breast cancer bone metastatic 4T1.2 cells were transfected with a validated shST6GAL1 #1 and shControl for 24 h with Lipofectamine P3000 reagents (Invitrogen), according to the company's instruction. 5000 cells transfected with shControl and shST6GAL1 were cultured in the 24-well tissue culture plate in the medium containing 5% serum for another 48 h and 72 h, and cell proliferation was determined with WST-8 reagent. Normalized A450 nm readings were plotted (n = 3). Data are means ± s.e., Student's t-test, **p < 0.001. A duplicate 48 h culture of 4T1.2 cells was used for Crystal Violet staining (B, lower panel). Representative live-cell images are shown for shControl and shST6GAL1. A Duplicate culture of 4T1.2 cells was used for SYBR-Green-qPCR and protein analyses for the ST6GAL1 gene. ST6GAL1 mRNA levels from the shControl and shST6GAL1 samples were normalized with the house-keeping gene GAPDH, and the normalized ST6GAL1 levels were calculated using the Delta-Delta Ct method, N = 3, data are ±s.e., Student's t-test, p < 0.001. A representative blot was shown for ST6GAL1 western blot analysis (N = 3), and GAPDH was used for housekeeping control for equal loading (A, right panels). (C, upper panels) 20,000 4T1.2 cells transfected with shST6GAL1 vs. shControl were plated on 12-well low-adherent tissue culture plates mixed with growth factor reduced Matrigel for 72 h to obtain aggressive tumor-cell invasion in a 3D cell culture setting. The experiments were repeated three times, and representative phase-contrast images (×10 magnification) of tumor-cell invasions are shown for each condition; the yellow line indicates the invasion area on the spheroid body. (C, lower panel) Histograms represent the invading area's quantification and the protrusion's average length (N = 10 per condition, scale bar = 250 µm, data are ±s.e., Student's t-test, p < 0.001. D–F Human TNBC BT-549 cells were transfected with shControl and shST6GAL1 #1 (Sigma Cat# TRCN0000035432) or shST6GAL1 #2 (Sigma Cat# TRCN0000035429), as mentioned before. 3D cell invasion assays (F, upper panels), quantification of cell invasion (F, lower panel), and cell proliferation (E) assays were performed. Duplicate cultures were used for qPCR and Western blot analysis of ST6GAL1 (D). For qPCR analysis, gene levels were normalized by GAPDH, N = 3, data are ±s.e., Student's t-test, p < 0.05. Experiments were repeated at least three times, and representative blots and images were shown. For Western blotting, tubulin as a loading control for equal loading was used for housekeeping control for equal loading. Histograms data for invasion assays are means ± s.e., scale bar 250 µm, one-way ANOVA, and post-hoc test for pairwise comparison, p < 0.001.

CMP-Sia, the ST6GAL1 donor substrate, alone did not affect any effect on MDA-231 proliferation. However, self-exosomes from a different culture of MDA-231 and additional rST6G have resulted in significant MDA-231 cell growth. In addition, adding exosomes that were recovered from the conditioned media of BT-549, natively ST6GAL1-high cells, resulted in significantly greater MDA-231 cell growth, which could be further enhanced in the presence of both BT-549 exosomes and rST6G. Maximal growth was observed when the added BT-549 exosomes were doubled. At the increased level of BT-549 exosomes, the additional presence of rST6G did not result in greater MDA-231 growth (Fig. 5A). These observations supported the idea that extracellular or extrinsic ST6GAL1 affects the biology of the TNBC cancer cell MDA-231. MDA-231 cell surface SNA reactivity was also increased in the presence of BT-549 exosomes; SNA signal was increased further when both BT-549 exosomes and rST6G were present (Fig. 5B), supporting the idea that extracellular ST6GAL1, in the forms of BT-549 exosome encapsulated ST6GAL1 as well as free recombinant enzyme, can act on target MDA-231 in the construction of α2,6-sialic acid epitopes.

