DNA Encoded Libraries (DEGL) of Glycan Antigens to Detect Antibodies: An Approach Towards Next Generation Functional Glycomics

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DNA ENCODED LIBRARIES (DEGL) OF GLYCAN ANTIGENS TO DETECT ANTIBODIES: AN APPROACH TOWARDS NEXT GENERATION FUNCTIONAL GLYCOMICS

by

AISHWARYA PARAMESWARAN

Under the Direction of Peng George Wang, PhD

ABSTRACT

Structure and functional study of glycans are highly challenging due to the difficulties in analyzing glycans and limited availability of samples for study. These limitations could be resolved by attaching DNA barcode to the glycan, which virtually represent glycan in further application, by increasing the sensitivity of detection by polymerase chain reaction (PCR), requiring minimal samples for analysis. Assuming bigger arena of DNA Encoded Glycan Libraries (DEGL) in future, we propose here a method for uniquely coding all glycans using computer program that can convert the structural information of glycans to DNA barcode. A unique and universal coding for glycans will benefit both synthesis and analysis of DEGLs. As a proof of principle study, a small DNA Encoded Glycan Library (DEGL) of blood and globo series glycan antigen and its application was demonstrated in detecting blood group and breast cancer from plasma.

INDEX WORDS: DNA Encoded Glycan Library (DEGL), Glyco-PCR, Inverse Blood typing, Blood antigens, Globo Series Glycans, Click Chemistry
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AISHWARYA PARAMESWARAN

A Thesis Submitted in Partial Fulfillment of the Requirements for the Degree of Master of Science in the College of Arts and Sciences Georgia State University 2017
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by

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Electronic Version Approved:

Office of Graduate Studies
College of Arts and Sciences
Georgia State University
August 2017
DEDICATION

I would like to thank my family, for their intense support throughout the journey of my life so far. To my parents for their love, suggestions, and continuous support as always. They give me great comfort and encouragement whenever I’m depressed.

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To Arya, who helped me acquire knowledge and information, to enhance my skills and advance towards and set goals for my career.

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# List of Abbreviations

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<th>Abbreviation</th>
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<tr>
<td>GlcNAc</td>
<td>N-acetylglucosamine</td>
</tr>
<tr>
<td>Glc</td>
<td>Glucose</td>
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<tr>
<td>Gal</td>
<td>Galactose</td>
</tr>
<tr>
<td>Man</td>
<td>Mannose</td>
</tr>
<tr>
<td>Neu5Ac</td>
<td>N-acetylneuraminic acid</td>
</tr>
<tr>
<td>Fuc</td>
<td>Fucose</td>
</tr>
<tr>
<td>GalNAc</td>
<td>N-acetylgalactosamine</td>
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<tr>
<td>Xyl</td>
<td>Xylose</td>
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<tr>
<td>GlcA</td>
<td>Glucuronic acid</td>
</tr>
<tr>
<td>Ido</td>
<td>Idose</td>
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<tr>
<td>DEGL</td>
<td>DNA Encoded Glycan Library</td>
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<tr>
<td>DNA</td>
<td>Deoxyribonucleic Acid</td>
</tr>
<tr>
<td>RNA</td>
<td>Ribonucleic Acid</td>
</tr>
<tr>
<td>GBP</td>
<td>Glycan Binding Proteins</td>
</tr>
<tr>
<td>PCR</td>
<td>Polymerase Chain Reaction</td>
</tr>
<tr>
<td>qPCR</td>
<td>quantitative Polymerase Chain Reaction</td>
</tr>
<tr>
<td>RT-PCR</td>
<td>Real Time Polymerase Chain Reaction</td>
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<tr>
<td>GAG</td>
<td>Glycosaminoglycans</td>
</tr>
<tr>
<td>PG</td>
<td>Proteoglycans</td>
</tr>
<tr>
<td>CHO</td>
<td>Chinese Hamster Ovary</td>
</tr>
<tr>
<td>GP</td>
<td>Glycoprotein</td>
</tr>
<tr>
<td>NGS</td>
<td>Next Generation Sequencing</td>
</tr>
<tr>
<td>EDC</td>
<td>1-Ethyl-3-(3-dimethylaminopropyl)-carboiimide</td>
</tr>
<tr>
<td>NHS</td>
<td>N-hydroxysuccinimide</td>
</tr>
<tr>
<td>BoNT</td>
<td>Botulinum neurotoxin</td>
</tr>
<tr>
<td>CuAAC</td>
<td>Copper-catalyzed azide-alkyne Cycloaddition</td>
</tr>
<tr>
<td>SPAAC</td>
<td>Strain promoted alkyne-azide cycloaddition</td>
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<tr>
<td>A</td>
<td>Adenine</td>
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</table>
T       Thymine
C       Cytosine
G       Guanine
Cq      Quantification cycle
Ct      Threshold cycle
RBC     Red Blood Cells
a       alpha
b       beta
IUPAC   International Union of Pure and Applied Chemistry
µg      Microgram
µL      Microliter
µM      Micromolar
nM      Nano molar
pM      Picomolar
mM      Millimolar
Tm      Melting Temperature
°C      Degree Celsius
TBST    Tris Buffered Saline- Tween 20
kDa     kilodalton
RPM     Revolution Per Minute
G+DNA   Glycan + DNA
NTC     No Template Control
Ab      Antibody
Ag      Antigen
s       seconds
G       Glycan
HS      Healthy Serum
NS      No Serum
BC      Breast Cancer
1 INTRODUCTION

Glycans are important biomolecules along with proteins, lipids, and nucleic acids. Glycans can be divided into three main categories: O-linked glycans, N-linked glycans and glycolipids. Majority of glycans are made up of N-acetylgalactosamine (GlcNAc), Glucose (Glc), Galactose (Gal), Mannose (Man), N-acetylneuraminic acid (Neu5Ac), Fucose (Fuc), N-acetylgalactosamine (GalNAc), Xylose (Xyl), Glucuronic acid (GlcA), Idose (Ido).

Each cell in a multicellular organism is covered with dense and complex glycans. Most cellular secretions and extracellular matrices are found to be high in glycans and glycosylated. It is even seen that an envelope of a virus that bud from an infected cell to carry the glycosylation pattern of the host cell.

Glycans and glycoconjugates play a very important role in many areas of biological processes including cell-cell interactions, pathogenicity, mediating immunological processes, stabilizing proteins and cancer. Expression patterns of glycans are highly altered in diseases like cancer, atherosclerosis, diabetes, retrovirus infection, thrombosis, arthritis, neurodegeneration and many other diseases. Glycan structures are the important part of the antigenic determinants during microbial infections to which the host immune cells recognizes and mount immune responses against the infection. For example, breast cancer can be detected by Globo-H glycan, an important biomarker seen in breast cancer. Therefore, glycans have a pivotal role in molecular recognition and signaling, making sugar-receptor interaction at the structural level highly necessary; and this interaction of polysaccharide with receptors are major area of focus in academic and industrial research.

Functional Glycomics; a systematic study of the structures and functions of carbohydrates is yet to reach the level attained by the genomics and proteomics despite so much importance of...
glycans. Template driven biosynthesis of DNA-RNA-Protein by nature itself also known as the Central Dogma Theory of Molecular Biology made the researchers apply and adopt this knowledge in the synthesis and analysis of these molecules. Apart from this, the easy availability of experimentation methods and materials for the synthesis and analysis of DNA, RNA, and proteins made the researchers focus on genomics and proteomics and take up those areas to what they are today. However, such free-flowing information is not available for carbohydrate making functional glycomics exceptionally challenging. Since glycans are less organized without a template driven machinery, it is necessary to develop a specific method for synthesis and analysis for each glycan. Although with the recent advancement in technology allowed the development of solid phase automated and enzymatic synthesis, which reduced the difficulties in obtaining the glycans, but the materials and experimentations are still extremely difficult and complex especially for complex glycans. Therefore, there is a need for a development of versatile and ultra-sensitive detection technology as it is of high importance in the field of carbohydrate research for detection and characterization of glycan-protein interaction.

The methods now used for studying glycan-protein interaction are mainly carbohydrate microarrays,\textsuperscript{11,12} surface plasmon resonance,\textsuperscript{13} enzyme linked lectin assay,\textsuperscript{12} isothermal calorimetry,\textsuperscript{14} glycosylated conductive polymer,\textsuperscript{15} and crystallographic studies.\textsuperscript{16} In carbohydrate research, widely used method is microarray with carbohydrate derivatives by various means of immobilization technology.\textsuperscript{11,17-18} However, all the above methods have its own limitations, mainly because these techniques uses immobilized glycans that do not mimic the biological environment of binding. The detection also requires tagging of the target, glycan modifications and specific binding of antibodies, in addition, the interaction of glycans with targets are also very
weak, which makes the generation sugar library having a glycodendrimer on a slide with optimum distance for specific binding highly challenging.

