Acute Lyme disease IgG N-linked glycans contrast the canonical inflammatory signature

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Lyme disease (LD) infection is caused by *Borrelia burgdorferi sensu lato* (Bb). Due to the limited presence of this pathogen in the bloodstream in humans, diagnosis of LD relies on seroconversion. Immunoglobulins produced in response to infection are differentially glycosylated to promote or inhibit downstream inflammatory responses by the immune system. Immunoglobulin G (IgG) N-glycan responses to LD have not been characterized. In this study, we analyzed IgG N-glycans from cohorts of healthy controls, acute LD patient serum, and serum collected after acute LD patients completed a 2- to 3-week course of antibiotics and convalesced for 70-90 days. Results indicate that during the acute phase of Bb infection, IgG shifts its glycosylation profile to include structures that are not associated with the classic proinflammatory IgG N-glycan signature. This unexpected result is in direct contrast to what is reported for other inflammatory diseases. Furthermore, IgG N-glycans detected during acute LD infection discriminated between control, acute, and treated cohorts with a sensitivity of 75-100% and specificity of 94.7-100%.

KEYWORDS
Lyme disease, IgG N-glycans, glycosylation, immunoglobulin, diagnostics - clinical characteristics, biomarker

Introduction

In 1975, a cluster of children and adults in the small community of Lyme, Connecticut experienced unusual arthritic symptoms. Today, Lyme disease (LD) is the most prevalent vector-borne disease in the US, with an estimated 476,000 annual cases (1). The causative agent, *Borrelia burgdorferi sensu lato* (Bb), is a spirochete transmitted...
from the *Ixodes* tick into the human host dermis during its blood meal (2). The spirochete leaves the blood and disseminates into multiple organ systems in as little as two weeks post-infection (3–5). Disseminated LD is more challenging to diagnose, and delayed treatment can lead to long-term disability or death (6, 7). The disease is endemic in the Northeastern US, and incidence rates continue to rise. The US’s annual treatment and diagnostic cost is over 4.8 billion USD (8).

Antibiotic treatment for LD in the acute phase is often curative (9–12). However, untreated patients and a subset of treated patients progress to disseminated disease (13, 14). Disseminated disease can result in facial nerve palsy (15), Lyme Carditis (16), Lyme Arthritis (17), Lyme Neuroborreliosis (18, 19), and long-term disability (20, 21). Persistent symptoms are reported by 10-20% of patients diagnosed and treated during the acute phase of LD. Persistent symptoms include joint pain, fatigue, and neurocognitive deficits (10, 22). This highlights the need for an accurate early diagnosis and the ability to track disease resolution.

Current diagnosis is complicated because testing relies on indirect methods. Direct PCR and blood culture methods often fail due to the spirochete’s limited presence in the bloodstream, low bacterial counts in circulation, slow replication cycle, the requirement for complex growth media, and specialized microscopy requirements (23, 24). Hence, indirect methods that rely on the patient’s serological response are the principal method of confirming an infection (25). These indirect methods of acute LD diagnostics based on ELISA and western immunoblot technologies suffer from low sensitivity and a high false-positive rate. Thus, while advances in LD detection research are being made (26–29), clinicians currently lack a sensitive method to diagnose early disease. Furthermore, clinical assays are unable to determine treatment efficacy, track disease resolution (30) or diagnose subsequent infections.

Serum protein glycosylation is often altered during inflammatory and autoimmune diseases. The glycosylation profile of immunoglobulins is dynamic and offers a novel immunologic insight into the host’s response (31–34). Glycosylation is the most abundant complex post-translational serum protein modification (35) and plays a significant role in protein structure and function in *vivo* (36, 37). IgG has a well-characterized N-glycosylation site on the constant fragment (Fc-Asn-297) region (38). This site contains complex biantennary glycans with varying degrees of galactose, bisecting N-acetyl-glucosamine, and sialic acid residues. Most notably, the glycans are highly core-fucosylated. IgG glycosylation is dynamic and the glycans present can affect the binding avidity to various Fcγ receptors, rate of complement activation, and release of cytokines (39–42). Previous studies have profiled IgG N-glycans in sera obtained from patients with inflammatory disease (Table 1). Reports indicate a trend for IgG N-glycan reduction in terminal galactose and sialic acid content during inflammatory diseases and this glycan signature is linked to a pro-inflammatory phenotype (52–58).