Mouse E0771 cells with a medium level of native ST6GAL1 expression (Fig. 2) was examined next. E0771 cell proliferation was elevated with added rST6G to the culture medium, as well as self-exosomes (e.g., exosomes isolated from a separate culture of E0771 cells). Maximum E0771 proliferation was observed when both E0771 exosomes and rST6G were present in the culture medium (Fig. 5C). The synergistic effects of rST6G and exosomes (see Fig. 5B SNA reactivity and Fig. 5C) strongly suggest the...
in vitro. 4T1.2 cells subjected to shRNA knockdown of endogenous ST6GAL1 expression strikingly muted invadopodia formation earlier (Fig. 3), we showed that shRNA knockdown of native ST6GAL1 for invasiveness ex vivo. Extracellular ST6GAL1 compensates cell-intrinsic native cancer cell invasion, treatments, including rST6G only. Representative 10× magnification light microscopy images are shown on a scale of 50 µm. Mouse breast cancer E0771 cells were cultured in the serum-free medium and treated with a control medium, 1 µg exosomes (BT-549), or exosome particles (BT-549) mixed with 1 µg rST6G for 20 min in a serum-free medium. SNA-lectins were stained (Green), and nuclei were stained with DAPI (blue color). Representative overlay images are shown. Invadopodia formation of human BT-549 cells was enhanced dramatically in the presence of both exosomes and rST6G (Fig. 5E). Since BT-549 already has significantly more ST6GAL1 in invadopodia formation than the shST6GAL1#1 spheroids, promoting invadopodia. In contrast, shST6GAL#1 exosomes are unable to restore invadopodia in the shST6GAL1#1 spheroids, implicating the ST6GAL1 cargo within the exosomes as the driver for invadopodia formation. A prominent role for extracellular ST6GAL1 in invadopodia formation was confirmed when either shControl or shST6GAL1#1 cultures were exposed to their respective self-exosomes with the further addition of rST6G. We observed the same dependence for extracellular ST6GAL1 for invadopodia formation in BT-549 cells (Fig. 6B). For BT-549, we show the results using two different sets of ST6GAL1 knockdown RNAs. As in 4T1.2, BT-549 exosomes from wild-type (shControl), but not from knockdown cultures, promoted invadopodia formation. The additional presence of recombinant ST6GAL1 further drives invadopodia formation. Supplementary Figure S5 panel A and panel B shows that shRNA knockdown of 4T1.2 cells and BT-549 resulted in an almost complete reduction of ST6GAL1 mRNA and protein levels. Additional breast cancer cells, 4T1 and E0771, were also tested, with similar results (Supplemental Fig. S4). Together these observations point to a role of extracellular ST6GAL1, either as encapsulated cargo within the exosomes or in freely soluble enzyme, in promoting invasiveness phenotype. Moreover, and most importantly, extracellular ST6GAL1 can be present of exosome cofactor(s) contributing to the extracellular ST6GAL1 mediated α2,6-sialylation of the target cell surface.

The mouse 4T1.2, the human MDA-231, and BT-549 cells that form invadopodia in an in vitro 3D culture setting were used to assess how extracellular ST6GAL1 might affect aggressive cancer cell’s invasiveness. In 4T1.2 cells, invadopodia formation was doubled when rST6G was included in the culture medium; invadopodia formation was enhanced further when CMP-Sia, the donor substrate to ST6GAL1, was also included (Fig. 5F). This observation supports the idea that extracellular ST6GAL1-mediated catalysis can drive invasive tumor behavior in vitro. Invadopodia formation of human BT-549 cells was enhanced marginally by the addition of self-exosomes (isolated from a separate culture of BT-549 cells), but invadopodia formation was dramatically enhanced in the presence of both exosomes and rST6G (Fig. 5E). Since BT-549 already has significant native ST6GAL1 expression, this observation further confirms the ability of extracellular ST6GAL1 in driving BT-549 biology and the putative existence of cofactor(s) in exosomes to augment extracellular ST6GAL1 action. MDA-231 cells with low endogenous ST6GAL1 further confirm that self-exosomes (isolated from a separate culture medium of MDA-231 cells) and additional rST6G but not self-exosomes alone significantly augment MDA-231 cell 3D invasiveness (Fig. 5D).

**Extracellular ST6GAL1 compensates cell-intrinsic native ST6GAL1 for invasiveness ex vivo**

Earlier (Fig. 3), we showed that shRNA knockdown of native ST6GAL1 expression strikingly muted invadopodia formation in vitro. 4T1.2 cells subjected to shRNA knockdown of endogenous ST6GAL1 (shST6GAL1#1), compared to control knockdown using irrelevant RNA sequences (shControl), had strikingly diminished invadopodia formation.

