Glycophenotypic Alterations of Urinary Protein as New Potential Diagnosis Indicators for Diabetic Nephropathy

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Received 17 July 2016; Revised 24 December 2016; Accepted 4 January 2017; Published 19 March 2017

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Diabetic nephropathy is a major cause of chronic kidney disease and end-stage kidney disease. However, so little is known about alterations of the glycopatterns in urine with the development of diabetic nephropathy. Presently, we interrogated glycopatterns in urine specimens using a lectin microarray. The results showed that expression levels of Sia\(\alpha\)2-6Gal/GalNAc recognized by SNA exhibited significantly increased tendency with the development of diabetic nephropathy; moreover, SNA blotting indicated glycoproteins (90kDa, 70kDa, and 40kDa) in urine may contribute to this alteration. Furthermore, the glycopatterns of (GlcNAc)\(2\)–4 recognized by STL exhibited difference between diabetic and nondiabetic nephropathy. The results of urinary protein microarray fabricated by another 48 urine specimens also indicated (GlcNAc)\(2\)–4 is a potential indictor to differentiate the patients with diabetic nephropathy from nondiabetic nephropathy. Furtherly, STL blotting showed that the 50kDa glycoproteins were correlated with this alteration. In conclusion, our data provide pivotal information to monitor the development of diabetic nephropathy and distinguish between diabetic nephropathy and nondiabetic renal disease based on precise alterations of glycopatterns in urinary proteins, but further studies are needed in this regard.

1. Introduction

Diabetic nephropathy (DN) is a serious complication of diabetes mellitus and now the major cause of chronic kidney disease (CKD) and end-stage kidney disease (ESRD) throughout the world. However, renal damage associated with diabetes can be classified as either DN or nondiabetic renal disease (NDRD). There are significant differences in the treatment and prognosis of these two types of renal disease. Renal pathological diagnosis is currently the gold standard for distinguishing DN from NDRD, although renal biopsy is invasive.

Urine has some unique advantages (collected noninvasively, a large number of proteins in urinary proteome also belong to plasma proteome); certain proteins in urine not only reflect the status of the entire body, but also correlated with diseases, which is good resource for the analysis of disease processes [1–4]. Valuated urinary proteome analysis is a tool for prediction and monitoring of DN. Transferrin is a glycoprotein with a molecular weight of 76.5 kDa, which was demonstrated that the urinary transferrin could reflect the degree of interstitial fibrosis and tubular atrophy in patients with DN [5]. The low-molecular weight proteome in urine was profiled by capillary electrophoresis-coupled mass spectrometry (CE-MS), and results indicated that the specific collagen fragments in urine could effectively assess DN risk at an early stage [6]. For type 2 diabetic nephropathy, it is reported that urinary cystatin C and NAP (nonalbumin protein) could...
reflect the progression of type 2 diabetic nephropathy [7]. Liver-type fatty acid-binding protein (L-FABP), an intracellular carrier protein of free fatty acids, is expressed in the liver and kidney, which could serve as a predictive marker for renal and cardiovascular prognosis in type 2 diabetic patients without advanced nephropathy [8]. It is interesting that L-FABP also could be an independent predictor of progression at all stages of type 1 diabetic nephropathy irrespective of stage [9].

Glycosylation is an important posttranscription modification; more than half of known proteins are glycoproteins. It is well known that the glycan attached on glycoproteins has many biological functions, such as maintaining conformation and flexibility of protein, oligosaccharide modification involved in biological processes including cell adhesion, signal transduction, and endocytosis. Abnormal glycosylation had demonstrated correlation with occurrence and development of diseases [10, 11]. Lectin microarray is a powerful technology that uses a panel of lectins immobilized on solid phase substrate for high-throughput analysis of glycans and glycoproteins related to diseases such as cancer and inflammatory diseases [12–14]. Alterations of sialylated MUC1 glycosylation detected by lectin microarray could be indispensable for the development of cholangiocarcinoma [15]. Glycosylated patterns of A1AT (alpha-1-antitrypsin) may serve as potential biomarkers for early detection [16]. For DN, nonenzymatic glycosylation product such as glycated albumin was strongly associated with incident diabetes and its microvascular complications. In addition, the glycated albumin: HbA1c ratio and glycated albumin may be superior to HbA1c associated with the presence of DN [17, 18]. However, there were few reports about enzymatic glycosylation on glycoproteins that altered in development of DN.