DNA-encoded glycan libraries (DEGL) can be considered as a potential alternative or adjunct to the prevailing glycan detecting techniques. DEGL adopt DNA sequences specifically for each glycan in the library, which enable sensitive detection by signal amplification PCR and identifying by DNA sequencing. For functional glycomics study, DEGL of interest is constructed and interacted with specific proteins. Unbound glycans are removed by washing, and bound ones are amplified using quantitative Polymerase Chain Reaction (qPCR), also known as real time PCR (RT-PCR) and sequenced to reveal identities.

Next generation technologies in genomics extended the applications of DNA beyond genetics, the most prominent one being DNA encoding. The concept of DNA encoding was introduced in 1992, but it gained momentum recently with the advancement of Next Generation Sequencing (NGS) technologies.\textsuperscript{20-22} The major advantage of the technology is its efficiency in handling thousands of molecules with uniquely generated sequences and their high throughput analysis. The DNA tagging concept was initially applied in glycomics by Kwon and his coworkers, demonstrating Glyco-PCR, a method to detect Glycan-Binding Proteins (GBP) using DNA-conjugated glycan with high sensitivity.\textsuperscript{23-24} In one of the previous work reported by Kwon and his coworkers, they have shown glyco-quantitative polymerase chain reaction (Glyco-qPCR) assay platform that allows the ultrasensitive detection and quantification of glycan in biological samples. They used therapeutic carbohydrates like chondroitin sulfate (a GAG), and Proteoglycans (PG) from Chinese hamster ovary cells (CHO) separated by capillary electrophoresis, Glycoprotein obtained from PNGaseF treatment of decorin core Glycoprotein (GP). They used EDC/NHS chemistry to covalently link the DNA tag and glycan, and unbound DNA was washed using
streptavidin-coated magnetic beads as the glycans they used were biotinylated. They also did a proof of concept study, in GAG-binding proteins, where interaction was tested between Heparin-Antithrombin and Chondroitin-Antithrombin. With this work, they proved that glyco-qPCR can be used to analyze the glyco-interactome and by applying PCR amplification glycan as low as 1 zettamoles can be detected.\textsuperscript{23}

Later on, they also demonstrated glyco-PCR in the detection of active botulinum neurotoxin (BoNT). They used Sialyllactose, which interacts with binding-domain of BoNTs, sialyllactose-DNA conjugate was used as binding-probe for active BoNT and recovered through BoNT-immunoprecipitation. Glyco-PCR analysis of the bound sialyllactose-DNA is then used to detect low attomolar concentrations of BoNT and attomolar to femtomolar concentrations of BoNT in honey. They also did a comparative study through the dual binding of antibodies and monovalent or multivalent SL-DNA conjugates to toxins.\textsuperscript{24}

But these studies were limited to certain types of glycan and there was no rationale behind the selection of DNA sequences. Thus, developing a general platform for the DNA encoding of glycans and subsequent detection has a significance role to play in the functional glycomics. In this work, we demonstrated three fundamental aspects of DNA encoding of glycans, a universal methodology for the coding of glycans based on the structure of the glycan, an easy and efficient chemistry for the coupling of the DNA with the glycan and finally PCR and qPCR-based methods for the quantitative detection of the glycans.

1.1 Purpose of the Study

Glycans play a very important role in many biological processes including cell- cell responses, pathogenicity and immune responses etc. Despite so much importance, there is still a need for an efficient method for detection and analyzing glycan-protein interactions in the field of
carbohydrate research. Currently, carbohydrate microarray is successfully being used with the help of immobilization technology. But this also has its own limitations, most important one is its efficiency of mimicking the biological environment of the binding. Hence, we envision here developing a successful method for DNA-encoded Glycan Libraries (DEGLs).

There has already been reported evidence,23-24 for demonstrating Glyco-PCR, as a method to detect Glycan- Binding Protein (GBP). But these studies do not have any rationale behind the selection of DNA tags. Our aim is to develop a broadly accepted general platform for DNA encoding in glycomics. This could be achieved by proper selection of DNA codes, DNA-Glycan conjugation by click chemistry and method of detection of DNA-Glycan conjugates using qPCR.

This work attempted to address all three steps generating a DNA code specifically for each glycan by a developing a software, which could be used in future for any DEGL containing any specific glycans, generation of DNA tags for each element in the library by adding forward and reverse primer binding on each end for PCR amplification, and 5’ terminal alkyne modification to offer coupling with the glycan using click chemistry (CuAAC) and to develop a method of detection of DNA-Glycan conjugates using qPCR as a proof principle application study.

Generating a DNA code specific for each glycan-based on its structure was attained by developing a software using basic computing principles in a computer language Python. The development and the logic underlying in generating a DNA code specific for each glycan is described elaborately under experiment section later.

1.1.1 **Click Chemistry**

Lack of abundant samples to play with demands a near perfect conjugation chemistry for the synthesis of DNA glycan conjugates. We tested few of the well-known bioorthogonal reactions widely used in chemical biology, glycan analysis and detection many cell surface glycans.25-26
Well developed bioorthogonal reactions include Staudinger Ligation,\textsuperscript{27} Copper-catalyzed azide-alkyne Cycloaddition (also called as CuAAC),\textsuperscript{28} strain promoted alkyne-azide cycloaddition (also called as Cu-free click chemistry or SPAAC).\textsuperscript{29}

![Chemical Reaction Diagram]

\textbf{R1,R2 = Biomolecules, Bioactive Molecules, Fluorophores, Affinity Tags etc.}

\textit{Figure 1.1 Staudinger Ligation}

Staudinger ligation was the first bioorthogonal reaction reported by Staudinger and co-workers in 1919.\textsuperscript{27} This is bioorthogonal ligation that transforms azides to primary amines by phosphines.\textsuperscript{30} Later some modifications were made to this reaction, such as using modified reagent. Accompanied by those changes, Staudinger ligation suffered from slow kinetics and easily oxidized phosphine reagents during the reaction process, thus making the reaction not a perfect candidate for glycan analysis and detection.\textsuperscript{31}

The most popular and common bioorthogonal method is copper-catalyzed click chemistry.\textsuperscript{28,32} This is a copper-catalyzed cycloaddition between alkyne and azide to form a 5 membered ring.\textsuperscript{33} In early 2000’s, Sharpless group and Meldel group discovered that this reaction could be accelerated by copper dramatically, and this reaction can take place in an aqueous system.\textsuperscript{27,31} This copper-catalyzed azide-alkyne cycloaddition (CuAAC) is used widely in chemical biology, cell biology, cell surface glycan analysis and detection. This reaction being very simple and easy to perform it is widely been used by researchers both in industry and academia. During our study, we found most satisfying results with the click reactions and we adopted this chemistry for all our future work.
Another important part of our study is PCR, which is highly essential for the detection of DNA that is coupled with glycan which in turn symbolizes the glycan presence. The advent of the Polymerase Chain Reaction (PCR) radically transformed biological science from the time it was first discovered by Mullis in 1990. PCR based strategies have propelled huge scientific endeavors such as Human Genome Project. The technique is currently used by clinicians and researchers to diagnose diseases, clone and sequence genes, and carry out sophisticated quantitative and genomic studies in rapid and very sensitive manner. One of the most important medical applications of PCR method is the detection of pathogens. In addition, it is also widely used in forensics for the identification of criminals.\textsuperscript{46}

PCR can be performed using source DNA from a variety of tissues and organisms, including peripheral blood, skin, hair, saliva, and microbes. Only trace amounts of DNA are needed for PCR to generate enough copies to be analyzed using conventional laboratory methods. For this reason, PCR is a sensitive assay. Each PCR assay requires the presence of template DNA, primers, nucleotides, and DNA polymerase. The DNA polymerase is the key enzyme that links individual nucleotides together to form the PCR product. The nucleotides include the four bases – adenine, thymine, cytosine, and guanine (A, T, C, G) – that are found in DNA. These act as the building blocks that are used by the DNA polymerase to create the resultant PCR product. The primers in the reaction specify the exact DNA product to be amplified. The primers are short DNA fragments with a defined sequence complementary to the target DNA that is to be detected and
amplified. These serve as an extension point for the DNA polymerase to build on. The above-mentioned components are mixed in a test tube or 96-well plate and then placed in a thermal cycler that allows repeated cycles of DNA amplification to occur in three basic steps. The reaction solution is first heated above the melting point of the two complementary DNA strands of the target DNA, which allows the strands to separate, a process called denaturation. The temperature is then lowered to allow the specific primers to bind to the target DNA segments, a process known as hybridization or annealing. Annealing between primers and the target DNA occurs only if they are complementary in sequence (e.g. A binding to G). The temperature is raised again, at which time the DNA polymerase is able to extend the primers by adding nucleotides to the developing DNA strand. With each repetition of these three steps, the number of copied DNA molecules increases exponentially. The PCR product is analyzed using agarose gel electrophoresis, which separates the DNA product based on the size and charge. The two main methods used for visualization of DNA are staining the product using ethidium bromide dye, which intercalates between the two strands of the DNA. However, due to the prevailing carcinogenic and serious health effects of ethidium bromide, it’s use has been reduced and replaced by SYBR™, a safe DNA gel stain.