Here in Fig. 6A, we present evidence that invadopodia formation can be further enhanced in shControl, the cells with wild-type native ST6GAL1 expression when these cells are incubated with exosomes generated by a parallel shControl culture. This observation indicates a role for exosome contents in promoting invadopodia. In contrast, shST6GAL1#1 exosomes are unable to restore invadopodia in the shST6GAL1#1 spheroids, implicating the ST6GAL1 cargo within the exosomes as the driver for invadopodia formation. A prominent role for extracellular ST6GAL1 in invadopodia formation was confirmed when either shControl or shST6GAL1#1 cultures were exposed to their respective self-exosomes with the further addition of rST6G. We observed the same dependence for extracellular ST6GAL1 for invadopodia formation in BT-549 cells (Fig. 6B). For BT-549, we show the results using two different sets of ST6GAL1 knockdown RNAs. As in 4T1.2, BT-549 exosomes from wild-type (shControl), but not from knockdown cultures, promoted invadopodia formation. The additional presence of recombinant ST6GAL1 further drives invadopodia formation. Supplementary Figure S5 panel A and panel B shows that shRNA knockdown of 4T1.2 cells and BT-549 resulted in an almost complete reduction of ST6GAL1 mRNA and protein levels. Additional breast cancer cells, 4T1 and E0771, were also tested, with similar results (Supplemental Fig. S4). Together these observations point to a role of extracellular ST6GAL1, either as encapsulated cargo within the exosomes or in freely soluble enzyme, in promoting invasiveness phenotype. Moreover, and most importantly, extracellular ST6GAL1 can be...
released from distal sources; non-cell-autonomous ST6GAL1 can compensate for the lack of native ST6GAL1 expression.

**Extracellular ST6GAL1 promotes enhanced expression of mesenchymal and stemness markers**

Here, we examine the molecular contribution of ST6GAL1, particularly the ST6GAL1 in the extracellular spaces, on breast tumor cells. shST6GAL1 knockdown BT-549 and 4T1.2 cells were compared to their respective shControl, showing increased caspase 3 and PARP activation upon serum depletion (Fig. 7A), which is consistent with the idea that apoptosis is promoted in ST6GAL1 insufficiency. Moreover, self-exosomes (shST6GAL1 exosomes with diminished ST6GAL1 cargo) and additional rST6G compensated for diminished endogenous ST6GAL1 expression by dramatically suppressing cleaved Caspase 3 and PARP expression in both BT-549 and 4T1.2, supporting the idea that extracellular ST6GAL1 can compensate cell-intrinsic ST6GAL1 and render protection from cell death (Fig. 7A). Extracellular ST6GAL1 also supplements cell-intrinsic ST6GAL1, driving mesenchymal and stemness marker expression. 4T1.2 cells with knockdown ST6GAL1 have elevated E-cadherin (an epithelial marker) and diminished N-cadherin (a mesenchymal marker), and decreased Snail (a mesenchymal associated transcription factor) and other mesenchymal stemness-associated markers (Fig. 7B and Supplemental Fig. S6). In T47D cells with a knockdown ST6GAL1, we observed increased E-cadherin (an epithelial marker) on the mRNA and protein level (Fig. 7C, D). The addition of self-exosomes together with rST6G dramatically suppressed E-cadherin expression, elevated N-cadherin, concomitant with elevated expression of EMT transcriptional factors Slug and Twist, and stemness-associated cMYC, Oct4, and Nanog. CD44 and Aldh1, other CSC genes related to stemness, were also elevated (Fig. 7C, D and Supplemental Fig. S6). Together, our observations support a role of ST6GAL1 in promoting the maintenance of the mesenchymal state, and ST6GAL1 insufficiency promotes a switch from mesenchymal to epithelial properties and decreases cell mortality. Most importantly, our data show extracellular ST6GAL1, not necessarily of cell-autonomous origins, can complement cell-intrinsic enzymes in maintaining mesenchymal phenotypes and survival.

**DISCUSSION**

The sialyltransferase ST6GAL1 modifies glycans and glycoproteins important not only in cancer progression [3, 8, 50] but also in the maintenance of cancer stem cell phenotype [51]. Generally, elevated ST6GAL1 and dysregulated α2,6-sialylation are prominently correlated with increased malignancy, especially in the lung, colorectal, pancreatic, and breast cancers [52, 53]. Canonicall, ST6GAL1 resides in the intracellular ER-Golgi secretory complex and cell-autonomously sialylated nascent cell surface and secreted components. However, far less attention was paid to the extensive reservoir of catalytically active ST6GAL1 in the extracellular milieu, but published reports have implicated the extracellular ST6GAL1 as an important systemic regulator of blood cell development and function [16–19, 35].

