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Differentiation of Cancer Cell Origin and Molecular Subtype by Plasma Membrane N-Glycan Profiling

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Abstract

In clinical settings, biopsies are routinely used to determine cancer type and grade based on tumor cell morphology, as determined via histochemical or immunohistochemical staining.

Unfortunately, in a significant number of cases, traditional biopsy results are either inconclusive or do not provide full subtype differentiation, possibly leading to inefficient or ineffective treatment. Glycomic profiling of the cell membrane offers an alternate route towards cancer diagnosis. In this study, isomer-sensitive nano-LC/MS was used to directly obtain detailed profiles of the different N-glycan structures present on cancer cell membranes. Membrane N-glycans were extracted from cells representing various subtypes of breast, lung, cervical, ovarian, and lymphatic cancer. Chipbased porous graphitized carbon nano-LC/MS was used to separate, identify, and quantify the native N-glycans. Structure-sensitive N-glycan profiling identified hundreds of glycan peaks per cell line, including multiple isomers for most compositions. Hierarchical clusterings based on Pearson correlation coefficients were used to quickly compare and separate each cell line according to originating organ and disease subtype. Based simply on the relative abundances of broad glycan classes (e.g. high mannose, complex/hybrid fucosylated, complex/hybrid sialylated, etc.) most cell lines were readily differentiated. More closely-related cell lines were differentiated based on several-fold differences in the abundances of individual glycans. Based on characteristic N-glycan profiles, primary cancer origins and molecular subtypes could be distinguished. These results demonstrate that stark differences in cancer cell membrane glycosylation can be exploited to create an MS-based biopsy, with potential applications towards cancer diagnosis and direction of treatment.

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Supporting Information Available: This material is available free of charge via the Internet at http://pubs.acs.org.

Keywords

mass spectrometry; LC/MS; N-glycans; cell membrane; cancer; molecular subtype

INTRODUCTION

The mammalian cell membrane is coated with a dense thicket of glycans — complex sugar chains that are post-translationally attached to most membrane proteins. These glycans serve as a primary point of contact between the cell and the rest of the world, mediating cell interactions with small molecules, microorganisms, and even other cells.¹ As a result, changes in glycosylation are often indicative of irregularities in the cellular machinery, and have previously been correlated with a number of human diseases — most famously, cancer. Several decades of research have now firmly established aberrant glycosylation as one of the hallmarks of cancer.^{2, 3} Glycans have been shown to mediate many cancer-related processes,^{4–8} and cell membrane glycans, in particular, play primary roles in tumorigenesis,^{9–11} metastatic colonization,^{12, 13} and chemotherapeutic drug resistance.^{14, 15}

Numerous studies on the human serum glycome (i.e. from secreted glycoproteins) have already identified potential biomarkers for multiple cancer types.^{16–19} However, the cell membrane glycome remains a largely untapped source of potential biomarkers, due in large part to the difficulty of isolating cell membrane glycoproteins from those found in the cytoplasm, nucleoplasm, or cellular organelles. Previous studies on cell membrane glycosylation have been limited in scope, relying largely on affinity interactions (either lectin- or antibody-based) to detect specific glycosylation motifs or epitopes.^{12, 20–22} Unfortunately, because these affinity interactions typically target only terminal monosaccharides, the analytical data gleaned from affinity-based studies is often inconclusive or subject to extensive interpretation. Additionally, due to the extraordinary structural heterogeneity of glycans, many glycan structures simply cannot be detected (or distinguished) by currently-available affinity-based methods.