Figure 1.3 Key Components Of PCR
1) Denaturation: the reaction is heated to 95°C-98°C for 20-30 secs to break the hydrogen bonds; 2) Annealing: The reaction temperature is lowered to 50°C-65°C for 20-40 secs (preferably 2°C-4°C less than the Tm of primers) to allow the primers to anneal to the template strands; 3) Elongation: The temperature is increased (optimum temperature of DNA polymerase preferably 72°C-78°C) to allow the addition of dNTP’s. At end of each cycle, the target sequence is doubled.

The next generation advancement of the PCR technology was the development of RT-qPCR. While in end point, PCR requires that PCR products were detected and quantified by gel electrophoresis after completion of the reaction, real-time qPCR technology allows quantification of PCR products in “real time” during each PCR cycle, yielding a quantitative measurement of PCR products accumulated during the course of the reaction. Real-time reactions are carried out in a thermocycler that permits measurement of a fluorescent detector molecule, which decreases post-processing steps and minimizes experimental error. This is most commonly achieved through the use of fluorescence based technologies. Three main fluorescence based technologies are, probe sequences that fluoresce upon (i) hydrolysis (TaqMan; Applied Biosystems, Foster City, CA, USA) or hybridization (LightCycler; Roche, Indianapolis, IN, USA); (ii) fluorescent hairpins; or (iii) intercalating dyes (SYBR Green). As SYBR™ Green binding is not specific for a target sequence this system can be readily used for different gene assays, is flexible, inexpensive, and accurate results can be obtained provided validation of the specificity by melt curve (or dissociation curve) analysis. The TaqMan™ chemistry is more expensive than DNA binding dye assays, but the presence of the hydrolysis probe ensures that only specific amplicons are measured. In addition, multiplexing reactions are possible, although their set-up requires an important optimization phase.47,48

In brief, similar to end-point PCR, qPCR consists a succession of amplification cycles in which the template nucleic acid is denatured, annealed with specific oligonucleotide primers, and extended to generate a complementary strand using a thermostable DNA polymerase. This results in an exponential increase of amplicons (amplification products) that, in contrast with end-point
PCR, can be monitored at every cycle (in real time) using a fluorescent reporter. The increase in fluorescence is plotted against the cycle number to generate the amplification curve, from which a quantification cycle Cq (often described as Ct for cycle threshold) value can be determined. Cq corresponds to the number of cycles for which the amount of fluorescence (hence, of the template) is significantly higher than the background fluorescence. Therefore, the Cq value can be linked to the initial concentration of target nucleic acid and serves as a basis for absolute or relative template quantification. All steps of (RT-)qPCR may introduce experimental errors. qPCR is a robust technique, but due to its high sensitivity, very small variations can induce non-negligible differences in the results. To measure intra-assay variability, which follows a statistical distribution, RT-qPCR can be performed in triplicate (experimental replicates). Inter-assay variability can be estimated using a “reference” sample that will be included in each experiment. 

**Figure 1.4 Phases of qPCR.**

*The PCR process goes through three main phases as the number of cycles and the amount of product generated increase. Initially, when the amount of product is small and enzyme and reagents are not limiting, product generation is exponential and the reaction is closest to 100% efficiency. This exponential growth is hard to detect initially through real-time fluorescence because the amount of product is small. During the linear phase, products continue to accumulate, but the reaction efficiency begins to fall and reagents*
become limiting. Finally, in the plateau phase of the reaction, accumulation of product ceases as the reaction is exhausted for a number of different reasons.\textsuperscript{47}

For overall method development we adopted two systems, initially, we used the most known blood glycan antibody system (ABO) for the early stage method optimizations and, later we used the Globo-series glycan analogs as a small DNA encoded library for the proof of principle demonstration via detecting the breast cancer from patient plasma.

\textbf{1.1.3 ABO Blood Group System}

The discovery of the ABO blood group, over 100 years ago, caused great excitement. Until then, all blood had been assumed to be the same, and the often-tragic consequences of blood transfusions were not understood. The modern basis of the ABO blood group system was coined by Karl Landsteiner in 1900 when he discovered that the blood group of certain people agglutinated red cells present in other individuals.\textsuperscript{34} Ever since it has been studied extensively and well characterized with the structures of antigens and antibodies involved. The terminal carbohydrate structures present on the red blood cells and their antibodies present in the blood plasma determine the blood group.

The ABO blood group antigens remain of prime importance in transfusion medicine—they are the most immunogenic of all the blood group antigens. The ABO blood group antigens also appear to have been important throughout our evolution because the frequencies of different ABO blood types vary among different populations, suggesting that a particular blood type conferred a selection advantage (e.g., resistance against an infectious disease.)

The ABO blood group antigens are encoded by one genetic locus, the ABO locus, which has three alternative (allelic) forms—A, B, and O. A child receives one of the three alleles from each parent, giving rise to six possible genotypes and four possible blood types (phenotypes).
The four basic ABO phenotypes are O, A, B, and AB. After it was found that blood group A RBCs reacted differently to a particular antibody (later called anti-A1), the blood group was divided into two phenotypes, A1 and A2. RBCs with the A1 phenotype react with anti-A1 and makeup about 80% of blood type A. RBCs with the A2 phenotype do not react with anti-A1 and they make up about 20% of blood type A. A1 red cells express about 5 times more A antigen than A2 red cells, but both types of red cell react with anti-A, and A1 and A2 blood groups are interchangeable.

The immune system forms antibodies against whichever ABO blood group antigens are not found on the individual's RBCs. Thus, a group A individual will have anti-B antibodies and a group B individual will have anti-A antibodies. Blood group O is common, and individuals with this blood type will have both anti-A and anti-B in their serum. Blood group AB is the least common, and these individuals will have neither anti-A nor anti-B in their serum.

### Table 1.1 Possible Blood types in offspring

<table>
<thead>
<tr>
<th>ABO Alleles Inherited from Mother</th>
<th>ABO Alleles Inherited from Father</th>
<th>ABO Alleles</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>A</td>
<td>A</td>
</tr>
<tr>
<td>AB</td>
<td>B</td>
<td>AB</td>
</tr>
<tr>
<td>B</td>
<td>AB</td>
<td>B</td>
</tr>
<tr>
<td>O</td>
<td>A</td>
<td>B</td>
</tr>
<tr>
<td></td>
<td></td>
<td>O</td>
</tr>
</tbody>
</table>

### Table 1.2 Antibodies and Antigen present in ABO Blood Typing System

<table>
<thead>
<tr>
<th>RBC of different blood group</th>
<th>Antibodies in Serum</th>
<th>Antigen in RBC surface</th>
<th>Genotype</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Ab B</td>
<td>A</td>
<td>AO or AA</td>
</tr>
<tr>
<td></td>
<td>Ab A</td>
<td>B</td>
<td>BO or BB</td>
</tr>
<tr>
<td></td>
<td></td>
<td>A and B</td>
<td>AB</td>
</tr>
<tr>
<td></td>
<td></td>
<td>None</td>
<td>OO</td>
</tr>
</tbody>
</table>
The ABO locus is located on chromosome 9 has three main allelic forms: A, B, and O. The A allele encodes for a glycosyltransferase that produces the A antigen (N-acetylgalactosamine is its immunodominant sugar), and the B allele encodes for a glycosyltransferase that creates the B antigen (D-galactose is its immunodominant sugar). The O allele encodes for an enzyme with no function, and therefore neither A or B antigen is produced, leaving the underlying precursor (the H antigen) unchanged. These antigens are incorporated into one of four types of oligosaccharide chain, type 2 being the most common in the antigen-carrying molecules in RBC membranes.