A number of important conclusions can be drawn from our observations. First, the data suggest the importance of cancer cell ST6GAL1 in several important cancer cell phenotypes, including cell proliferation and invadopodia formation. Overall this is fully consistent with the known importance of ST6GAL1-mediated α2,6-sialic acid-linked glycans/glycoproteins in tumor progression and metastasis [1–3, 6, 9, 54].

Second, an extensive panel of mouse and human breast cancer cell lines had highly heterogenous ST6GAL1 expression. While we observed this heterogeneity is consistent with clinical breast
cancer, how this heterogeneity contributes to overall malignancy and cancer progression is not yet clear. Breast cancer is a heterogeneous disease [55]. TNBC as the most aggressive and metastatic subtype of breast cancer and has limited therapeutic options. Aggressive metastatic diseases are killing more than 90% of breast cancer patients [56]. Therefore, understanding the role of sialylation in invasive, aggressive breast cancer is clinically relevant. Although ST6GAL1 transcripts levels are lower in breast tumors than in normal adjacent tissues in TCGA breast cancer cohort, however among cancer patients, ST6GAL1 transcripts...
CHC was used for equal loading and transfer. Experiments were repeated at least three times, and representative blots were shown.

Levels are significantly elevated in triple-negative clinical cases compared with ER+ patients (Fig. 1A, B, and Supplementary Fig. S1A). By analyzing TCGA breast cancer data, we found that ST6GAL1 transcripts were not associated with aggressive stages of breast cancer in patients; however, they were linked to a higher grade of cancer patients (Fig. 1C and Supplementary Fig. S1B, C). In agreement with the cancer-promoting role of ST6GAL1, recent data revealed that ST6GAL1 is upregulated in metastatic pancreatic cancer cell models and displayed enrichment of gene networks associated with stemness, EMT, and hypoxia pathways [9]. Gene enrichment data analysis using both the METABRIC and TCGA breast cancer cohort suggested that higher levels of tumor ST6GAL1 transcripts were linked with metastatic progression of cancer pathways gene sets, including Hedgehog, EMT, and hypoxia (Fig. 1F–H and Supplementary Fig. S1D–F). In sync with the notion that ST6GAL1 is critical for breast cancer, we found that knockdown of ST6GAL1 with shRNA targeted to the different sites of ST6GAL1 significantly reduced aggressive breast cancer cell proliferation and invasive properties in a 3D cell culture setting (Fig. 3). We also observed that tumor ST6GAL1 transcripts are significantly associated with higher intra-tumor heterogeneity scores in breast cancer patients in TCGA breast cancer cohort (Fig. 1D). Poorly understood intra-tumor heterogeneity, with reference to the tumor-specific subclonal driver gene mutations and copy number alteration, is thought to be involved in aggressive cancers [57, 58]. ST6GAL1 might just provide another layer of heterogeneity in aggressive breast cancer.

Generally, there is a correlation between ST6GAL1 mRNA and protein in tumor cells; however, there is an exception [18]. Transcriptional regulation of the st6gal1 gene is complex in tumor cells, and we have noticed that human breast tumor cells have a low level of ST6GAL1 mRNA compared with mouse breast cancer cells. In some experiments, we have also noted that the level of intracellular ST6GAL1 protein does not always correspond with mRNA levels in breast cancer cells. Since most of the ST6GAL1 is continually secreted, it is likely that decreased mRNA will principally affect released ST6GAL1 rather than the retained enzyme. Indeed, that is what we observed in some shRNA-mediated ST6GAL1-knockdown experiments in breast cancer cells. However, we feel a study of the kinetic relationship between released and retained ST6GAL1 is beyond the scope of this manuscript and was not pursued here.