Recently-developed LC/MS-based methods of glycomic profiling, in contrast, offer direct and comprehensive access to cell membrane glycosylation. An et al recently developed and validated a method to isolate and enrich cell membrane N-glycans, enabling quantitative, reproducible, and (most importantly) LC/MS-compatible analysis.²³ LC/MS compatibility is crucial to glycomic analyses, particularly when complex, structurally heterogenous analytes (such as cell membrane glycans) are involved. Over 200 glycosyltransferases and glycosidases work in concert to decorate proteins with glycosylation, and even slight perturbations in this delicate balance can result in widespread changes.²⁴ Glycomic profiling of these changes can be an extremely sensitive indicator of disease progression; however, detailed analytical methods are necessary in order to accurately capture the full structural complexity of the glycome. LC/MS fits this requirement exceedingly well, providing two-dimensional separation of complex glycan mixtures according to accurate mass, size, and three-dimensional conformation.

Nano-LC/MS methods for separation and profiling of N-glycans have recently been developed and optimized by Hua et al.¹⁶ These methods utilize chip-based porous graphitized carbon (PGC) nano-LC to separate complex glycan mixtures in a structure-sensitive manner prior to high resolution, accurate mass TOF-MS analysis. In tests on serum N-glycans, the optimized PGC nano-LC/TOF-MS platform provided detailed and comprehensive glycan identification, miniscule sample consumption, and dramatically increased sensitivity over previous MALDI-MS or high-flow LC/MS methods.²⁵

The present study combines cell membrane N-glycan enrichment procedures with detailed nano-LC/MS glycan profiling to provide the first comprehensive survey of cancer cell N-glycosylation. Cell membrane N-glycan profiles were obtained for fifteen different cell lines isolated from ovarian, breast, lung, cervical, and lymphatic cancer patients (as well as one non-cancerous fibrocystic breast cell line). Cell lines were grouped and compared according to biologically- or clinically-relevant characteristics (e.g. cancerous vs non-cancerous cell lines, lung carcinoma vs cervical carcinoma cell lines, etc). Significant changes in membrane glycan profiles were confirmed by statistical tests such as Pearson correlations and hierarchical clustering. Major differences were uncovered amongst the membrane glycan profiles of the analyzed cell lines, in terms of both overall glycan biosynthetic class and individual glycan signals. LC/MS-based membrane glycan profiling was shown to be a powerful platform for differentiation and identification of different cancer cell types.

METHODS

Cell culture

Cell lines (Hs-578T, ZR-75-1, MCF-7, MCF-7/HER2, MCF10A, Raji, Ramos, NCI-H929, BCBL-1, ES-2, NCI-H358, A549, C-33A, CaSki, and HeLa) were obtained from ATCC. Cell cultures were performed using the ATCC-recommended growth media, subcultivation ratios, and medium renewal interval (available online at www.atcc.org).

During cell harvest, in order to preserve membrane glycoprotein integrity, cells were scraped rather than enzymatically lifted. For each cell sample, approximately 10 million cells were counted and collected. Duplicate samples were collected from the lymphoma (Raji; Ramos; NCI-H929; BCBL-1), cervical carcinoma (33A; CaSki; HeLa), and lung carcinoma (NCI-H358; A549) cell cultures; however, due to cell growth limitations, only single samples were collected from the breast (Hs-578T; ZR-75-1; MCF-7; MCF-7/HER2; MCF10A) and ovarian (ES-2) cell cultures.

Plasma membrane extraction

Isolation of the plasma membrane was performed according to optimized procedures published by An et al.²³ Briefly, cells were suspended a homogenization buffer consisting of 0.25 M sucrose; 20 mM HEPES-KOH, pH 7.4; and 1:100 protease inhibitor mixture (EMD Millipore) prior to sonication. Cell lysates were ultracentrifuged at 1,000 g and 200,000 g to remove the nuclear and cytosolic fractions, respectively. Supernatants, consisting of enriched plasma membranes, were collected for glycan extraction.

N-glycan release and enrichment

Enzymatic release and solid-phase extraction of N-glycans were performed according to optimized procedures published by Kronewitter et al.²⁶ Briefly, membrane glycoproteins were denatured by rapid thermal cycling (25–100 °C) in an aqueous solution of 100 mM ammonium bicarbonate and 5 mM dithiothreitol. Next, 2.0 μ L (or 1000 U) of peptide N-glycosidase F (New England Biolabs) were added and the mixture was incubated in a microwave reactor (CEM Corporation) for 10 minutes at 20 watts. Following the addition of 800 μ L of cold ethanol, the mixture was chilled at –80 °C for 1 hour, then centrifuged in order to precipitate out the deglycosylated proteins. The glycan-rich supernatant fraction was collected and dried *in vacuo*.