Scientists are beginning to understand how glycosylation reflects health status and influences protein structure and function. IgG glycosylation moieties are associated with specific functions. In general, lowered galactose and sialic acid residue content are reported in inflammatory states (Table 1), and conversely, increased galactose is associated with an anti-inflammatory state. Hence, to improve immunotherapy outcomes, pharmaceutical companies glycoengineer monoclonal antibodies with specific glycosylation features to treat cancers and chronic diseases. For example, the effectiveness of intravenous immunoglobulin (IVIG) therapy for autoimmune conditions including Gillian-Barre, Immune Thrombocytopenic Purpura, or Kawasaki disease patients is associated with the increased sialic acid content on the Fc N-glycans of the IVIG (59–61). In addition, therapeutic monoclonal antibodies are glycoengineered to contain specific a-fucosylated, agalactosylated N-glycans to promote superior half-life and treatment efficacy (62–65). IgG N-glycan modulation of the immune response during disease reveals another layer of physiologic crosstalk. Evidence indicates that the repertoire of N-glycans present on IgG produced in response to vaccines is dependent on many factors, including age, inflammatory state, and the type of adjuvant (58).

It is well accepted that N-glycans modulate the function of antibodies and are altered in disease states (66). We hypothesize that the IgG N-glycan profile of LD patients will reflect the immunological response to the acute LD infection. Thus, this first report of total IgG N-glycans associated with LD will provide insight into the inability of the host immune system to resolve Bb bacterial infection.

**Material and methods**

**Patient samples**

Serum samples (Table 2) were obtained from the Bay Area Lyme Disease Biobank and stored at ~80°C. Serology was determined at Stony Brook University and the Bay Area Lyme Disease Biobank (67). Human subject research IRB requirements were met (IRB #1808006553). Acute LD patients presented with erythema migrans (EM) rash(es) and donated their blood at the time of clinical diagnosis. Acute LD patients were included in the cohort when their acute LD diagnosis was confirmed using either two-tiered serological studies or PCR identification. Convalescent draws were obtained after patients completed their prescribed course of 14-21 days of antibiotics and convalesced for 70-90 days without further symptoms. We refer to the convalescent cohort as “treated”. In the pilot study, serum was pooled into three patient cohorts. Seven healthy controls were pooled, 5 acute LD patients were pooled, and 3 treated LD patients were pooled. The resulting three samples were analyzed within the pilot study using HPLC and the
GlycoTyper MALDI-TOF methods. In the subsequent confirmatory study using the GlycoTyper method, data was collected from each individual patient: healthy control (n=18), acute (n=18), and patient-matched treated (n=18). Further demographic details for the patient cohorts are presented in Supplemental Table 1.

HPLC IgG N-glycan analysis

IgG was purified from 5 μL of pooled serum using Protein A/G UltraLink Resin (Thermo Scientific, MA) according to the manufacturer’s directions. IgG Heavy Chains (Fc region) were isolated using 1D gel electrophoresis, stained with Coomassie stain and the 50kDa band was excised. Following gel de-staining, the glycans were enzymatically removed and fluorescently labeled following standard in-gel PNGase F and labeling protocols as previously described (68, 69). The labeled N-glycans were combined with 100% Acetonitrile (30:70) in an HPLC-compatible vial. Fluorescently labeled glycans were subsequently analyzed by high-performance liquid chromatography (HPLC) by using a TSK-Gel Amide 80 column (Tosoh Bioscience LLC). The mobile phase consisted of solvent A (50 mmol/L ammonium formate, pH 4.4) and solvent B (acetonitrile). The gradient used was as follows: a linear gradient from 20% to 58% solvent A at 0.4 mL/min for 152 min followed by a linear gradient from 58% to 100% solvent A for the next 3 minutes. The flow rate was increased to 1.0 mL/min, and