For A/B antigen synthesis to occur, a precursor called the H antigen must be present. In RBCs, the enzyme that synthesizes the H antigen is encoded by the H locus (FUT1). In saliva and other bodily secretions, the enzyme that synthesizes the H antigen is encoded by the Se locus

Figure 1.5 ABO Blood antigen structures.
A similar method was developed to determine Globo-H expression in breast cancer cells. Globo H was found to be overexpressed on the cell surface of several epithelial cancers such as breast, colon, endometrial, gastric, pancreatic, lung, and prostate cancers. It is also moderately expressed in normal epithelial tissues (lung, breast, prostate, stomach, pancreas, and ovary), but its distribution is restricted to apical epithelial cells at lumen borders. In breast cancer Globo H expression was observed in >60% of ductal, lobular, and tubular carcinoma, but not in nonepithelial breast tumors. Globo H is not expressed in normal tissue except for weak expression

### Table 1.3 Distribution of various Blood Groups among Population

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Caucasians</td>
<td>44%</td>
<td>33%</td>
<td>10%</td>
<td>9%</td>
<td>3%</td>
<td>1%</td>
</tr>
<tr>
<td>Africans</td>
<td>49%</td>
<td>19%</td>
<td>8%</td>
<td>20%</td>
<td>3%</td>
<td>1%</td>
</tr>
<tr>
<td>Asians</td>
<td>43%</td>
<td>27%</td>
<td>Rare</td>
<td>25%</td>
<td>5%</td>
<td>Rare</td>
</tr>
</tbody>
</table>

#### 1.1.4 Globo-H

Similarly, eukaryotic cells are covered by a glycocalyx: an extensive network of polysaccharides. The glycocalyx serves as a rich source of binding sites for receptors and ligands, as well as pathogens and toxins. The mammalian glycome is estimated to consist of a few hundred unique glycan structures on glycoproteins and glycolipids. One such glycan is Globo H.

Some of the Globo-series glycans are overexpressed in certain cancers and are proven as biomarkers for early detection of breast cancer and ovarian cancer. Recent developments in the Globo-H immunotherapy also needs high sensitive methods to quantitatively detect the antiglombo-H antibodies for monitoring the therapy and in clinical trials.

Globo H is a glycosphingolipid of the Globo series with a sugar terminus resembling the blood group antigen H determinant. First identified in human teratocarcinoma and breast cancer cells, Globo H was found to be overexpressed on the cell surface of several epithelial cancers such as breast, colon, endometrial, gastric, pancreatic, lung, and prostate cancers. It is also moderately expressed in normal epithelial tissues (lung, breast, prostate, stomach, pancreas, and ovary), but its distribution is restricted to apical epithelial cells at lumen borders. In breast cancer Globo H expression was observed in >60% of ductal, lobular, and tubular carcinoma, but not in nonepithelial breast tumors. Globo H is not expressed in normal tissue except for weak expression.
in the apical epithelial cells at lumen borders, a site that appears to be inaccessible to the immune system. Thus, Globo H has been considered as an ideal target for immunotherapy of many epithelial cancers and indeed two phase I trials of a Globo H-based vaccine in breast and prostate cancer, respectively, have shown promising results.45

Functionally, Globo H has been associated with tumor stem cells, to be a potent inducer of angiogenesis, and an immunosuppressor through Notch signaling. The high Globo H expression by only cancer and cancer stem cells made it an attractive target for the generation of therapeutic cancer vaccines. These vaccines underwent the long history of development and improvement and are currently being tested in clinical trials for treatment. Globo H has therefore been considered as one promising tumor associated glycan biomarker, in particular for breast cancer.36

Figure 1.6 Globo-H Structure
2 EXPERIMENT

2.1 Systematic Coding of Sugars

The first aim was to develop a unique coding method that is consistent with the representation of glycans, so it can be used in future for any DEGL with ease and without any uncertainty in future glycomics. Unlike peptides, proteins and DNA/RNA, representation of carbohydrate brings many challenges. A detailed representation should feature composition, sequence, branching position, modifications, and anomeric configuration. The use of symbols to represent the glycans was first proposed by Kornfield et.al.\textsuperscript{41} This approach has replaced the use of IUPAC naming for representation of glycan, with modifications adapted timely to fit the entire glycome.\textsuperscript{42-43} All these representations are based on symbols and color codes to represent the building blocks.

Table 2.1 Glycan Representation (CFG)

<table>
<thead>
<tr>
<th>Symbolic Representation (Cartoon Representation CFG)</th>
<th>IUPAC Naming (Linear Code)</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image" alt="Glycan Representation" /></td>
<td>GalNAc b1-4 (Fuc a1-2) GlcNAc b1-4 Man a1-3 (GalNAc b1-4 (Fuc a1-2) GlcNAc b1-4 Man a1-6) Man b1-4 GlcNAc b1-4 (Fuc a1-6) GlcNAc</td>
</tr>
</tbody>
</table>

With the above approach, it is pragmatic to use nucleotide bases to represent the glycans. But this also brings prevailing challenges for using nucleotide bases for glycans as there are many symbols and color codes are available for representation, but there are only 4 nucleotides bases A, T, G, and C are available for coding. To have enough nucleotide codes to cover the entire the glycome we had to depend on permutation and combination approach. There are many sugar monomers which are incorporated in many ways in a glycan, making glycan structural information...
very complex. These inherit difficulties lead to the development of a dedicated computer program, which can manage the structural information of the glycan to coding dictionaries and transform the IUPAC name of glycan to single stranded DNA code. For easy acceptance of the codes, the codes should be of minimal length storing all the glycan information and enable both PCR and qPCR amplification.

For a perfect structure based coding, there must be a controlled vocabulary for all the structural components. For attaining this goal, the structural components of the carbohydrate were split into the structural components monosaccharides, linkages, and modifications etc. Three independent libraries were generated namely library A for monomer, library B for linkage and library C for modification. Next, all the monomers were gathered in library A and assigned each one with unique four-letter code. There are about 256 combinations generated using A, T, G, and C. About 70 monomers are added to this library as of now using the randomly generated codes. In a similar way, all possible linkages were accounted for library B, using codes based on 3 letters. Currently, there are 36 linkages in this library. For library C about 100 modifications were identified and incorporated in the library C and assigned with 4 letter codes which were not used in the library A. Special characters like ‘(‘ in the IUPAC naming for branching were assigned special codes. All these libraries were then incorporated into a Python-based program and the program was tested for about 100 glycans, for which the program successfully delivered the codes. The libraries can be extended and modified for addition as and when needed, to make this program a universal one and available for research community.
Table 2.2 Glycan Representation with DNA Codes

<table>
<thead>
<tr>
<th>Symbolic Representation (Cartoon Representation CFG)</th>
<th>IUPAC Naming (Linear Code)</th>
<th>DNA Code obtained through the program</th>
</tr>
</thead>
<tbody>
<tr>
<td>GalNAc b1-4 (Fuc a1-2) GlcNAc b1-4 Man a1-3 (GalNAc b1-4 (Fuc a1-2) GlcNAc b1-4 Man a1-6) Man b1-4 GlcNAc b1-4 (Fuc a1-6) GlcNAc</td>
<td>ACGCTGTTTTAGAAAAAGAAA AAGCTGTAATAAACCTTTACG CTGTGTTTAGAAAAAGAAAAAG CTGTAATAACAAAAATATG TAAGCTGTTTTAGAAAAAGAAA AAAGC</td>
<td>ACGCTGTTTTAGAAAA AAGAAAAAGCTGTA AAAAACTTTAC GCTGTTTTAG AAAAGAAAA AGCTGTAATAAAC AAAGC</td>
</tr>
</tbody>
</table>

A sequence length between 50 -100 is best for PCR applications. Among 100 glycans tested most fell within this limit, but some exceeded this. Through careful observation of the sample library, there were many repeating fragments in the glycans (e.g. Six sugar core of N-glycans). Hence, by taking this as a single block would reduce the length overall length of the codes of the long glycans, so a fourth library (Library D) was included in the program containing most of the common building blocks found in the glycans. Similar to other libraries, library D can also be extended, so as to cover the entire glycome and make it a universal one.