In this study, we preformed lectin microarray to investigate altered glycopatterns during the development of DN and evaluate whether glycopattern in urine was a potential biomarker to differentiate DN from NDRD.

2. Materials and Methods

2.1. Study Population. Our study subjects were patients who underwent renal biopsy at the Chinese People’s Liberation Army (PLA) General Hospital. The inclusion criteria were as follows: patients aged between 20 and 70 years; patients diagnosed with type 2 diabetes mellitus (type 2) and with persistent overt proteinuria (defined as urinary albumin excretion ≥ 300 mg/24 h or urinary protein excretion ≥ 500 mg/24 h by at least two tests without evidence of urinary tract infection); patients with serum creatinine less than 1.41 μmol/L; and patients who were voluntarily admitted to the hospital to undergo renal biopsy. The exclusion criteria were the following: patients who had a confirmed diagnosis of renal diseases before they were diagnosed with type 2 diabetes; with a clinically confirmed diagnosis of NDRD, including lupus nephritis and Henoch-Schönlein purpura nephritis; with familial hereditary nephropathies, such as autosomal dominant polycystic kidney disease; and with an uncertain pathological diagnosis.

The tissue was examined by at least two pathology experts together with another two nephrologists; then the pathological diagnosis was determined. Renal biopsy indications: in clinical practice, the possible diagnosis of NDRD was confirmed by renal biopsy when the following situations occurred: gross/microscopic hematuria; elevated serum creatinine without obvious proteinuria; persistent massive proteinuria with normal renal function and no diabetic retinopathy. The renal biopsy indications for the suspected diagnosis of NDRD at our centre were in accordance with those listed for NDRD in the 2007 Kidney Disease Outcomes Quality Initiative (KDOQI) guidelines. Thus, 19 cases were diagnosed as DN and 18 cases were diagnosed as NDRD which included 10 (membranous nephropathy) MN and 8 IgAN. To evaluate whether glycopattern in urine was a potential biomarker to differentiate DN from NDRD, the presence of other kidney diseases accompanied with DN or NDRD and the patients who had pathological diagnosis of DN combined with NDRD were excluded in this study.

In order to evaluate potential biomarkers for distinguishing DN from NDRD, all the patients were divided into DN and NDRD group. To assess the possibility of urinary glycopatterns as potential biomarkers for monitor the process of DN, we separate the DN group into two subgroups according to the eGFR level. The eGFR level was calculated using the Modification of Diet in Renal Disease (MDRD) Study formula. The DN group I defined as the eGFR > 60 mL/min/1.73 m², along with low level 24 h urinary protein (1.41 ± 0.61 g/24 h), means the early stage of kidney disease. The DN group II defined as eGFR < 60 mL/min/1.73 m², along with high level 24 h urinary protein (2.64 ± 1.41 g/24 h), means the patients have renal insufficiency; indicating a worse prognosis.

All healthy volunteers (HVs) had no recent illness or medication. All females had no menstrual cycle at the time of collection. The collection of urine samples was done prior to treatment and before the biopsy or at least 1 month after the biopsy. The clinical information was summarized in Table 1.

2.2. Study Approval. The collection and use of all human pathology specimens for research presented here were approved by the ethical Committee of Northwest University (Xi’an, China), the Department of Nephrology, Chinese PLA General Hospital, Chinese PLA Institute of Nephrology, State Key Laboratory of Kidney Diseases, National Clinical Research Center of Kidney Diseases, Beijing Key Laboratory of Kidney Disease (Beijing, China). Written informed consent was received from participants for the collection of their urine. This study was conducted in accordance with the ethical guidelines of the Declaration of Helsinki.

2.3. Processing of Urine. 15 mL of urine was collected and stored at 4°C immediately. Then, the samples were centrifuged at 3000 rpm for 20 minutes at 4°C to remove cells and debris. The supernatant was frozen at –80°C in 1 mL aliquots. The method of trichloroacetic acid (TCA)/acetone precipitation was utilized to prepare urinary protein; the method of urinary protein precipitation was described elsewhere with
little modification [19–21]. Briefly, 3 mL mixed urine specimen was mixed with 30 mL TCA 10%-acetone solution 90% (v/v), followed by overnight incubation at −20°C. The mixture was centrifuged at 10,000g for 15 min at 4°C; after that, protein precipitate was resuspended by 2 mL ice-cold acetone; after centrifugation, the protein was left to dry on ice for 1 h. Pellet resuspension was performed with 500 µL 10 mM PBS buffer (pH 7.4) with 1% Tween-20 at ambient temperature for 15 min, sonicated for another 15 min in ultrasound bath. The protein concentration was determined by Bradford assay (Sigma-Aldrich, St. Louis, MO).