Graphitized carbon solid-phase extraction was performed using an automated liquid handler (Gilson). Graphitized carbon cartridges (150 mg, 4.0 mL, Grace Davison) were washed with 80% acetonitrile / 0.10% trifluoroacetic acid (v/v) in water, then conditioned with pure water. Aqueous N-glycan solutions were loaded onto the cartridges and washed with pure

water to remove salts and buffer. Membrane N-glycans were eluted with sequential addition of 10% acetonitrile, 20% acetonitrile, and 40% acetonitrile / 0.05% trifluoroacetic acid (v/v) in water. Samples were dried *in vacuo*.

Nano-LC/MS analysis and data processing

MS analysis was performed according to optimized procedures published by Hua et al.¹⁶ Briefly, aqueous glycan solutions were injected by autosampler onto a chip-mounted nano-LC column (Agilent Technologies), consisting of a 9×0.075 mm i.d. enrichment column and a 43×0.075 mm i.d. analytical column, both packed with 5 µm porous graphitized carbon as the stationary phase. A rapid glycan elution gradient was delivered to the analytical column at 0.4 µL/min using solutions of (A) 3.0% acetonitrile and 0.1% formic acid (v/v) in water, and (B) 90.0% acetonitrile and 0.1% formic acid (v/v) in water, ramping from 5% to 35.9% B over the course of 16.5 min. Remaining non-glycan compounds were flushed out with 100% B prior to re-equilibration.

MS spectra were acquired by a TOF MS detector (Model 6210, Agilent Technologies) in positive ionization mode over a mass range of m/z 600–2000 with an acquisition time of 1.5 seconds per spectrum. Mass correction was enabled using reference masses of m/z 622.029, 922.010, 1221.991, and 1521.971 (ESI-TOF Calibrant Mix G1969-85000, Agilent Technologies).

Raw LC/MS data was filtered with a signal-to-noise ratio of 5.0 and parsed into a series of extracted ion chromatograms using the Molecular Feature Extractor algorithm included in the MassHunter Qualitative Analysis software (Version B.04.00, Agilent Technologies). Using expected isotopic distribution and charge state information, extracted ion chromatograms were combined to create extracted compound chromatograms (ECCs) representing the summed signal from all ion species associated with a single compound (e.g. the doubly protonated ion, the triply protonated ion, and all associated isotopologues). Thus, each individual ECC peak could be taken to represent the total ion count associated with a single distinct compound.

Each ECC peak was matched by accurate mass to a comprehensive library of all possible complex, hybrid, and high mannose glycan compositions based on known biosynthetic pathways and glycosylation patterns.^{27, 28} Deconvoluted masses of each ECC peak were compared against theoretical glycan masses using a mass error tolerance of 20 ppm and a false discovery rate of 0.6%. As all samples originated from human cell lines, only glycan compositions containing hexose (Hex), N-acetylhexosamine (HexNAc), fucose (Fuc), and N-acetylneuraminic acid (NeuAc) were considered.