Table 2.3 Long and Short DNA codes of the Glycan

<table>
<thead>
<tr>
<th>IUPAC Naming (Linear Code)</th>
<th>Long DNA Code obtained through the program</th>
<th>Short DNA Code obtained through the program</th>
</tr>
</thead>
<tbody>
<tr>
<td>GalNAc b1-4 (Fuc a1-2) GlcNAc b1-4 Man a1-3 (GalNAc b1-4 (Fuc a1-2) GlcNAc b1-4 Man a1-6) Man b1-4 GlcNAc b1-4 (Fuc a1-6) GlcNAc</td>
<td>ACGCTGTTTTAGAAAAAGAAA AAGCTGTAATAAACCTTTACG CTGTGTTTAGAAAAAGAAAAAG CTGTAATAACAAAAATATG TAAGCTGTTTTAGAAAAAGAAA AAAGC</td>
<td>ACGCTGTTTTAGAAAA AAGAAAAAGCTGTA AAAAACTTTAC GCTGTTTTAG AAAAGAAAA AGCTGTAATAAAC AAAGC</td>
</tr>
</tbody>
</table>

Another probable issue encountered was with short glycans, which gives small codes of length less than 25. This will face practical difficulty while doing qPCR. To resolve this issue a random DNA sequence of length 20 was added to the code with a mention in the program to avoid
the misreading. These two modifications made sure to keep the sequences within the desired limits to be used in the qPCR.

Table 2.4 DNA code for short glycans

<table>
<thead>
<tr>
<th>IUPAC Naming (Linear Code)</th>
<th>DNA Code obtained through the program</th>
<th>Expanded DNA Code obtained through the program</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rha</td>
<td>ATAG Length – 4</td>
<td>ATAGCCCCCCAGTCAGGCCTAACGTA Length - 24</td>
</tr>
</tbody>
</table>

2.2 Coding and Decoding of glycans through Program

2.2.1 Coding

For getting nucleotide sequence, glycan sequence is given in the input box with spaces in between each character of the glycan sequence. When a glycan sequence is inputted the program looks for each character from left to right. It checks for match in monomer dictionary (Library A), linkage dictionary (Library B) and modification dictionary (Library C). If a match is found, the corresponding nucleotide is placed under the name ‘nuc’.

If the length of the nucleotide is less than 20, then an R nucleotide sequence is added to the ‘nuc’ to increase the length the nucleotide sequence and output is published.

If length of ‘nuc’ is more than 20 then the whole length of ‘nuc’ is searched for any matching sequences in the core dictionary (Library D). If there is match then a lengthy nucleotide core is replaced by a short 5 length sequence. This is done to reduce a lengthy nucleotide to short one for cost effectiveness. Reduced ‘nuc’ output is published.

2.2.2 Decoding

For decoding the nucleotide sequence back to glycan sequence, in the input box nucleotide sequence is given. The nucleotide sequence is searched for any matching five-length sequences in
the core dictionary (Library D). If there is any matching sequence then the length of nucleotide is extended to the original lengthy nucleotide sequence.

![Flowchart describing Coding](image)

*Figure 2.1 Flowchart describing Coding*

If length of nucleotide is more than 0; first four nucleotide sequence are taken and match is found in the carb dictionary (Library A) and corresponding carb is taken for the output. Next
three nucleotide sequences are taken and searched for a match in linkage dictionary (Library B); if there are no match then next four nucleotides are taken and searched in modification dictionary (Library C) and next three nucleotides are again searched for a match in linkage dictionary (Library C). Then next three nucleotides are taken and seen if it is ‘AAA’ or ‘TTT’, if it is so then a corresponding branching is taken for output. Then next three nucleotides are taken and seen if it is ‘AAA’ or ‘TTT’, if it is so then a corresponding branching is taken for output. This is done to cover the double branching cases. This continues for the complete length of the nucleotide sequence, before publishing the output.

Figure 2.2 Flowchart describing decoding
2.3 Materials

All chemicals and biological reagents were purchased from Thermo Fisher unless otherwise mentioned. Maxima SYBR Green/ROX qPCR Master Mix, Platinum Pfx DNA Polymerase were purchased from Life Technologies (Carlsbad, CA). Micro Bio-Gel P-30 Chromatography Columns were purchased from Bio-Rad (Hercules, CA). Tris[(1-Benzyl-1H-1,2,3-Triazol-4-yl) methyl] amine (TBTA) Click chemistry Ligand were purchased from TCI (Tokyo, Japan). MicroAmp 96 well Fast PCR Reaction Plate, MicroAmp Optical Adhesive Film, MicroAmp Fast Reaction Tubes, Strips were purchased from Applied Biosystems (Foster City, CA). All oligonucleotides were purchased from Integrated DNA Technologies (Coralville, IA). Blood Group A Antigen-Antibody HE-193 (MA1-19693), Blood Group B Antigen-Antibody HEB-29 (MA1-19691) and Blood Group ABH Antigen-Antibody HE-10 (MA1-19694) were purchased from Thermo Fisher.

2.4 Gradient PCR and Tm Determination

Gradient PCR was performed to figure out the optimal melting temperature to be used in the further quantitation experiments. Platinum Pfx DNA Polymerase was used for the PCR, approximately 26 µg of template, 1 µL of primers (10 µM), 1 mM dNTP, 10X amplifying buffer, MgSO₄, Polymerase and nuclease free water was used for a typical 25 µL reaction. Standard
thermocycling condition were used with different temperature during annealing phase of the cycle (95°C for 10 min, X°C for 30s, 95°C for 15s, 13 cycles) wherein X represents the annealing phase and temperature gradient from 50 °C - 60 °C were tested during this phase. The amplified PCR products were analyzed using 4% agarose gel electrophoresis. Tₘ determined using this method was used for qPCR.

<table>
<thead>
<tr>
<th>Component</th>
<th>25 uL</th>
<th>50 uL</th>
<th>Final Concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td>10X Amplifying Buffer</td>
<td>5 uL</td>
<td>10 uL</td>
<td>-</td>
</tr>
<tr>
<td>MgSO₄</td>
<td>1 uL</td>
<td>2 uL</td>
<td>-</td>
</tr>
<tr>
<td>dNTP(10mM)</td>
<td>0.75 uL</td>
<td>1.5 uL</td>
<td>0.3mM</td>
</tr>
<tr>
<td>Forward Primer(10uM)</td>
<td>0.75 uL</td>
<td>1.5 uL</td>
<td>0.3uM</td>
</tr>
<tr>
<td>Reverse Primer(10uM)</td>
<td>0.75 uL</td>
<td>1.5 uL</td>
<td>0.3uM</td>
</tr>
<tr>
<td>Template(1uM)</td>
<td>1 uL</td>
<td>2 uL</td>
<td>0.04</td>
</tr>
<tr>
<td>Polymerase</td>
<td>0.3 uL</td>
<td>0.5 uL</td>
<td>-</td>
</tr>
<tr>
<td>Nuclease-Free Water</td>
<td>15.4 uL</td>
<td>31 uL</td>
<td>-</td>
</tr>
</tbody>
</table>

2.5 Synthesis of Glycan(Antigen) – DNA Conjugates

Glycan(Antigen)- DNA Conjugates were synthesized using Azido-Alkyne cycloaddition click reaction. The 5’-/5Hexynyl-terminated DNA was procured from the commercial suppliers (IDT) with standard desalting purified. All glycans were synthesized via chemo enzymatic method with the azido propyl linker at the reducing end.

2.5.1 Click reaction procedure

5’-Hexynyl-terminated DNA (500µM, 20 µL, final concentration of 50 µM), azido glycan (500µM, 30 µL, final concentration of 75µM), 2M TEA buffer (pH 7, 20 µL, final concentration of 0.2M), 5 mM of freshly prepared Ascorbic acid solution (20 µL, final concentration 0.5mM), 10 mM of copper-TBTA in 55% DMSO (10 µl, final concentration 0.5 mM) and 50 % volume DMSO (100 µL) were mixed together in a tube. The reaction mixture was vortexed and kept at
room temperature for overnight. The reaction mixture was purified using Micro Bio-Gel P-30 Chromatography Columns (20 base pair cut-off). The concentration of the Glycan (Antigen)-DNA Conjugates was determined by absorbance at 260/280 using Nanodrop (ThermoFisher).

![Diagram of molecular structures]

*Figure 2.4 Click Conjugation of 5'-Hexynyl DNA and azido modified glycans*

### 2.6 General protocol for qPCR

#### 2.6.1 For a typical 12.5 µL reaction

The glycan(Antigen)-DNA Conjugates were added to 6.25 µL 2X Maxima SYBR Green/ROX qPCR Master Mix with 1 µL primers (Forward Primer – 500 nm and Reverse Primer- 300 nm). qPCR was performed with Applied Biosystems Stepone System (50 °C for 2 mins (Holding), 95°C for 10 mins (Holding), 95 °C for 15 s, 60°C for 10 s, 72 °C for 10 s for 40 cycles).