### TABLE 2 Patient Samples.

| Sample Group | Sample ID | Age | Sex | Ethnicity | Treatment | Serum Collection |
|--------------|-----------|-----|-----|-----------|-----------|-----------------|
| Pilot Set    | Control   | 46.4| 42.9| 42.9      | 0         | 100             | Not Applicable  |
| Pilot Set    | Lyme disease | 51.4| 40.0| 40.0      | 0         | 100             | Yes            |
| Confirmatory Set | Control | 51.3| 40.9| 22.7      | 18.2      | 81.8            | Not Applicable  |
| Confirmatory Set | Lyme disease | 53.1| 40.0| 20.0      | 20.0      | 80.0            | Yes            |

Healthy Control and Lyme disease LD with matched Acute and Post-treatment Demographics of Bay Area Lyme Disease Biobank Serum Samples. Sample ID numbers are linked to demographic details including age, sex, ethnicity, and if the patient donated a post-treatment convalescent serum sample.
the column was washed in 100% solvent A for 5 minutes. Following the wash step, the column was equilibrated in 20% solvent A for 22 minutes in preparation for the next sample run. HPLC analysis was performed using the Waters Alliance HPLC System, complemented with a Waters fluorescence detector, and quantified by the Millennium Chromatography Manager (Waters Corporation). Glycan structures were identified by calculating the glucose unit and GlycoStore database as previously described (70). The HPLC method overview is provided in Supplemental Figure 1A.

Removal of sialic acids for HPLC and MALDI comparison

Due to the inherent challenges of detecting glycans containing sialic acid using mass spectrometry, HPLC data was collected using samples that were desialylated to allow a direct comparison with the MALDI-FT-ICR method. Beginning with 13μL of the labeled and chromatographically clean 2-AB glycans, 4μL 5X pH 6.0 Enzyme Buffer was added before pipette mixing 3μL of Sialidase A (ProZyme (now AdvanceBio), OH). The 20μL final volume was incubated at 32°C for 12 hours. Next, the 20μL solution was added to a 10K MWCO concentrator column (Corning, NY) and centrifuged at 12,000 rpm for 10 minutes. The flow-through was collected and combined with an additional 25μL of dH2O before being added back to the 10K MWCO column and centrifuged at 12,000 rpm. This process of collecting the flow-through and combining it with additions of 25μL of dH2O was repeated to serially enrich the labeled N-glycans while filtering out the sialidase enzyme. The sample was subsequently dried down via SpeedVac holding a vacuum at -28inHg with out additions and re-suspended in 30μL dH2O. Sialidase-treated samples were then analyzed as outlined for non-sialidase-treated sample in HPLC.

MALDI-FT-ICR mass spectrometry N-glycan analysis

SolarisX Legacy 7T FT-ICR mass spectrometer equipped with a Matrix-Assisted Laser Desorption/Ionization (MALDI) (Bruker) analysis of total IgG N-glycans is detailed in the literature (48, 71–73). In brief, 1μL serum diluted in 99μL 1X PBS was incubated with 0.2mg/ml spotted anti-IgG capture antibodies (Bethyl Laboratories Inc., Tx, Cat. Number A80-104) treated with Sialidase A (produced in-house by MUSC), sprayed with PNGase F (produced by N-Zyme, PA) to release N-glycans from captured targets, coated with a matrix, and analyzed for glycan abundance at specific m/z peaks by MALDI-FT-ICR MS using SCI淡淡的 Lab software 2022a (Bruker). A capture antibody treated with PBS served as a blank to subtract the N-glycans released from the capture antibody from the final analysis. A MALDI-FT-ICR (referred to as MALDI) method overview is provided in Supplemental Figure 1B and is also referred to as the “GlycoTyper” method.