2.4. Lectin Microarrays. To reduce the differences between subjects and to tolerate individual variation, urinary protein was pooled equivalently according to sample classification, and each class was divided into three biological replications. The urinary proteins (80 µg) were mixed with equal volume of Na₂CO₃ solution (100 mM, pH 9.3) and then labeled with Cy3 fluorescent dye (GE Healthcare, Biosciences, Piscataway, NJ, USA); the mixture was incubated for 3 h at ambient temperature in darkness. After that, the labeled protein was purified with Sephadex G-25 columns (GE Healthcare) according to the manufacturer’s instructions to remove uncombined fluorescent dye. The lectin microarray was fabricated by using 37 exogenous plant lectins as purchased from Vector Laboratories (Burlingame, CA), from Sigma-Aldrich (St. Louis, MO), and from Calbiochem Merck (Darmstadt, Germany) with different binding preferences covering N- and O-linked glycans. The lectins were dissolved to a concentration of 1 mg/mL in the manufacturer’s recommended buffer, which contained 1 mMol/L appropriate monosaccharide. Lectins were spotted on the homemade epoxysilane-coated slides with Stealth micro spotting pins (SMP-10B) (TeleChem, USA) by a Capital smart microarrayer (CapitalBio, Beijing). Please refer to Qin et al. [22] for the sugar-binding specificities of the lectins, and layout of the lectin microarray was showed in Figure 1(a). Each lectin was spotted in triplicate per slide. After that, the slides were incubated in a humidity-controlled incubator at 50% humidity overnight and then put in vacuum dryer for 2 h at 37°C to immobilize. Then, the slides were blocked with the blocking buffer containing 1% (w/v) BSA in 10 mM PBS (pH 7.4) for 1 h at 25°C and then rinsed twice with 1 × PBST (0.2% Tween-20 in 10 mM PBS, pH 7.4), followed by a final rinse in 10 mM PBS. Previously, we had determined that 4 µg labeled protein was used to ensure that all binding signals were produced in the linear binding range of each lectin for the lectin microarrays [22]. Therefore, 4 µg of Cy3-labeled protein mixeed with 0.5 mL of incubation buffer containing 0.2% (w/v) BSA, 500 mM glycine, and 0.1% Tween-20 in 10 mM PBS was applied to incubate with the lectin microarrays, and an incubation was performed in the chamber at 25°C for 3 h in a rotisserie oven set at 4 rpm. After incubation, the slide was washed with 1 × PBST twice for each for 10 min and washed twice with 10 mM PBS for 5 min and then dried by centrifugation at 600 rpm for 5 min at room temperature. The microarrays were scanned with the 70% photomultiplier tube and 100% laser power settings using a GenePix 4000B Microarray Scanner (Axon Instruments, USA).

2.5. Data Acquisition and Analysis. The fluorescence signal intensity of each spot was extracted by GenePix 3.0 software (Axon Instruments, Inc., USA). To avoid the nonspecific adsorption, if the signal intensity of one lectin spot was less than average background ± standard deviation (SD), it is regarded as invalid spot and removed. After data filtered, the median of the effective data points of each sample was counted. To eliminate fluorescence bias between different data sets, the fluorescence signals of each lectin microarray were treated by max-normalization [23–25]. Briefly, the signals of each lectin were normalized to the highest signal intensity among 37 lectins. After normalization, the processed data of the parallel data sets were compared with each other based upon fold changes according to the following criteria: fold changes ≥ 1.5 or ≤0.67 and p < 0.05 in the pairs indicated up- or downregulation of certain kind of glycopatterns, respectively. Differences between the arbitrary two data sets or multiple data sets were tested using Student’s t-test or Kruskal-Wallis test; the histogram and ROC
2.6. Urinary Protein Microarrays. In order to verify the results of the lectin microarrays, a urinary protein microarray was produced by using another 48 individual urine samples; the detailed sample information was showed in Table S1 (see Supplementary Material available online at https://doi.org/10.1155/2017/5728087). The fabrication and incubation of urinary protein microarrays were similar with lectin microarrays. Briefly, urinary proteins were dissolved in spotting buffer containing 0.5 mg/mL BSA in 10 mM PBS (pH 7.4) to a concentration of 1 mg/mL and spotted on the homemade epoxysilane-coated slides with Stealth micro spotting pins (SMP-10B) by a Capital smart microarrayer. A urinary protein microarray was produced by spotting 48 individual urine samples (7 HVs, 9 T2DM patients, 15 patients with DN (7 DN group I, 8 DN group II), and 17 patients with NDRD (9 MN and 8 IgAN)) on the surface of the epoxy slide. Each urine specimen was spotted in triplicate per slide and incubated in a humidity-controlled incubator at 50% humidity overnight for immobilization. Then, the slides were blocked with the blocking buffer for 1 h; after washing, the blocked slide incubated with 4 μg Cy3 labeled lectin diluted in 0.5 mL of incubation buffer for 3 h at room temperature in the dark. The slide was washed with 1 × PBST twice for each for 10 min and washed once with 10 mM PBS for 5 min and then dried by centrifugation at 600 rpm for 5 min. The slides were scanned with the 70% photomultiplier tube and 100% laser power settings using a GenePix 4000B Microarray Scanner and the signal intensity was extracted by GenePix 3.0 software.