RESULTS AND DISCUSSION

Separation and quantitative profiling of cell membrane N-glycans

LC/MS-based glycan profiling provides a comprehensive look at the different glycan compositions and structures present on the cell membrane. On average, cell membrane glycan profiles yielded over 250 N-linked glycan compound peaks with over 100 distinct N-linked glycan compositions, spanning five orders of magnitude. Each of the identified compositions include two or more peaks corresponding to either structural and/or linkage isomers (regioisomers) or, in some cases, anomeric isomers. For example, Figure 1a shows chromatograms of cell membrane N-glycans identified on non-CD4 T-cells from human blood. From this data, the relative abundances of individual glycan compositions or structures were readily quantified, simply by integrating the ion counts associated with each peak and normalizing to the total (summed) ion count of all glycans detected in the sample. Where applicable, the results of duplicate analyses were averaged. Previous studies have

already validated the quantitative reproducibility of the cell membrane glycan extraction procedure as well as the PGC nano-LC/MS platform, enabling rapid profiling and comparison of individual glycan abundances.^{16, 23}

In addition to *individual glycan profiling*, overarching differences in cell membrane glycosylation can be detected through *glycan class profiling*. In Figure 1a, the glycan compositional assignments have been used to sort glycan signals into different biosynthetic glycan classes, each represented by a different color — green for high mannose (high Man) glycans; blue for undecorated complex/hybrid (C/H) glycans; red for fucosylated complex/hybrid (C/H-F) glycans; purple for sialylated complex/hybrid (C/H-S) glycans; and orange for fucosylated-sialylated complex/hybrid (C/H-FS) glycans. Based on the summed ion counts of the individual glycan signals, relative abundances were assigned to each glycan class, creating a *glycan class profile* for the membrane N-glycans of each cell line (Figure 1b).

The ability of LC/MS-based glycomic analysis to detect unique cell membrane signatures at both the *individual* glycan level as well as the glycan *class* level has a number of potential clinical applications. To demonstrate this point, we hereby present a few clinically-relevant examples in which different cancer cell types are compared and differentiated.

Differentiation of cancerous and non-cancerous cells

When radiological imaging techniques detect the presence of tumors or lesions within a patient, doctors must determine whether these abnormal masses are malignant (cancerous), or benign (non-cancerous). While histological characteristics can often indicate whether a tumor is currently benign or cancerous, molecular markers can potentially increase accuracy, and might even indicate the prognosis and future inclinations of a tumor (e.g. likelihood of metastasizing).^{12, 15, 29, 30}

To demonstrate the utility of cell membrane glycan profiling towards tumor diagnosis, glycan class profiles (Figure 2) were compared amongst four cancerous breast cell lines (Hs-578T, ZR-75-1, MCF-7, and MCF-7/HER2) and one non-cancerous breast cell line (MCF-10A). All four cancerous cell lines originated from ductal breast carcinomas, whereas the non-cancerous MCF-10A cell line originated from fibrocystic breast tissue. Cursory inspection of the glycan class profiles reveals that all four breast carcinoma cell lines have much higher relative abundances (48–63%) of membrane high mannose glycosylation than the single non-cancerous fibrocystic breast cell line (23%). These results are consistent with previous research in which high mannose glycosylation is reported to increase during breast cancer.¹⁷

In addition to differentiating between cancerous and non-cancerous breast tissue, the glycan class profile (Figure 2) may also be used to differentiate amongst the four different breast carcinoma lines. For instance, cell line Hs-578T has much higher relative abundances of sialylated and fucosylated-sialylated complex/hybrid N-glycans compared to the other three breast carcinoma lines. Some of this increase may be associated with the presence of sialyl Lewis X (sLe^X) or sialyl Lewis A (sLe^A) epitopes, both of which are fucosylated-sialylated. Studies have shown that sLe^X and/or sLe^A are often found on the N-glycans of metastatic tumor cells, including carcinomas.^{12, 13} In clinical settings, direct detection of sLe^X and/or sLe^A on breast cancer carcinomas could help predict the metastatic potential of a tumor.

While informative at a macro level, the glycan class profiles shown in Figure 2 are ultimately just simplified representations of each cell's membrane glycosylation, summed from hundreds of individual glycan signals. To more precisely quantify the similarities and differences between the different cell lines, these individual glycan signals were

incorporated into pairwise Pearson correlations.³¹ Figure S1 shows a typical comparison between two breast cell lines: Hs-578T (cancerous) and MCF-10A (non-cancerous). Each point represents a different N-glycan, with the *x*-coordinate corresponding to the relative abundance of that glycan in Hs-578T cell membranes, and the *y*-coordinate corresponding to the relative abundance of that glycan in MCF-10A cell membranes. Based on these points, a best-fit line is drawn, and the Pearson correlation coefficient *R* is calculated. The low *R*-value of 0.217 indicates that the membrane glycosylation profiles of these two cell lines are quite different.