Statistical analysis

One-way ANOVA analysis with post-hoc Tukey’s multiple comparisons test was employed to examine the triplicate datasets for statistically significant differences between cohorts. P<0.05 was considered statistically significant and figures are denoted as having *p<0.05, **p<0.01, ***p<0.001. Figures and statistical analysis were completed using GraphPad Prism 8. Receiver Operating Characteristic (ROC) curves analyzed individual N-glycan abundance levels for diagnostic utility. Results are described using the Oxford nomenclature (74).

Results

We demonstrate that total serum IgG N-glycosylation of acutely infected Lyme disease patients contrast with the typical pro-inflammatory signature often found on other inflammatory diseases. First, a proof-of-concept study was performed using pooled serum to identify the IgG N-glycan signature of healthy control, acute LD, and patient-matched antibiotic-treated LD serum using high-pressure liquid chromatography separation (HPLC) and detection of fluorescently labeled glycans (75). The samples were subsequently analyzed using the recently developed GlycoTyper MALDI method (73) which pairs a specific total-IgG capture antibody with subsequent MALDI-FT-ICR imaging (MALDI). The trends in the glycan signatures were reproducible between both platforms, and thus we proceeded to analyze a larger confirmatory set of individual serum samples using the GlycoTyper platform.

Acute LD IgG N-glycans gain terminal galactose and sialic acid

HPLC analysis of the glycan signature revealed several statistically significant differences between control and acutely infected patients (Figure 1). Individual glycans were quantitated as a percent of the total glycan profile and compared across cohorts using one-way ANOVA (Figure 1A). Twoagalactosylated glycan species, F(6)A2G0 and F(6)A2BG0, decreased in the acute and treated pooled cohorts when compared to controls. There was also an observed decrease in the mono-galactosylated F(6)A2G1 glycan. In addition, we observed significant increases in three glycan species. N-glycans containing terminal di-galactose, with and without core fucose increased significantly (A2G2, F(6)A2G2), as did the core fucosylated mono-sialylated glycan (F(6)A2G2S1). Next, the abundance of IgG N-glycans with specific characteristics were summed to analyze the glycans by class. IgG
N-glycans containing bisecting GlcNAc, core-fucose, no galactose, a mono- or digalactose, or a mono- or di-sialylated were respectively summed to present N-glycan class data (Figure 1B). The decrease in the smaller agalactosylated class and an increase in the terminally galactosylated and sialylated glycans was observed. However, bisecting and core-fucosylated N-glycan classes did not significantly vary across cohorts. The only statistical difference observed between the acute and treated patient cohorts was a continued increase in the presence of terminal galactose.

**MALDI-FT-ICR and HPLC detect similar trends of IgG N-glycans in LD**

IgG N-glycans were desialylated and the HPLC analysis was compared to the MALDI-FT-ICR method. The IgG N-glycans were grouped by terminal sugar moiety into 4 glycan classes. These groups showed the same trends when using both platforms (Figure 2). Significant reductions in agalactosylated N-glycans and significant increases in terminal galactose were identified in the HPLC (Figure 2A) and MALDI glycan analysis platforms (Figure 2B). Both methods indicate there is no significant difference in the abundance of bisecting or core-fucosylated N-glycans when comparing controls, acute and treated patient cohorts. The reproducible shifts in IgG N-glycan abundance within the desialylated glycan classes promoted the use of a larger confirmatory set of LD serum to be analyzed in a high-throughput manner using the MALDI method.

**MALDI analysis identifies IgG N-glycosylation signatures of patients with LD**

IgG N-glycans from a confirmatory set of healthy control, acute LD, and acute re-drawn serum post-antibiotic treatment patient sera were examined. MALDI analysis of the desialylated IgG N-glycans identified statistically significant shifts of N-glycan species (Figure 3). Several significant differences were detected when comparing control and acute LD. Acute LD patients exhibit a significant decrease in the core fucosylated agalactosylated N-glycan F(6)A2G0 when compared to controls (Figure 3A). Conversely, there was a statistically significant increase in the core fucosylated di-galactosylated N-glycan F(6)A2G2. IgG N-glycans terminating in total galactose or digalactose significantly increased during acute LD compared to healthy controls (Figure 3B). There was no significant difference in the total bisecting or core-fucosylated N-glycans when comparing Acute to healthy control IgG.