2.7. SDS-PAGE and Lectin Blotting. Urinary glycoproteins were further analyzed using lectin blotting. Firstly, the pooled urinary proteins were analyzed by SDS-PAGE. Briefly, urinary proteins were mixed with 5× loading buffer and boiled for 5 min at 100°C; after that, the samples were spun down and loaded on a 10% polyacrylamide resolving gel and a 3% stacking gel, and protein molecular weight marker (Thermo Scientific, Waltham, USA) was run in gel simultaneously. After being immobilized, the gel was stained by alkaline silver. For lectin blotting analysis, the proteins in gels were transferred to a PVDF membrane (Millipore, MA, USA) at 100 V for 90 min in wet transfer device (Beijing Liuyi Instrument Factory, China). Subsequently, the membranes were washed twice with TBST buffer (150 mM NaCl, 10 mM Tris-HCl, and 0.05% v/v Tween-20, pH 7.5) and blocked for 1 h with Carbo-Free Blocking Solution (Vector, Burlingame, CA) for 1 h at ambient temperature. The membranes were then incubated with 3 μg/ml Cy5 (GE Healthcare, Biosciences, Piscataway, NJ, USA) labeled lectins in previous blocking solution with gentle shaking overnight at 4°C below protection from light. The membranes were washed twice each for 10 min with TBST and scanned by Storm 840 PhosphorImager (Molecular Dynamics, Sunnyvale, CA) in red fluorescence channel 635 nm excitation/650 LP emission.

3. Result

3.1. Alterations of Urinary Glycopatterns with the Development of DN. About 30–40% of patients with type 1 or type 2 diabetes develop evidence of nephropathy, responsible for
40~50% of all ESRD. The glycopatterns in urine from HVs and patients were evaluated by the lectin microarrays. The glycopatterns of Cy3-labeled pooled urinary proteins from HVs and all patients bound to the lectin microarrays were shown in Figure 1(b). The normalized fluorescent intensities (NFIs) for each lectin were summarized as the mean values ± SD in Table S2.

Firstly, we investigated the glycopattern of urinary glycoproteins in HVs, patients with T2DM, and different degree of DN separately. The NFIs of three biological replicates were imported EXPANDER 6.4 for hierarchical clustering analysis (HCA). The NFIs values mapped to the range of 2 to –2 interval were color coded, with red and green indicating an increase and decrease in the abundance of glycopatterns, respectively. The HCA revealed that both DN groups were clustered into one class, whereas the HVs and T2DM cluste-tered into another class, and lectins were categorized into three independent groups according to the NFIs of lectins showing the positive signals in HVs and patients (Figure 2(a)). The results indicated that the abundance of glycan patterns was different between HVs, T2DM, and DN groups. Further, the NFIs of each lectin from HVs and patients were compared based on fold change in pairs, with all p values below 0.05, to evaluate whether the glycopatterns of urinary glycoproteins were altered with the development of DN (Table 2). Compared the alterations of glycopatterns in urine between HVs and T2DM, the results showed that the α-Fuc binder AAL, LacNac, poly-LacNac and (GlcNAc)_n binder LEL, Galβ1-3GalNAcα-Ser/Thr (T-antigen) binder ACA, and multivalent Sia and (GlcNAc)_n binder WGA exhibited significantly decreased NFIs in T2DM (all fold change = 0, p < 0.001). In addition, the α-Gal/GalNac binder BS-I, (β-1,4)-linked GlcNAc binder DSA, and α- or β-linked terminal GalNac binder SBA showed significantly decreased NFIs in T2DM (all fold change ≤ 0.60, p < 0.05) (Figure 2(b)).