Pearson correlations were calculated for each possible pair of breast cell lines (Figure 2, **inset**), for a total of ten comparisons. In general, comparisons *amongst* cancerous breast cell lines show high correlation (0.700 < R < 0.932). Meanwhile, comparisons *between* cancerous and non-cancerous breast cell lines show low correlation (0.217 < R < 0.545). Based on these *R*-values, breast cell lines have been grouped into hierarchical clusters (Figure 2, **inset**). For example, closely-related cancerous breast cell lines MCF7 and MCF7/ HER2 (R = 0.932) are grouped together into a single cluster. Meanwhile, non-cancerous breast cell lines, providing rapid differentiation between non-cancerous and cancerous breast cells.

Identification of cancer cell origin and type

Once doctors identify a cancerous tumor in a patient, one of the first steps towards devising an effective treatment is to determine the primary origin of the cancer. However, in approximately 3% of patients, the primary cancer site is never found. Often, these cancers are poorly differentiated, lacking key morphological features that can be used to identify the origin of more developed cells.^{32–34} To make matters worse, poorly differentiated (high grade) cancers are typically the most aggressive, growing faster and spreading more rapidly than well-differentiated (low grade) cancers.³⁵ Lymphomas, which originate from the lymphatic system, commonly exhibit poor differentiation.^{32, 36}

In Figure 3, four lymphoproliferative cancer cell lines (Raji, Ramos, NCI-H929, and BCBL-1) are compared with a single ovarian carcinoma cell line (ES-2). The Raji and Ramos cell lines originate from Burkitt's lymphomas; the NCI-H929 from a multiple myeloma; the BCBL-1 from a primary effusion lymphoma; and the ES-2 from a poorly-differentiated ovarian clear cell carcinoma. As before, cursory inspection of the glycan class profiles (Figure 3) reveals stark differences between the different cell lines. For example, using just the relative abundances of the high mannose glycosylation, it is possible to separate the cell lines into three general groupings — the Raji and Ramos cell lines, at 33–34% high mannose glycosylation; the NCI-H929 and BCBL-1 cell lines, at 59–61% high mannose glycosylation; and the ES-2 cell line, at 7% high mannose glycosylation. These groupings are reflected in the Pearson correlations and hierarchical clustering (Figure 3, **inset**), in which the Raji and Ramos cell lines occupy one cluster, with very high correlation to each other; the NCI-H929 and BCBL-1 cell lines ES-2 cell line is grouped apart from all of the lymphoma cell lines, with very low correlation to any of them.

The incredibly high correlation of the Raji and Ramos cell lines (Figure 4) is quite significant biologically, because both of these cell lines derive from Burkitt's lymphoma patients. However, the patients were different from each other in almost every other respect — Raji was isolated from an African in Nigeria, whereas Ramos was isolated from a Caucasian in the United States; Raji represents the endemic form of Burkitt's lymphoma, whereas Ramos represents the spontaneous form; Raji was isolated in the early 1960s, whereas Ramos was isolated more than a decade later.^{37, 38} Yet, despite these differences, the individual glycan signals isolated from the Raji and Ramos cell membranes exhibit a

startlingly high correlation (R = 0.974) — suggesting that, in at least some cases, glycosylation patterns can be linked to specific cancer variants even in spite of confounding genetic and environmental factors.

Distinguishing between epithelial carcinomas from different organs

Identification of a primary cancer site enables doctors to focus diagnostic testing and treatment, significantly improving a patient's chances for survival; however, radiological observations are often obfuscated by metastatic tumors. To make matters worse, carcinomas, which derive from a variety of epithelial tissues, often exhibit very similar histologic and biochemical characteristics, even when they originate from different organs. This similarity creates significant problems when doctors need to differentiate between primary and metastatic carcinomas.