We observed a continued perturbation of IgG glycosylation in the post-treatment cohort that does not return towards
healthy control baselines for many detected N-glycans. Treated LD patients maintained a significant decrease in the agalactosylated F(6)A2G0 observed in acute LD (Figure 3A). An additional significant drop was seen in two glycan structures, F(6)A2G1 and F(6)A2BG2, when compared with control and acute. Several glycans showed significant increases. Increases that were limited to the treated cohort include the A2G1 and F(6)A2BG1 structures. The F(6)A2G2 continued to increase above the already elevated acute LD cohort level. Grouping the treated timepoint IgG N-glycans revealed a drop in total core-fucose as well as a decrease in the digalactosylated grouped N-glycans (Figure 3B). There was no difference observed for the total bisecting N-glycans between acute and treated timepoints.
These findings align with previous assays from the pooled pilot study experiments. Comparisons of IgG N-glycosylation dependence on age, sex, location of collection, or ethnicity were assessed. There was no statistically significant difference within or between cohorts when applying these metrics (data not shown) with one exception. Healthy control males had 2.3% higher F(6)A2G0 N-glycan profiles compared to their healthy control female counterparts. While IgG N-glycan signatures change during ageing and between sexes, the age- and sex-matched cohorts permitted comparisons across and between the cohorts (76).

IgG N-glycans discriminate between healthy control and LD timepoints

Seroconversion is often employed as a LD diagnostic, yet there is great need for a more sensitive, early indicator of disease. Moreover, serological assays for LD cannot differentiate between a patient with acute LD compared to a patient that has recently convalesced from LD. Figure 4 reports the efficacy of selected IgG N-glycans to discriminate between healthy and LD patient cohorts. Thresholds for listed N-glycan classes were determined using receiver operating characteristic (ROC) curve. The performance of the discrimination was reported in a confusion matrix (Supplemental Figures 2-4). Healthy controls are differentiated from acute serum samples using four N-glycan classes: F(6)A2G0, F(6)A2G2, percent total terminal galactose, and percent terminal di-galactose; resulting in 75% sensitivity, 100% specificity, and 85.7% accuracy. Healthy controls are differentiated from treated LD patient serum using the N-glycan classes: F(6)A2G0, A2G1, F(6)A2G1, F(6)A2G2, F(6)A2BG2, percent total fucose, and percent total terminal galactose; resulting in 100% sensitivity, 94.7% specificity, and 97.3% accuracy. Lastly, acute LD serum is differentiated from the treated serum using the N-glycan classes: A2G1, F(6)A2G1, F(6)A2G2, F(6)A2BG1, F(6)A2BG2, total percent fucose, and percent G2 galactosylation; resulting in 100% sensitivity, specificity, and accuracy.

Lyme disease subverts IgG response – working hypothesis

Results presented in Figure 5 show the increased terminal galactose observed on IgG N-glycans detected during LD within the context of the humoral immune response. IgG N-glycans detected in acute LD patients contain significantly higher levels of galactose and conversely lower agalactosylated structures. Most IgG N-glycans responding to inflammatory diseases present increased agalactosylated N-glycans to promote downstream pro-inflammatory immune responses. Thus, the LD IgG N-glycans may be induced through the Borrelia burgdorferi’s disruption of the germinal center and antibody response.