Evaluating the alterations of urinary glycopatterns with the development of DN, the results showed that there were 6 lectins (e.g., the Siaα2-3Galβ1-4Glc(NAC)/Glc binder MAL-II, GalNac and GalNacα-1,3Gal binder PTL-I, and terminal in GalNac and Gal binder SJA) that exhibited significantly increased NFIs in DN group I compared with HVs and T2DM, and high-Mannose, Manα1-6(Manα1-3)Man binder Con A showed significantly increased NFIs in DN group II compared with HVs and T2DM (p < 0.01). However, the α- or β-linked terminal GalNac binder SBA and Fucα1-6GlcNAc binder PSA revealed significantly decreased NFIs in DN group II compared with HVs and T2DM (all fold change = 0, p < 0.001). The (GlcNAc)_2–4 binder DSA and Siaα2-6Gal/GalNac binder SNA exhibited significantly increased NFIs (all fold change ≥ 2.88, p < 0.05) in both DN groups compared with HVs and T2DM, and the Manα1-3Man binder GNA, α-Gal/GalNac binder BS-I, α-Fuc binder AAL, and α- or β-linked terminal GalNac binder SBA exhibited increased NFIs (all fold change ≥ 1.61, p < 0.05) in each DN group I compared with T2DM. However, the (GlcNAc)_2–4 binder STL exhibited significantly decreased NFIs in both DN groups compared with HVs and T2DM (all fold change = 0, p < 0.001). And the α-Gal/GalNac binder BS-I and (GlcNAc)_n and branched (LacNac)_n binder PWM showed decreased NFIs (fold change = 0.54, p < 0.05) in DN group II compared with HVs (shown in Figure 2(b)).

### 3.2. Alterations of Urinary Glycopatterns Associated with Different Degree of DN.

Although the albuminuria and eGFR could reflect the process of DN, it also needs to find new sensitive biomarkers for monitoring the development of DN. The glycopatterns of urinary glycoproteins in different degree of DN were analyzed. The NFIs of each lectin from DN group I and group II were compared based on fold change (Table 2), and the results demonstrated that there were 9 lectins (e.g., the Siaα2-3Galβ1-4Glc(NAC)/Glc binder MAL-II, GalNac and GalNacα-1,3Gal binder PTL-I, and terminal in GalNac and Gal binder SJA) that exhibited significantly decreased NFIs in DN group II compared with group I (all fold change = 0, p < 0.001). However, the high-Mannose, Manα1-6(Manα1-3)Man binder Con A showed significantly increased NFIs in DN group II compared with DN group I (p < 0.001) (shown in Figure 1(b)). Besides that, the Siaα2-6Gal/GalNac binder SNA showed significantly increased NFIs in DN group II compared with group I (fold change = 2.54, p < 0.01). The α-Gal/GalNac binder BS-I, (GlcNac)_n and branched (LacNac)_n binder PWM, and Manα1-3 man binder GNA exhibited significantly decreased NFIs (all fold change ≤ 0.55, p < 0.05) in DN group II compared with DN group I (Figure 2(b)).

### 3.3. Alteration of Glycopatterns between DN and NDRD.

Currently, it is difficult to distinguish DN from NDRD in clinical practice. Renal biopsy is still a reliable clinical detection for diagnosis of DN. Here, the differences of urinary protein glycopatterns were investigated between DN (DN group I and DN group II) and NDRD (MN and IgAN). As a result of HCA, DN groups and NDRD were separated and lectins were categorized into four independent groups according to the NFIs of lectins, which revealed that the glycopatterns in urine showed difference between DN and NDRD (Figure 3(a)).

The fold change between DN and NDRD groups was summarized in Table 3.