*Ct* or threshold cycle is a measurement of signal intensity for qPCR experiments. In a qPCR experiment, PCR is performed in presence of a fluorogenic intercalating dye (SYBR Green in our case). The dye intercalates with double-stranded DNA, as more double-stranded DNA is produced in each PCR cycle the dye increases the fluorescence intensity. Once the fluorescence intensity
reaches a threshold level, the cycle number is recorded by the instrument as Ct value. Therefore, sample having a large amount of DNA will have a lower Ct value compared to those samples containing relatively less amount of DNA.

<table>
<thead>
<tr>
<th>Component</th>
<th>Volume (in μL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2x Maxima SYBR Green/ROX qPCR Master Mix</td>
<td>6.25</td>
</tr>
<tr>
<td>Forward Primer(500nm)</td>
<td>1</td>
</tr>
<tr>
<td>Reverse Primer(300nm)</td>
<td>1</td>
</tr>
<tr>
<td>Template (Gradient as stated above)</td>
<td>1</td>
</tr>
<tr>
<td>Nuclease Free Water</td>
<td>3.25</td>
</tr>
</tbody>
</table>

2.7 Filtration of Antibody-Antigen DNA Conjugates

10 μL of glycan(antigen)-DNA Conjugate (5 μM) was incubated with 2 μL of antibody (if plasma is used then 2 μL of plasma is diluted to 8 μL using water) for 2 hours at room temperature. After two hours, the reaction volume was made up to 100 μL using TBST buffer. The reaction solution was washed using 100 μL TBST buffer for 7 times in 100kda cutoff centrifugation filtration filter tubes at 5000g for 10 mins. After washing the filtered and concentrated solution was used as a template for qPCR analysis.

2.8 Immunoprecipitation of Antibody-Antigen-DNA Conjugates

10 μL of glycan(antigen)- DNA Conjugate (5 μM) was incubated with 2 μL of antibody (if plasma is used then 2 μL of plasma is diluted to 8 μL using water) for 2 hours at room temperature. After two hours, the reaction volume was made up to 100 μL using TBST buffer and added to 20 μL of prewashed A/G protein bead and incubated for one hour at room temperature with occasional shaking at an interval of 15 mins. After one hour, the beads were washed thoroughly for 7 times using TBST buffer. After washing the beads were reconstituted in 20 μL of TBST buffer. The beads were directly used a template for qPCR analysis.
2.9 Blood Collection

Blood was collected from 10 volunteers by finger pricking, and collected to heparin-coated blood collection tubes. The collected blood was centrifuged at 5000 rpm for about 5 mins for obtaining plasma from blood. The collected plasma is stored at -20°C until use.
3 RESULTS AND DISCUSSION

In this study, I have developed a program which can generate DNA code specifically for each glycan (described in systematic coding of sugars under experimental section). Next steps were to successfully couple DNA and glycan using click chemistry, develop methods for glycan and protein binding, and analyze the glycan using qPCR.

3.1 DNA sequences

Using the developed Python-based program DNA sequences for blood group antigens and Globo glycan series were obtained.

Table 3.1 DNA sequences
For all 5’ to 3’ DNA Sequences BOLD is the forward primer and BOLD ITALIC is the reverse primer binding site.

<table>
<thead>
<tr>
<th>Name</th>
<th>5’Mod</th>
<th>Sequence</th>
<th>3’Mod</th>
</tr>
</thead>
<tbody>
<tr>
<td>O DNA</td>
<td>5Hexynyl</td>
<td>AATGATACGGCGACCACCGAAAGAAAAGACGATCTAGACTCTTACGATGTAAGACGACCACCGAAACGCACTTTTAGAAAAGAGAAAACGATCTCTACGATGTAACACGTCGAAACTCCAGTCAC</td>
<td>None</td>
</tr>
<tr>
<td>A DNA</td>
<td>5Hexynyl</td>
<td>AATGATACGGCGACCACCGAAACGCACTTTTAGAAAAGAGAAAACGATCTCTACGATGTAACACGTCGAAACTCCAGTCAC</td>
<td>None</td>
</tr>
<tr>
<td>B DNA</td>
<td>5Hexynyl</td>
<td>AATGATACGGCGACCACCGAAACGCACTTTTAGAAAAGAGAAAACGATCTCTACGATGTAACACGTCGAAACTCCAGTCAC</td>
<td>None</td>
</tr>
<tr>
<td>Gb-H</td>
<td>5Hexynyl</td>
<td>AATGATACGGCGACCACCGAAAGAAAAGACGATCTACGCTACGATGTAAGAAGACGATCTACGCTACGATGTAACACGTCGAAACTCCAGTCAC</td>
<td>None</td>
</tr>
<tr>
<td>Gb-5</td>
<td>5Hexynyl</td>
<td>AATGATACGGCGACCACCGAAAGAAAAGACGATCTACGCTACGATGTAAGAAGACGATCTACGCTACGATGTAACACGTCGAAACTCCAGTCAC</td>
<td>None</td>
</tr>
<tr>
<td>Gb-4</td>
<td>5Hexynyl</td>
<td>AATGATACGGCGACCACCGAAAGAAAAGACGATCTACGCTACGATGTAAGAAGACGATCTACGCTACGATGTAACACGTCGAAACTCCAGTCAC</td>
<td>None</td>
</tr>
<tr>
<td>Gb-3</td>
<td>5Hexynyl</td>
<td>AATGATACGGCGACCACCGAAAGAAAAGACGATCTACGCTACGATGTAAGAAGACGATCTACGCTACGATGTAACACGTCGAAACTCCAGTCAC</td>
<td>None</td>
</tr>
<tr>
<td>Gb-2</td>
<td>5Hexynyl</td>
<td>AATGATACGGCGACCACCGAAAGAAAAGACGATCTACGCTACGATGTAAGAAGACGATCTACGCTACGATGTAACACGTCGAAACTCCAGTCAC</td>
<td>None</td>
</tr>
<tr>
<td>Bb-4</td>
<td>5Hexynyl</td>
<td>AATGATACGGCGACCACCGAAAGAAAAGACGATCTACGCTACGATGTAAGAAGACGATCTACGCTACGATGTAACACGTCGAAACTCCAGTCAC</td>
<td>None</td>
</tr>
<tr>
<td>Bb-3</td>
<td>5Hexynyl</td>
<td>AATGATACGGCGACCACCGAAAGAAAAGACGATCTACGCTACGATGTAAGAAGACGATCTACGCTACGATGTAACACGTCGAAACTCCAGTCAC</td>
<td>None</td>
</tr>
<tr>
<td>Bb-2</td>
<td>5Hexynyl</td>
<td>AATGATACGGCGACCACCGAAAGAAAAGACGATCTACGCTACGATGTAAGAAGACGATCTACGCTACGATGTAACACGTCGAAACTCCAGTCAC</td>
<td>None</td>
</tr>
<tr>
<td>FW Primer</td>
<td>None</td>
<td>AATGATACGGCGACCACCGAA</td>
<td>None</td>
</tr>
<tr>
<td>RV Primer</td>
<td>None</td>
<td>GTGACTGGAGTTCAAGACGTG</td>
<td>None</td>
</tr>
</tbody>
</table>
The obtained DNA sequences were flanked with primer binding domains and 5’ Hexynyl modification were procured from IDT.

3.2 Tm Determination

Tm was determined using the protocol for gradient PCR and Tm Determination (under experimental section) for further quantitation experiments. After PCR to determine the optimum temperature the PCR products were analyzed in using 4% agarose gel.

![Figure 3.1 Tm Determination](image)

*Figure 3.1 Tm Determination*

Gel analysis of the PCR product (Figure 3.1) showed the optimum temperature to be in a range of 52°C-60°C. 60°C was used as annealing temperature for further experiments.

3.3 Glycan (Antigen) – DNA Conjugates

After Tm determination, the 5’-Hexynyl terminated DNA and azido glycan were coupled together using click chemistry according to the protocol mentioned under synthesis of Glycan(Antigen) – DNA Conjugates (under experimental section).

A PCR comparison was conducted between pure DNA and G+DNA to make sure that the DNA can be used for further PCR reaction after conjugation with glycan. PCR was conducted
under the same conditions as stated under Gradient PCR and Tm Determination and analyzed using 4% agarose gel electrophoresis.

From the gel analysis of the PCR product (Figure 3.2) a clear concrete band was visible in both DNA and G+DNA, which made sure that the DNA can be used for the further analysis after conjugation with the glycan.