Third, breast cancer cells release bioactive exosome- and exomere-like particles, but the particles from different breast cancer lines have vastly different amounts of ST6GAL1 cargo. This is in agreement with earlier findings that released cargo of exosome particles are enriched with α2,6-sialylated glycoproteins [59] and ST6GAL1 [23]. Earlier studies in colon cancer cell models suggested that variable functional ST6GAL1 in exosomes released by tumor cells are linked to the source cells [23]. The study also suggested that tumor cells with a higher level of endogenous ST6GAL1 protein released exosomes can transfer ST6GAL1 into the lower natively expressing ST6GAL1 tumor cells and enhance ST6GAL1-mediated cellular signaling in the recipient cells. Our early work suggests that the liver mainly releases soluble ST6GAL1 that is found in circulation. Here, our data suggest that certain cancer cells have the capacity to significantly raise the local extracellular ST6GAL1 levels. In the form of a recombinant protein (rST6GAL1) in combination with the exosomes released by tumor cells, these are capable of impacting target tumor cells and compensate for endogenous ST6GAL1 functions. Fourth, the exosome ST6GAL1 cargo compensates for deficiency in cell-native ST6GAL1 expression in restoring proliferation and invadopodia formation. In the TNBC clinical specimens, we observed that ST6GAL1 is heterogeneously expressed (Fig. 1E), which could correlate with the heterogeneous expression of α2,6-sialic acid-linked lectins [11]. The biology driving this α2,6-sialylation heterogeneity is unknown, and how it contributes to the overall pathobiology of the cancer is also completely unknown. The heterogeneity may underscore the current confusion of why ST6GAL1 expression is observed to be beneficial but sometimes detrimental to patient outcome. Using KM plotter web-based data analysis [60], our results suggested that tumor ST6GAL1 mRNA was associated with favorable disease-free survival of TNBC patients (Fig. 1I), suggesting a complex mechanism of ST6GAL1-mediated cancer progression, warrant further investigation involving tumor microenvironment (TME). The ability of extracellular ST6GAL1 to compensate for the deficiency in cell-native ST6GAL1 expression may provide a new perspective in understanding the role of this sialyltransferase in cancer.

Fifth and most unexpectedly, although cancer exosomes devoid of ST6GAL1 cargo are unable to promote tumor cell survival, cell proliferation, and invadopodia formation, these properties are restored upon the addition of soluble recombinant ST6GAL1 (Fig. 6 and Supplementary Fig. S4). Notably, the addition of exogenous sources of ST6GAL1 can induce a more mesenchymal and CSC phenotype of breast cancer cells (Fig. 7 and Supplementary Fig. S6). This observation strongly implicates a cofactor in exosomes facilitating extracellular ST6GAL1 action. While the identity of this cofactor and its mechanism of action remains to be formally elucidated, an earlier report indicated that activated platelets release vesicle-like particles that supply the sialic acid donor substrate needed for the extracellular sialyltransferase catalysis [35].

Taken together, our data further support a role for ST6GAL1 in breast cancer progression. Breast cancer is a heterogeneous disease with multiple subtypes. We were using multiple TNBC and ER+ cell lines to establish that extracellular ST6GAL1-mediated novel signaling is important for tumor cell survival, proliferation, invasion, and EMT progression. Canonically, ST6GAL1 resides within the intracellular ER-Golgi secretory complex, where the sialyltransferase cell-autonomously glycosylates nascent glycoproteins in biosynthetic transit. However, our data demonstrate the existence of an extracellular mechanistic pathway for ST6GAL1 function in breast cancer. At this time, it is unclear whether the intrinsic and extrinsic mechanisms are redundant pathways or...
whether these mechanisms address exclusive needs unmet by the other. Moreover, mechanistically it is not clear whether the external ST6GAL1 needs to be internalized by the target cells or the external ST6GAL1 acts directly on the cell surface of the targets. A recent study has elegantly demonstrated that tumor cells secrete ST6GAL1 in exosome vesicles or exomere smaller particles and are capable of transferring into the recipient cell as a functional ST6GAL1 in colon cancer cell models [23, 61].

On the other hand, other reports highly suggest that extracellular ST6GAL1 directly sialylates the cell surfaces of target cells [16–18, 35, 62, 63]. Whichever the mechanistic conduit for extracellular ST6GAL1 action, at least some biologic functions, namely promoting cell proliferation and invasiveness, can be restored by extracellular ST6GAL1 in lieu of adequate cell-native ST6GAL1 expression. While our data suggest the role of ST6GAL1 in driving aggressive cancer cell behavior, our data also indicate that the intrinsic ability of individual cancer cells to cell-autonomously express ST6GAL1 might not be absolutely necessary. Lastly, there is also the intriguing possibility that cancer cell natively expressed ST6GAL1 needs to be first excreted into the extracellular space.