The results showed that there were 5 lectins (the Siaα2-3Galβ1-4Glc(NAC)/Glc binder MAL-II, GalNac and GalNacα-1,3Gal binder PTL-I, terminal in GalNac and Gal binder SJA, GalNacα1-3((Fucα1-2))Gal binder DBA, and Gal and T-antigen binder PTL-II) to show high NFIs in DN group I and high-Mannose, Manα1-6(Manα1-3)Man binder Con A to show high NFIs in DN group II compared with NDRD (p < 0.001). However, the (GlcNac)_2–4 binder STL showed high NFIs in NDRD compared with DN groups, and α-Fuc binder AAL, α- or β-linked terminal GalNac binder SBA and Fucα1-6GlcNAc binder PSA showed high NFIs in NDRD compared with DN groups (p < 0.001). In addition, the Galβ1-3GalNacα-Ser/Thr(T) binder Jacalin and Fucα1-2Galβ1-4Glc(NAC) binder UEA-I exhibited high NFIs in MN compared with both DN groups (p < 0.001). Besides that, the β-D-GlcNAc binder DSA showed significantly higher NFIs in both DN groups than that in patients with IgAN (fold change = 3.28, p < 0.001), the Siaα2-6Gal/GalNac binder SNA showed significantly increased NFIs in DN group II compared with NDRD (fold change ≥ 2.56, p < 0.001), and...
Table 2: Fold change of the urinary glycopatterns based upon ratio of the NFI s of each lectin between HVs, T2DM, and DN groups.

<table>
<thead>
<tr>
<th>Lectin name</th>
<th>Specificity</th>
<th>T2DM/HVs</th>
<th>DN group I/HVs</th>
<th>Fold change&lt;sup&gt;a&lt;/sup&gt;</th>
<th>DN group II/T2DM</th>
<th>DN group II/DN group I</th>
<th>DN group II/DN group I</th>
</tr>
</thead>
<tbody>
<tr>
<td>MAL-II</td>
<td>Siaα2-3Galβ1-4Glc(NAc)/Glc</td>
<td>—</td>
<td>***</td>
<td>***</td>
<td>—</td>
<td>—</td>
<td>0***</td>
</tr>
<tr>
<td>PTL-I</td>
<td>GalNAc, GalNAcα1-3Gal, GalNAcα1-3Galβ-1,3/4Glc</td>
<td>—</td>
<td>***</td>
<td>***</td>
<td>—</td>
<td>—</td>
<td>0***</td>
</tr>
<tr>
<td>SJA</td>
<td>Terminal in GalNAc and Gal, anti-A and anti-B human blood group</td>
<td>—</td>
<td>***</td>
<td>***</td>
<td>—</td>
<td>—</td>
<td>0***</td>
</tr>
<tr>
<td>AAL</td>
<td>α-Fuc, Fucα1-6GlcNAc (core fucose)</td>
<td>0*** 3.30**</td>
<td>***</td>
<td>0***</td>
<td>—</td>
<td>—</td>
<td>0***</td>
</tr>
<tr>
<td>LEL</td>
<td>LacNAc and poly LacNAc, (GlcNAc)&lt;sub&gt;n&lt;/sub&gt;</td>
<td>0*** 0***</td>
<td>—</td>
<td>0***</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>DBA</td>
<td>αGalNAc, Tn antigen, GalNAcα1-3(Fucα1-2)Gal (blood group A antigen)</td>
<td>—</td>
<td>***</td>
<td>***</td>
<td>—</td>
<td>—</td>
<td>0***</td>
</tr>
<tr>
<td>RCA120</td>
<td>β-Gal, Galβ1-4GlcNAc (type II), Galβ1-3GlcNAc (type I)</td>
<td>0.799 1.05 1.31 1.17 1.46 1.12</td>
<td>—</td>
<td>0***</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>STL</td>
<td>(GlcNAc)&lt;sub&gt;n&lt;/sub&gt;, core (GlcNAc) of N-glycan</td>
<td>1</td>
<td>0***</td>
<td>0***</td>
<td>0***</td>
<td>0***</td>
<td>—</td>
</tr>
<tr>
<td>BS-I</td>
<td>α-Gal/GalNAc, Galα1-3Gal</td>
<td>0.60* 0.97 1.61* 0.54* 0.89 0.55*</td>
<td>—</td>
<td>0***</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>ConA</td>
<td>High-Mannose, Manα1-6(Manα1-3)Man, terminal GlcNAc</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>/***</td>
<td>/***</td>
<td>/***</td>
</tr>
<tr>
<td>PTL-II</td>
<td>Gal, blood group H, T-antigen</td>
<td>—</td>
<td>***</td>
<td>***</td>
<td>—</td>
<td>—</td>
<td>0***</td>
</tr>
<tr>
<td>DSA</td>
<td>(β-1,4)-linked GlcNAc</td>
<td>0.21** 2.88*** 13.61*** 2.88*** 13.61*** 1</td>
<td>—</td>
<td>0***</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>SBA</td>
<td>α- or β-linked terminal GalNAc, (GalNAc)&lt;sub&gt;n&lt;/sub&gt;, GalNAcα1-3Gal, blood-group A</td>
<td>0.32* 1.16 3.63* 0*** 0*** 0***</td>
<td>—</td>
<td>0***</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>PSA</td>
<td>Fucα1-6GlcNAc, α-D-Man, α-D-Glc</td>
<td>1.31 1.52 1.16</td>
<td>0***</td>
<td>0***</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>ACA</td>
<td>Galβ1-3GalNAcα1-Ser/Thr (T antigen), sialyl-T(ST)</td>
<td>0*** 0***</td>
<td>—</td>
<td>0***</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>WGA</td>
<td>Multivalent Sia and (GlcNAc)&lt;sub&gt;n&lt;/sub&gt;</td>
<td>0*** 0***</td>
<td>—</td>
<td>0***</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>PWM</td>
<td>(GlcNAc)&lt;sub&gt;n&lt;/sub&gt;, branched (LacNAc)&lt;sub&gt;n&lt;/sub&gt;</td>
<td>0.59 1.04 1.76</td>
<td>0.54*</td>
<td>0.92 0.521*</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>GNA</td>
<td>Manα1-3Man</td>
<td>1.46 2.39** 1.64* 1.17 0.80 0.488*</td>
<td>—</td>
<td>0***</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>PHA-E + L</td>
<td>Bisecting GlcNAc, biantennary N-glycans, tri- and tetraantennary complex-type N-glycan</td>
<td>—</td>
<td>/***</td>
<td>/***</td>
<td>—</td>
<td>—</td>
<td>0**</td>
</tr>
<tr>
<td>SNA</td>
<td>Sia2-6Gal/GalNAc</td>
<td>1.37 4.34** 3.16* 11.03*** 8.05*** 2.54***</td>
<td>—</td>
<td>/***</td>
<td>/***</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