To make sure that the Glycan-DNA conjugate can be used for binding the protein with the glycan and DNA can be used for PCR analysis, any presence of residual unreacted DNA was further tested by PCR amplification and gel electrophoresis. Pure DNA_A and G+DNA_A was incubated with the specific antibody for two hours and washed with the washing buffer, and later eluted with the elution buffer. Both wash and elution buffers were collected and performed a PCR. Gel analysis of the PCR product (Figure 3.3) did not show a band in the wash buffer (residual unreacted DNA) but a strong band was seen in elution buffer indicating near perfect efficiency of coupling protocol.
3.4 qPCR Reactions and Standard Curve

The success of DEGL lies in the readiness of applying the DNA codes to the standard PCR and qPCR protocols. It is important to have the glycan-coupled DNA (G+DNA) achieve similar PCR efficiency to the native DNA, so compared the G+DNA conjugates with the corresponding DNA strands to verify the efficiency of amplification in both PCR and qPCR. In all the above sections, PCR was performed on the both DNA and G+DNA using standard thermocycling with different annealing temperatures and Immunoprecipitation methods. Next, experiment was carried out to determine the qPCR detection limit of both pure DNA and G+DNA. Both pure DNA and G+DNA were serially diluted to provide different concentrations of templates, with final concentrations ranging from 2.4 nM to 16 pM. Standard curve qPCR was carried out to obtain the corresponding Ct values, Critical threshold value (Ct) is generally used to compare the concentration of the templates in the qPCR reaction mixture, a low Ct indicating high template concentration. Ct values of the conjugates from 2.4 nM to 16 pM concentrations were observed in the range of 5-25, while the negative control (no template control, NTC) gave a Ct value above 30 (Figure 3.4). The test was not conducted for further low concentrations worrying about the
interference of primer dimer formation, a potential challenge in amplifying templates smaller than 100 bases. Nevertheless, a standard plot of the Ct value vs log concentration of the DNA and G+DNA conjugate (Figure 3.5) did show the linear relation indicating the successful application of G+DNA for quantitative detection up to the picomolar level. Both DNA and DNA-Glycan conjugates gave similar amplification, assuring that the DNA tag can be an alternative to detect the glycan itself. Encouraged by the result clearly suggesting the detection of glycans from even a picomolar level, proceeded to the final goal, screening of glycans against glycan binding proteins.

Figure 3.4 qPCR limit of detection comparison of glycan DNA conjugate and pure DNA
A & C amplification plot and Ct value plot of glycan DNA conjugate; B & D amplification plot and Ct value plot of pure DNA
3.5 Blood Glycan – Antibody Sensitivity study: Filtration

Since G+DNA could be applied in both screening and detection, DEGLs enable interrogation of the glycans against target proteins in solution, while known glycan-DNA interactions could be used to detect a specific target with high sensitivity. Two set of experiments were done for validating these two aspects of DEGL; initially demonstrated the detection of specific glycan binding proteins by using the familiar blood glycan-antibody interactions. As a proof of principle study to demonstrate the significance of DNA Encoded Glycan Library (DEGL), the blood group antigen-antibody interactions were chosen.

Structures of the O (or H), A, and B antigens are well known (Figure 3.6). This system was selected for the study mainly due to two reasons, i) It is most studied and accepted among all the glycan interactions, and ii) They offer the opportunity to study both the sensitivity and specificity of the technique because of the structural similarities. Three blood group antigens A, B and O sugars were synthesized chemo-enzymatically. Synthesis of glycan-DNA conjugate was achieved via click chemistry. A, B and O glycans were coupled with the corresponding 5’ alkyne modified DNA (DNA code + Primers).
Figure 3.6 Blood Group ABO antigen-DNA conjugate structures
All three G-DNA conjugates (G1+A DNA, G2+B DNA and G3+O DNA, with G1, G2 and G3 representing A, B, and O glycan antigens, respectively) were interrogated with the commercially-procured human blood antibodies (IgM, mouse). Concentrations of 5 µM, 2.5 µM, and 1 µM of the G+DNA were incubated with 2 µL of the antibodies; unbound DNA was eliminated by filtration with 100k cut off centrifuge tubes. The filtrate after incubating with the antibodies were subjected to qPCR assay, and Ct values of the G+DNA were compared with the Ct values of negative control where antibody was omitted from the incubation. A Ct difference of more than 10 points was observed at all three G+DNA concentrations indicating the requirement of low concentration of glycans for the accurate results. The filtration results were consistent with the concentration of antigens used, and were found to be more sensitive (Figure 3.7). The higher
sensitivity of the filtration method may have been due to the size difference of IgM antibodies and DNA strands (900 kDa and 25 kDa, respectively). From the above experiment, it was proven that DEGL can be used for detection of specific glycan binding proteins by using the familiar blood glycan-antibody interactions over a range of concentration (5 µM, 2.5 µM, 1 µM). With the success of above experiment, 2.5 µM was used for further demonstration of selectivity and specificity of the glycan binding proteins by using blood glycan-antibody interactions.

Next aspect of DEGL is to screen for the glycan binding proteins and, for addressing this mixed all three antibodies (1:1:1) and interrogated this mixture with each of the G+DNA conjugates (2.5 µM) separately.

As seen in figure 3.8a, Ct values of 8.4 and 7.7 respectively for G1+A DNA and G2+B DNA indicate the high affinity binding of A and B glycans with their corresponding antibodies Ab-A and Ab-B. But a similarly tested G3+O DNA showed relatively higher Ct value compared to both A and B conjugates but still less than that of the no antibody controls indicating weak interaction of antigens with the Ab-O.
Figure 3.7 Comparison of Filtration Sensitivity Study

A) G1+A DNA against the antibody A at different concentrations (5 µM, 2.5 µM, 1 µM); B) G2+B DNA against the antibody B at different concentrations (5 µM, 2.5 µM, 1 µM); C) G3+O DNA against the antibody O at different concentrations (5 µM, 2.5 µM, 1 µM); D) Ct values plot for G+DNA at different concentrations (5 µM, 2.5 µM, 1 µM).

This result indicates that DEGL could also work with many closely related target proteins around without any loss of selectivity. The specificity of the method was also tested, for this investigated a single G+DNA (G2+B DNA) against the three antibodies, a clear amplification was visible only when both Ab B and G2+B DNA were in the mix and all the remaining tests were poorly amplified. This result clearly indicates that the amplification is indeed from the target
specific binding of glycan and there is no significant interference of nonspecific binding of DNA and target proteins.

![Figure 3.8 Filtration study with G+DNA conjugates and Antibody](image)

A) G+DNA conjugates against the mixture of antibodies having equal concentration of Ab-A, Ab-B and Ab-O B) G2+B DNA against each antibody separately

3.6 Blood Glycan – Antibody Interaction Comparison: Filtration and Immunoprecipitation

The next step was to try with natural blood serum, for this blood was collected using finger pricking as stated in blood collection (under experimental section). Since the natural blood has IgG antibodies, a comparison between filtration and immunoprecipitation was conducted to find the best method. For interrogating this aspect G+DNA at concentration of 2.5 µM was incubated with 2 µl of diluted plasma (2 µl of plasma was diluted with 8 µl water) with two different blood plasma (Blood Groups O and B). The unbound DNA was washed by both filtration and immunoprecipitation methods. The filtrate/eluent after incubating with the blood plasma (washed of unbound DNA) were subjected to qPCR assay, and Ct values of the G+DNA were compared with the Ct values of negative control where blood plasma was omitted from the incubation. A Ct difference of more than 10 points was observed at filtration results. The filtration results were consistent with the concentration of antigen used (figure 3.7), and were found to be more sensitive. The higher sensitivity of the filtration method may have been due to the size difference of IgM
antibodies and DNA strands (150 kDa and 25 kDa, respectively). A Ct difference of about 10 points was observed at immunoprecipitation results. The immunoprecipitation was consistent with the concentration of antigen used, and was found to be more accurate and specific from the amplification curve (Figure 3.10). The higher accuracy and specificity of the immunoprecipitation may have been due to the compatibility of beads with the qPCR. Ct value of 20.3 in G1+A DNA and 22.2 in G2+B DNA (Immunoprecipitation) and Ct value of 4.5 in G1+A DNA and 6.8 in G2+B DNA (Filtration) was obtained with blood group B (Figure 3.9).

Similarly, Ct value of 17.6 in G1+A DNA and 18.4 in G2+ B DNA (Immunoprecipitation) and Ct value of 4.9 in G1+A DNA and 4.8 in G2+B DNA (Filtration) was obtained with blood group O. A lower Ct value in G1+ A DNA with Blood Group B and almost same Ct value in G1+A DNA and G2+B DNA with Blood Group O proved this method can be used for inverse blood group typing.
Figure 3.10 Amplification plot comparison
A) Amplification Plot Blood Group B Immunoprecipitation; B) Amplification Plot Blood Group O Immunoprecipitation; C) Amplification Plot Blood Group B Filtration; D) Amplification Plot Blood Group O Filtration.