DATA AVAILABILITY
Publicly available datasets were analyzed in this study.

REFERENCES
1. Pinho SS, Reis CA. Glycosylation in cancer: mechanisms and clinical implications. Nat Rev Cancer. 2015;15:540–55.
2. Magalhaes A, Duarte HO, Reis CA. Aberrant glycosylation in cancer: a novel molecular mechanism controlling metastasis. Cancer Cell. 2017;1;735–3.
3. Dorsett KA, Marcelli MP, Kwang J, Ankenbauer KE, Bhalerao N, Belfis SL. Regulation of ST6GAL1 sialyltransferase expression in cancer cells. Glycobiology. 2021;31:530–9.
4. Swendall AF, Londono-Joshi AI, Schwartz MJ, Fineberg N, Buchsbaum DJ, Belfis SL. ST6GAL1 protein expression is upregulated in human epithelial tumors and correlates with stem cell markers in normal tissues and colon cancer cell lines. Cancer Res. 2013;73;2368–78.
5. Hsieh CC, Shyr YM, Liao WY, Chen TH, Wang SE, Lu PC, et al. Elevation of beta-galactoside alpha2,6-sialyltransferase 1 in a fructoseoverresponsive manner promotes pancreatic cancer metastasis. Oncotarget. 2017;8;7691–709.
6. Wichter B, Milde-Langosch K, Galatenko V, Schmalfeldt B, Oliveira-Ferrer L. Prognostic role of the sialyltransferase ST6GAL1 in ovarian cancer. Glycobiology. 2018;28;899–903.
7. Rao TC, Beggs RR, Ankenbauer KE, Kwang J, Ma VP, Salata K, et al. ST6GAL1-mediated sialylation of the epidermal growth factor receptor modulates cell mechanics and enhances invasion. J Biol Chem. 2022;298;101726.
8. Garnham P, Scott E, Livermore KE, Munkley J. ST6GAL1: a key player in cancer. Oncotell. 2019;18;1983–9.
9. Britain CM, Bhalerao N, Silva AD, Chakraborty A, Buchsbaum DJ, Crowley MR, et al. Glycosyltransferase ST6Gal-I promotes the epithelial to mesenchymal transition in pancreatic cancer cells. J Biol Chem. 2021;296;100034.
10. Smithson M, Irwin R, Williams G, Alexander KL, Smythies LE, Nearing M, et al. Siloxygenation in Brazilian glioma growth in vivo. Proc Natl Acad Sci USA. 2010;107;12646
11. Chin A, Qi Y, Poon E, Xie Y, Patel AK, Hain K, et al. siRNA-mediated silencing of ST6GAL1 efficiently in the sialyltransferase ST6GalI. Blood. 2006;108;3397
12. Shrestha S, Maiti A, Xu P, Qi Q, Kawaguchi T, Okano M, et al. Regulation of hypoxia-inducible factor functions in the nucleus by sialic acid-dependent RNAs for cancer diagnosis. ACS Appl Mater Interfaces. 2018;10;43375–86.
13. Sarkar S, Maceyka M, Halt NC, Paugh SW, Sankala H, Miliotis S, et al. Sphingosine kinase 1 is required for migration, proliferation and survival of MCF-7 human breast cancer cells. FEBS Lett. 2005;570;5313–7.
14. Luca R, Averna M, Zaffa F, Vecchi M, Bianchi F, La Fata G, et al. The fragile X protein binds mRNAs involved in cancer progression and modulates metastasis formation. EMBO Mol Med. 2013;5;1532–6.
15. Berens EB, Holy JM, Riegel AT, Weilstein A. A cancer cell spheroid assay to assess invasion in a 3D setting. J Vis Exp. 2015;105;34009–414.
16. Halt NC, Maiti A, Xu P, Qi Q, Kawaguchi T, Okano M, et al. Regulation of hypoxia-inducible factor functions in the nucleus by sialic acid–1–phosphate. FASEB J. 2020;34;4293–310.
17. Mali A, Qi Y, Poon E, Xie Y, Balaka H, Halt NC. Class I histone deacetylase inhibitor suppresses sarscoglogenic mimicry by enhancing the expression of tumor suppressor and anti-angiogenesis genes in aggressive human TNBC cells. Int J Cancer. 2019;155;1154–66.
18. Manhardt CT, Punch PR, Dougher CWL, Lau JTY. Extrinsic sialylation is dynamically regulated by systemic triggers in vivo. J Biol Chem. 2017;292;13514–5.
19. Schindelin J, Arganda-Carreras I, Frise E, Kaynig V, Piotzsch T, et al. Fiji: an open-source platform for biological-image analysis. Nat Methods. 2012;9;676–82.
20. Phansopa C, Kozak RP, Liew LP, Frey AM, Farmilo T, Parker J, et al. Characterization of a sialate-O-acetylesterase (NanS) from the oral pathogen Tannerella forsythia that enhances sialic acid release by NanS, its cognate sialidase. Biochem J. 2015;472;157–67.
21. Gao J, Aksoy BA, Dogrusoz U, Dresner G, Brown G, Sumer SO, et al. Integrative analysis of complex cancer genomics and clinical profiles using the cBioPortal. Sci Signal. 2013;6;pl1.
22. Curtis C, Shah SP, Chin SF, Taramizhali G, Rueda OM, Dunning MJ, et al. The genomic and transcriptomic architecture of 2,000 breast tumours reveals novel subgroups. Nature. 2012;486;346–52.
23. Rocco JW. Mutant allele tumor heterogeneity (MATH) and head and neck squamous cell carcinoma. Head Neck Pathol. 2015;9;1–5.
24. Subramanian A, Tamayo P, Mootha VK, Mukherjee S, Ebert BL, Gillette MA, et al. Gene set enrichment analysis: a knowledge-based approach for interpreting gene expression data. Proc Natl Acad Sci USA. 2005;102;15455–50.
25. Duarte HO, Rodrigues JG, Gomes C, Hensbergen PJ, Edvardsen ALH, de Ru AH, et al. ST6GAL1 targets the ectodomain of ErbB2 in a site-specific manner and regulates gastric cancer cell sensitivity to trastuzumab. Oncogene. 2021;40;3719–73.
26. Friedl P, Alexander S. Cancer invasion and the microenvironment: plasticity and reciprocity. Cell. 2011;147;992–1009.
44. Escrevente C, Grammel N, Kandzia S, Zeiser J, Tranfield EM, Conradt HS, et al. Sialoglycoproteins and N-glycans from secreted exosomes of ovarian carcinoma cells. PLoS ONE. 2013;8:e78631.