Cell membrane glycan profiling, however, may help determine the origin of a carcinoma. To demonstrate this concept, several cervical and lung carcinoma cell lines were selected for analysis. Cervical carcinomas commonly metastasize to the lung, forcing doctors to differentiate between primary lung carcinomas and metastatic cervical carcinomas prior to treatment.³⁹ Therefore, in Figure 5, N-glycan profiles were compared amongst two lung carcinoma cell lines (NCI-H358 and A549) and three cervical carcinoma cell lines (C-33A, CaSki, and HeLa). Glycan class profiling as well as hierarchical clustering (Figure 5, inset) reveals two major clusters: one containing both lung carcinoma cell lines, and the other containing two of the three cervical carcinoma cell lines (CaSki and HeLa). The placement of the third cervical carcinoma cell line, C-33A, is more intriguing — clustering actually reveals that cervical carcinoma cell line C-33A is more closely related to lung carcinoma cell lines NCI-H358 and A549 than to its fellow cervical carcinoma cell lines CaSki and HeLa. C-33A is rather unique amongst cervical carcinoma cell lines in that it is completely negative for human papillomavirus (HPV) DNA, whereas most cervical carcinomas (and cell lines derived therefrom) harbor significant quantities of HPV DNA.⁴⁰ For example, cervical carcinoma cell lines CaSki and HeLa are both HPV-positive and display highly correlated membrane glycosylation profiles (Figure S2), presumably due to mutual HPVinduced carcinogenesis. The cluster containing HPV-negative cervical carcinoma cell line C-33A and the two lung carcinoma cell lines, therefore, likely represents a grouping of cell lines for which carcinogenesis occurred due to largely genetic rather than environmental (e.g. viral) factors.

To distinguish amongst these individual cell lines, more detailed glycan profiling techniques were applied. Individual glycan species, identified and tracked by mass spectrometry, were employed as markers to differentiate amongst cell lines C-33A (cervical carcinoma), NCI-H358 (lung carcinoma), and A549 (lung carcinoma). Cervical carcinoma cell line C-33A was readily distinguished by elevated levels of glycans Man₄GlcNAc₂ and Man₃GlcNAc₂, which were present at 7-fold and 4-fold abundance, respectively, relative to the two lung carcinoma cell lines (Figure 6a). Lung carcinoma cell line NCI-H358, in turn, was readily distinguished from lung carcinoma cell line A549 by 6-fold elevated levels of glycans Man₃GlcNAc₂Fuc, Man₃GlcNAc₃Fuc, and Man₃GlcNAc₄Fuc (Figure 6b). Interestingly, all five marker glycans are biosynthetically related, each differing from the next by only one monosaccharide. Additionally, all represent relatively early steps in the glycan biosynthesis process, suggesting a lower degree of differentiation — one of the hallmarks of cancer glycosylation.^{2, 3}

CONCLUSION

We have demonstrated for the first time direct analysis of cancer cell membrane N-glycans by nano-LC/MS, revealing stark differences in the glycosylation of different cell types. The

platform is not only quantitatively reproducible, but also highly sensitive, with a large dynamic range.¹⁶ Additionally, porous graphitized carbon nano-LC provides structuresensitive separation of isomeric glycans.⁴¹ Though structural elucidation was not pursued in the present study, the platform's demonstrated structural separation capabilities may be easily coupled to tandem MS fragmentation for future glycan structure studies.^{31, 42, 43}

The observation of such stark differences in the membrane N-glycosylation patterns of different cancer cells reveals new avenues for the differentiation and diagnosis of cancer type/origin. Previous research on cancer cell membrane glycosylation focused mainly on O-glycans, perhaps as a matter of convenience — antibodies against specific O-glycans (such as sLe^X and sLe^A) are widely available, whereas few if any antibodies exist that target specific N-glycans.^{12, 29, 30} However, recent studies have shown that the vast majority of cell membrane glycosylation is in fact N-linked, rather than O-linked.²³ A shift of focus towards N-glycans, therefore, would provide a major boost in diagnostic biomarker candidates, especially now that a direct LC/MS analysis method has become available.