Discussion

This is the first report of total human IgG N-glycan profiles during early and treated LD. Our initial studies detected an unanticipated shift in IgG N-glycan profile which was confirmed and expanded upon using a larger set of samples. During many inflammatory states (Table 1), IgG N-glycan profiles respond...
with reduced terminal sialic acids and galactose and increased GlcNAc sugar exposure (39). This agalactosylated IgG phenotype has been demonstrated to promote pro-inflammatory responses and may aid in clearing the infection (52–58, 77, 78). In the case of LD, total IgG N-glycan profiles had the opposite trend. LD increases the galactose and sialic acid content of IgG N-glycans while further decreasing the terminal GlcNAc exposure (Figures 1–4). The result of such a shift requires further investigation, but in theory could contribute to ineffective host responses to LD infection by reducing humoral immune activation (77, 78). The ability to detect these changes within total IgG suggests that LD induces a large shift in the IgG N-glycan composition. Thus, this dysregulated immune response may at least in part explain why human IgG produced during an initial LD infection is not effective to clear the current infection or protect from a future re-infection with LD.

The pilot study served to compare the well-established HPLC IgG N-glycan analysis method (79) to the novel MALDI analysis method (73) using pooled cohorts of control, acute LD, and treated LD serum. Due to MALDI’s limited detection of unmodified sialic acids, samples were treated with a sialidase (72). The results presented in Figure 2 demonstrate that both methods detect comparable individual identified N-glycans with similar trends of terminal galactose exposure induced during LD and maintained during the post-antibiotic treatment time points. To confirm the pooled pilot study findings, three cohorts of 18 individual samples were analyzed in triplicate using the MALDI method. The GlycoTyper method was selected to analyze the confirmatory cohort due to its significantly higher throughput compared to HPLC. We found that the demographics of the age- and sex-matched patient cohorts did not lead to inherent differences in IgG N-glycosylation.

While the MALDI results confirmed the original trends observed in the pooled pilot study, a higher number of samples analyzed led to further trends emerging (Figures 3A, B). Agalactosylated structures are reduced while total N-glycans terminating with a galactose moiety continued to increase within the acute and treated cohorts as indicated within the pooled pilot experiments. N-glycans containing a core-fucose are markedly lowered in the antibiotic-treated cohort which suggests an increase in Antibody-Dependent Cellular Cytotoxicity (ADCC) abilities of post-treatment IgG (80). The increased level of digalactosylated (G2) N-glycans on IgG at the acute stage and subsequent return to healthy control levels is a potential biomarker reflecting the host’s response to successful antibiotic treatment. Following antibiotic therapy for Tuberculosis (TB), IgG N-glycan digalactose content returned to healthy control levels (43). Grace et al. found G2 N-glycans decreased during acute TB and subsequently increased after effective antibiotic treatment. In the case of LD, IgG N-glycans increase G2 content during infection and return to healthy control levels after effective antibiotic treatment (Figure 3B). This once more converse trend observed in LD suggests a subverted immune response during LD compared to TB infection. The implication of the IgG N-glycan response to LD is portrayed in Figure 5.

LD patients are discriminated from healthy controls with a high degree of sensitivity (75-100%) and specificity (94.7-100%) using total IgG N-glycan measurements (Figure 4). Blinded analysis of acute LD, post-treatment LD, and mimic diseases using these methods should be completed to validate these findings. Future LD tests incorporating total IgG N-glycan analysis could increase acute LD diagnostic sensitivity and track subsequent antibiotic treatment responses.

Mechanisms controlling the dynamic B-cell glycosyltransferase expression operating on IgG N-glycans during LD require examination. B-cell glycosyltransferase expression is known to respond to the cytokine micro-environment (81). Recent studies detected IgG N-glycosylation profiles are impacted by the type of adjuvant present during vaccination (82). Additionally, during the
onset of autoimmune disease, TH17 cells signal newly differentiated B cells with IL-22 and IL-23 to regulate glycosyltransferase expression, resulting in a pro-inflammatory agalactosylated, non-fucosylated IgG N-glycan (83). The increase in terminal galactose of IgG N-glycans during acute LD may be attributed to an upregulation of naïve B-cell beta-1,4-galactosyltransferase expression (66, 84–86).

Future studies should determine if LD induces a specific cytokine signaling pathway to affect the glycosyltransferase expression of plasma cells or B cells during LD infection.