45. Barres C, Blanc L, Bette-Bobillo P, Andre S, Mamoun R, Gabius HJ, et al. Galectin-5 is bound onto the surface of rat reticulocyte exosomes and modulates vesicle uptake by macrophages. Blood. 2010;115:696–705.

46. Poloni L, Orizio F, Busatto S, Radeghieri A, Bresciani R, Bergese P, et al. Exosomes secreted by HeLa cells shuttle on their surface the plasma membrane-associated sialidase NEU3. Biochemistry. 2015;56:6401–8.

47. Escrevente C, Keller S, Altevogt P, Costa J. Interaction and uptake of exosomes by ovarian cancer cells. BMC Cancer. 2011;11:108.

48. Lotvall J, Hill AF, Hochberg F, Buzas EI, Di Vizio D, Gardiner C, et al. Minimal experimental requirements for definition of extracellular vesicles and their functions: a position statement from the International Society for Extracellular Vesicles. J Extracell Vesicles. 2014;3:26913.

49. Tkach M, Kowal J, Thery C. Why the need and how to approach the functional diversity of extracellular vesicles. Philos Trans R Soc Lond B Biol Sci. 2018;373:20160479–87.

50. Costa AF, Campos D, Reis CA, Gomes C. Targeting glycosylation: a new road for cancer drug discovery. Trends Cancer. 2020;6:757–66.

51. Schultz MJ, Holdbrooks AT, Chakraborty A, Grizzle WE, Landen CN, Buchsbaum DJ, et al. The tumor-associated glycosyltransferase ST6Gal-I regulates stem cell transcription factors and confers a cancer stem cell phenotype. Cancer Res. 2016;76:3978–88.

52. Dall’Olio F, Malagolini N, Trinchera M, Chiricolo M. Sialosignaling: sialyl-transferases as engines of self-fueling loops in cancer progression. Biochim Biophys Acta. 2014;1840:2752–64.

53. Rodrigues E, Macauley MS. Hypersialylation in cancer: modulation of inflammation and therapeutic opportunitie. Cancers. 2018;10:207–25.