While the results of the present study provide a compelling argument for glycosylationbased cancer diagnosis, a crucial next step will be analysis of tumor samples from cancer patients. In our lab, preliminary experiments on patient T-cells (Figure 1) as well as microtomed epithelial tissue have already yielded glycan signals and profiles comparable to those of cultured cells, indicating the wide applicability of the membrane fractionation and LC/MS analysis procedures. Translation from cultured cell lines to patient tumors will provide an unprecedented glimpse of *in vivo* glycosylation, paving the way for a nextgeneration MS biopsy.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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Abbreviations

PGC	porous graphitized carbon	
ECC	extracted compound chromatogram	
Hex	hexose	
HexNAc	N-acetylhexosamine	
Fuc	fucose	
NeuAc	N-acetylneuraminic acid	
Man	mannose	
C/H	undecorated complex/hybrid	
C/H-F	fucosylated complex/hybrid	
C/H-S	sialylated complex/hybrid	
C/H-FS	fucosylated-sialylated complex/hybrid	

sLe ^X	sialyl Lewis X
sLe ^A	sialyl Lewis A
HPV	human papillomavirus

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Figure 1.

a) Extracted compound chromatograms of cell membrane N-glycans identified on non-CD4 T-cells from human blood. Colors denote the biosynthetic class of each glycan. Glycan structures are putative, based on known biosynthetic pathways. **b**) Relative abundances of each glycan biosynthetic class on non-CD4 T-cell membranes.



Figure 2.

Glycan class profiles of four cancerous breast cell lines (Hs-578T, ZR-75-1, MCF-7, and MCF-7/HER2) and one non-cancerous breast cell line (MCF-10A). Relative abundances are shown for each glycan biosynthetic class. **Inset:** Color-coded representation of the Pearson correlation coefficient (R) between each pair of cell lines, ranging from red (high correlation) to blue (low correlation), along with hierarchical clustering trees.



N-glycan type and decoration

Figure 3.

Glycan class profiles of four B-cell lymphoma cell lines (Raji, Ramos, NCI-H929, and BCBL-1) and one ovarian carcinoma cell line (ES-2). Relative abundances are shown for each glycan biosynthetic class. **Inset:** Color-coded representation of the Pearson correlation coefficient (R) between each pair of cell lines, ranging from red (high correlation) to blue (low correlation), along with hierarchical clustering trees.



Figure 4.

Scatterplot showing the relative abundances of each glycan signal in the Raji vs Ramos Bcell lymphoma cell lines. Each point represents an individual glycan signal. Pearson correlation coefficient (R) is calculated based on the best-fit line.



Figure 5.

Glycan class profiles of two lung carcinoma cell lines (NCI-H358 and A549) and three cervical carcinoma cell lines (C-33A, CaSki, and HeLa). Relative abundances are shown for each glycan biosynthetic class. **Inset:** Color-coded representation of the Pearson correlation coefficient (R) between each pair of cell lines, ranging from red (high correlation) to blue (low correlation), along with hierarchical clustering trees.



Figure 6.

Relative abundances of biosynthetically-related glycans with high fold differences between carcinoma cell lines C-33A (cervical), NCI-H358 (lung), and A549 (lung). **a**) Increased levels of Man₄GlcNAc₂ and Man₃GlcNAc₂ distinguish C-33A from NCI-H358 and A549. **b**) Increased levels of Man₃GlcNAc₂Fuc, Man₃GlcNAc₃Fuc, and Man₃GlcNAc₄Fuc distinguish NCI-H358 from A549. Glycans are shown in order of biosynthesis, from left to right.