3.7 DEGL: Real Life experiments – Inverse Blood Typing

From the successful results with two real blood experiments, blood typing experiments were conducted by collecting samples from healthy volunteers. Identifying the serum antibody present in the blood plasma will help to identify the blood group, person having the A antibodies alone in the blood have B antigen (group B), person with B antibodies alone are of A blood group, those who have both the antibodies belongs to the O and no antibodies in the blood indicating AB group (Table 3.2). This type of blood typing is called as inverse blood typing. However, the
conventional blood typing used widely is based on agglutination which is a direct blood typing using antigens (glycan antigen) on the surface of the RBC’s.

**Table 3.2 Blood Group Antigens and Antibodies**

<table>
<thead>
<tr>
<th>RBC of different blood group</th>
<th>A</th>
<th>B</th>
<th>AB</th>
<th>O</th>
</tr>
</thead>
<tbody>
<tr>
<td>Antibodies in Plasma</td>
<td>Ab B</td>
<td>Ab A</td>
<td>Ab (-)</td>
<td>Ab A Ab B</td>
</tr>
<tr>
<td>DEGL (G1+A DNA, G2+B DNA)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Results (Ct values)</td>
<td>A B</td>
<td>A B</td>
<td>A B</td>
<td>A B</td>
</tr>
</tbody>
</table>

Few microliters of blood were collected from 10 healthy volunteers using a finger prick, collected blood were centrifuged to separate the plasma and cellular components. The blood plasma was used to detect the blood group, 4 uL of the plasma was diluted to 20 uL and from this 2 uL each was incubated with A and B antibodies for 2 hours to allow the formation of G+DNA-Ab complexes. Further, G+DNA-Ab complexes were allowed to interact with the protein A/G magnetic beads for 1 hour, separated using a magnetic stand and washed seven times to discard any unbound G+DNA. After thoroughly washing the G+DNA-Ab-Bead complex, it was diluted to 10 uL with water and 1 uL was directly used as a template in the qPCR. Ct values of the G+DNA in the blood plasma was compared with the Ct value of G+DNA with no plasma. Among the 10 samples tested, two samples showed low Ct values for both G1+A DNA and G2+B DNA indicating the presence of both antibodies and hence belonged to O group. Five of the samples had a low Ct value for G2+B DNA means A group and three of the samples had a low Ct value for the
G1+A DNA implying B group (Figure 3.11). All the results were accorded with the conventional blood typing. The above results indicated that the DEGL can be used for inverse blood typing, which can be an alternative for conventional blood typing. However, using this method the rhesus group cannot be diagnosed as it is a protein antigen-antibody interaction.

3.8 Globo Series Library

After the successful completion of library synthesis and qPCR method development using blood glycans and inverse blood typing, next the concept was applied to Globo glycans. Here DEGL and glyco-PCR were used to profile the Globo-glycan binding antibodies and there by detecting the cancers expressing these antigens. Globo-glycans are part of the glycosphingolipids and their expression pattern is well correlated with the cancer metastasis and progression. Aberrant expression of Globo-glycans will also prompt the body to generate the anti-glycan antibodies targeting these epitopes. These antibodies are generally observed in the sera even before the cancer progression to the late stages and effectively act as biomarkers for certain types of cancer including breast, lung, prostate and ovarian cancer.\textsuperscript{7,36,45}
Figure 3.12 Globo glycan structures and Globo-glycan conjugates

Globo-H and its truncated analogs Gb-5, Gb-4, Gb-3, and Gb-2 were synthesized from the enzymatic extension of chemically synthesized Gb-2. To test the terminal epitope specificity alone; Bb-4, Bb-3 and Bb-2 were also synthesized via chemo-enzymatic strategy.

Initially, tested with the Globo-H glycan conjugate alone with the commercial anti Globo-H human antibody VK-9 in the same way as with the blood glycans, and the results are shown in the figure 3.13. Two concentrations of Globo-H glycan-DNA conjugate (2.5 µM and 5 µM) were incubated with 2 µL of VK-9 antibody and compared the Ct value with the no antibody control experiments. As expected, the amplification and Ct values were correlated with the concentration of the antigens present in the solution, a low Ct value of 7 and 19 was observed for 5 µM conjugate and comparatively higher Ct value of 14 and 21 was obtained for 2.5 µM conjugate.
Detection of glycan binding antibodies present in the cancer plasma

After gaining the confidence from the Globo-H and VK-9 interrogation studies, we conducted an elaborate study using all the Globo series glycan against two breast cancer plasma samples (sanguine bioscience BC1 and BC2), and compared the Ct values of each glycan conjugates with the sample treated with both healthy serum (HS) and no serum (NS). The results show that only GbH, Bb4 and Bb3 had significant amplification with the Ct values less than 8, while other glycan conjugates show relatively less amplification (Figure 3.14 and 3.15).
Figure 3.14 Ct Value Plot of different Globo glycan conjugate series with breast cancer serum (BC1 and BC2), Healthy Serum (HS) and No Serum (NS).

These results are consistent with the micro array based methods reported by Chi-Huey Wong group.

Though this approach accurately differentiated the cancer plasma from the healthy plasma based on the presence of glycan binding antibodies, it is not convenient to test every glycan conjugate separately and it is not practical when several plasma samples are needed to be tested. But this difficulty can be easily overcome by employing next generation sequencing (NGS), which allow mixing of the glycan conjugate and mixing of plasma samples with one additional step of barcode indexing. Hence NGS technology allows the high throughput detection of hundreds of plasma samples with even thousands of glycans in the DEGL.
Figure 3.15 Amplification Plot of different Globo glycan conjugate series with breast cancer serum (BC1 and BC2), Healthy serum (HS) and No serum (NS)
4 CONCLUSION

With the advancement of the technologies in the DNA synthesis and sequencing have brought the enormous potential of DNA in the applications beyond genetics. DNA encoded libraries (DEL) are of the modern concept in drug screening and lead optimization. This study recognized the importance of DEL technologies in the functional glycomics (DEGL), which is otherwise limited to the low sensitive detection techniques and highly difficult to obtain samples. Applying the highly sensitive qPCR detection will enhance the reach of functional glycomics to low nano gram samples and high throughput screening. Here a unique method has been described to code each glycan, accounting their structural information very similar to how proteins coded with triplet codons. The program which is available for the use of research community is aimed to provide and keep a uniformity in the codes for every single glycan in the future DEGL technologies.

A strict library of DNA codes will not only reduce the confusion in the community but also bring down the synthetic expenses using optional bulk synthesis. Along with the unique DNA codes, it also produces an agreement on the coupling chemistry and primer selection to have the ready-made glycan codes for research use. Also identified copper assisted alkyne-azide coupling (CuAAC) chemistry, also known as ‘click chemistry’ for the best result in because most DNA suppliers readily offer azide and alkyne modifications on their library.

In a proof of principle study, generated a small DEGL consisting the common blood antigens and their detection in solution were demonstrated. With the same principle did a blood group analysis with the DEGL to confirm the use of DEGL in detecting the glycan binding antibodies. Although inverse blood typing can’t replace the gold standard conventional direct blood typing based on the surface antigen, this study can be further extended to the next level
where the blood typing will be based on the analysis of P1PK, FORS, Se, Le and I blood glycan antigens. All the blood glycan antigens (ABO, P1PK, FORS, Se, Le and I) will form a complete blood glycan group typing. Some of them being highly rare antigens, this kind of analysis will make a place along with molecular blood typing and other rare antigen blood typing apart from the conventional ABO typing.

Finally, another DEGL consisting the Globo glycan series was generated and their detection in solution and breast cancer serum were demonstrated to confirm the use of DEGL in detecting the glycan binding antibodies which are of highly significant in the diagnostic applications. Here, Globo-glycans were analyzed in cancer serum which can be extended to several other diseases like Crohn’s disease, Rheumatoid arthritis, cancers and infection where disease-specific glycan biomarkers and determinants can be found. This application of DEGL can be promising for the analysis of disease-specific glycan biomarkers and determinants for early diagnosis of the disease.

With the successful results from the proof of principle study namely inverse blood typing and detection of Globo-glycans in cancer samples, it has been proven that the DEGL will be an efficient method in future years to come. DEGL will find its own space along with other prevalent technologies available.
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