54. Hsieh TH, Hsu CY, Tsai CF, Long CY, Wu CH, Wu DC, et al. HDAC inhibitors target HDACS, upregulate microRNA-125a-5p, and induce apoptosis in breast cancer cells. Mol Ther. 2015;23:656–66.

55. Jin X, Mu P. Targeting breast cancer metastasis. Breast Cancer. 2015;9:23–34.

56. Anders C, Carey LA. Understanding and treating triple-negative breast cancer. Oncology. 2008;22:1233–9; discussion 1239–43.

57. Thorsson V, Gibbs DL, Brown SD, Wolf D, Bortone DS, Ou Yang TH, et al. The immune landscape of cancer. Immunity. 2019;51:411–2.

58. Dento SC, Leshchiner I, Haase K, Tarabichi M, Wintersinger J, Deshwar AG, et al. Characterizing genetic intra-tumor heterogeneity across 2,658 human cancer genomes. Cell. 2021;184:2239–54 e439.

59. Zhang H, Freitas D, Kim HS, Fabijanic K, Li Z, Chen H, et al. Identification of distinct nanoparticles and subsets of extracellular vesicles by asymmetric flow field-flow fractionation. Nat Cell Biol. 2018;20:332–43.

60. Nagy A, Lancezy A, Menyhart O, Gyoryfi B. Validation of miRNA prognostic power in hepatocellular carcinoma using expression data of independent datasets. Sci Rep. 2018;8:9227.

61. Martins AM, Ramos CC, Freitas D, Reis CA. Glycosylation of cancer extracellular vesicles: capture strategies, functional roles and potential clinical applications. Cells. 2021;10:1099–33.

62. Punch PR, Irons EE, Manhardt CT, Marathe H, Lau JTY. The sialytransferase ST6GalNAC1 protects against radiation-induced gastrointestinal damage. Glycobiology. 2020;30:446–53.

63. Lee-Sundlov MM, Ashline DJ, Hanneman AJ, Grozovsky R, Reinhold VN, Hoffmeister KM, et al. Circulating blood and platelets supply glycosyltransferases that enable extrinsic extracellular glycosylation. Glycobiology. 2017;27:188–98.

64. Cancer Genome Atlas N. Comprehensive molecular portraits of human breast tumours. Nature. 2012;490:61–70.

65. Lee MM, Nasiri-Kenari M, Manhardt CT, Ashline DJ, Hanneman AJ, Reinhold VN, et al. Platelets support extracellular sialylation by supplying the sugar donor substrate. J Biol Chem. 2014;289:8742–8.

ACKNOWLEDGEMENTS

This work was supported by grants R01AI140736 and P01HL51333 to JL and the Roswell Park Health Research Incorporated (HRI) Start-Up Funds #714084-01 to NCH. We acknowledge funding support from the National Cancer Institute (NCI) of the National Institutes of Health (NIH) under award number R21CA235305 to YW. The content is solely the authors’ responsibility and does not necessarily represent the official views of NIH. We thank the support from National Science Foundation under award number CBET-1337860 which funds the nanoparticle tracking analysis system (NanoSight, LM10, Malvern Instruments Ltd). We would like to thank National Cancer Institute (NCI) R01 CA207504 (to JZ). We also acknowledge the support of NIH grants R34 GM134794 and R01 GM130915 to KWM to produce the ST6GAL1 enzyme. The core facilities of Roswell Park Comprehensive Cancer Center used in this work were supported in part by NIH National Cancer Institute Cancer Center Support Grant CA07606. We thank Mr. Donald MacFarland, Lead Electron Microscopist, University at Buffalo, Jacobs School of Medicine, Buffalo, NY, USA, for the TEM analysis.

AUTHOR CONTRIBUTIONS

Study concept and design: NCH and JTYL. Acquisition of data: NCH, AM, RW, VL-A, and CCH. Analysis and interpretation of data: NCH, AM, JTYL, YW, KT, MER, WB, and JZ. Bioinformatics: RW. Material support: DC and KWM. Study supervision: NCH and JTYL. Manuscript writing: NCH and JTYL. Manuscript editing: all the authors.

COMPETING INTERESTS

The authors declare no competing interests.

ADDITIONAL INFORMATION

Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s41417-022-00485-y.

Correspondence and requests for materials should be addressed to Nitiati C. Hait or Joseph T. Y. Lau.

Reprints and permission information is available at http://www.nature.com/reprints

Publisher’s note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2022