Lyme disease has been demonstrated to destroy germinal centers of lymph nodes during early infection (77, 78, 87, 88). These germinal centers are a vital structure that produces long-lived immunoglobulin responses through T-cell-dependent interactions (89). The destruction of the germinal center may explain why patients are liable to become re-infected with LD after treatment. Additionally, Bb has been demonstrated to gain entry inside human endothelial cells (90), alter innate immune responses after phagocytosis (91), suppress lymphocytes growth rate (92), and down-regulate major-histocompatibility complexes expression on Langerhans cells (93). Any one of these effects could be responsible for altering the IgG N-glycan profile during LD. For example, antigens presented in a T-cell independent manner promote increased sialylation on the Fc portion of IgG leading to an immunotolerant, less pro-inflammatory IgG N-glycan repertoire (94). Interestingly, murine studies have demonstrated a reduction in the general humoral response after LD infection as indicated by low titers of anti-viral antibodies produced post-vaccination and Bb-impaired helper T-cell mediated affinity maturation (87, 95). Lastly, because the LD cohort completed 2-3 week of oral doxycycline therapy, the serum N-glycome responses could have been altered in part due to changes in gut flora (96, 97).

Conclusion

Using the GlycoTyper MALDI-FT-ICR imaging approach, we detected an unexpected IgG N-glycan signature in humans during LD. We have demonstrated the IgG N-glycans produced during a Lyme disease infection lack the classic highly agalactosylated signature associated with most inflammatory diseases. Instead, LD induces IgG N-glycans containing larger, terminally galactosylated sugar moieties. Moreover, many IgG N-glycans detected at the acute LD timepoint remain altered from healthy control levels at the post-antibiotic treatment timepoint. Of note, the digalactose content of IgG N-glycans in the treated time point was comparable to the healthy baseline levels after antibiotic treatment. Furthermore, we detected a significant decrease in total core-fucosylation at the treated timepoint suggesting a possible increase in ADCO which may aid in clearing Bb from the host. IgG N-glycans offer numerous biomarkers, reflect acute disease state, response to treatment, and may improve the sensitivity of the acute diagnosis of LD above the current two-tiered testing protocol. This first examination of IgG N-glycan signatures associated with LD requires future study.

Data availability statement

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

Ethics statement

The studies involving human participants was reviewed and approved by the University of Drexel College of Medicine Review Board. This study involves secondary research using de-identified data and biospecimens not collected specifically for this study. The IRB 1808006553 was assigned an exempt status.

Author contributions

BH-G and MC were involved in conceptualization of assay, assay design, data analysis, and manuscript writing. BH-G was responsible for statistical analysis. AM and SG were involved in assay design and data analysis. BW was involved in editing the manuscript. All authors contributed to the article and approved the submitted version.

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Conflict of interest

Author AM is a founding partner in GlycoPath, Inc. and holds patents for the GlycoTyper methodology.

The remaining authors declare that the research was conducted in the absence of any commercial or financial interests.

Conclusion

Using the GlycoTyper MALDI-FT-ICR imaging approach, we detected an unexpected IgG N-glycan signature in humans during LD. We have demonstrated the IgG N-glycans produced during a Lyme disease infection lack the classic highly agalactosylated signature associated with most inflammatory diseases. Instead, LD induces IgG N-glycans containing larger, terminally galactosylated sugar moieties. Moreover, many IgG N-glycans detected at the acute LD timepoint remain altered from healthy control levels at the post-antibiotic treatment timepoint. Of note, the digalactose content of IgG N-glycans in the treated time point was comparable to the healthy baseline levels after antibiotic treatment. Furthermore, we detected a significant decrease in total core-fucosylation at the treated timepoint suggesting a possible increase in ADCO which may aid in clearing Bb from the host. IgG N-glycans offer numerous biomarkers, reflect acute disease state, response to treatment, and may improve the sensitivity of the acute diagnosis of LD above the current two-tiered testing protocol. This first examination of IgG N-glycan signatures associated with LD requires future study.
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Supplementary material

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fimmu.2022.949118/full#supplementary-material
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