<sup>a</sup>Fold changes of lectins bound to DN compared to healthy control and diabetic calculated with mean value of normalized florescence intensities of lectins obtained from 3 biological reduplicates. —, no significant difference; /, the denominator of the fold-change was infinite; 0, the numerator of the fold-change was zero. * versus p < 0.05; ** versus p < 0.01; *** versus p < 0.001; p value was calculated by Student’s t-test.
Figure 2: The alterations of urinary glycopatterns associated with development of DN. (a) Hierarchical clustering analysis of the 20 lectins with three biological replicates. Samples were listed in columns and the lectins were listed in rows. The color and intensity of each square indicated expression levels relative to other data in the row. Red: high, green: low, and black: medium. (b) Comparing the NFI s of the urinary glycopatterns based upon ratio of each lectin between HVs, T2DM, and DN groups. Significant differences between groups were analyzed according to Student’s t-test, respectively (* p < 0.05, ** p < 0.01, and *** p ≤ 0.001). The data were the averaged NFI s ± SD of three biological replicates.
**Table 3: Fold change of the urinary glycopatterns based upon ratio of the NFIs of each lectin between DN groups and NDRD.**

<table>
<thead>
<tr>
<th>Lectin name</th>
<th>Specificity</th>
<th>DN group I/MN</th>
<th>DN group I/IgAN</th>
<th>DN group II/MN</th>
<th>DN group II/IgAN</th>
</tr>
</thead>
<tbody>
<tr>
<td>Jacalin</td>
<td>Galβ1-3GalNAcα-Ser/Thr(T), GalNAcα-Ser/Thr(Tn)</td>
<td>0***</td>
<td>—</td>
<td>0***</td>
<td>—</td>
</tr>
<tr>
<td>MAL-II</td>
<td>Sia2-3Galβ1-4Glc(Nac)/Glc,</td>
<td>/***</td>
<td>/***</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>PTL-I</td>
<td>GalNAc, GalNAcα-1,3Gal, GalNAcα-1,3Galβ1-1,3/4Glc</td>
<td>/***</td>
<td>/***</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>SJA</td>
<td>Terminal in GalNAc and Gal, anti-A and anti-B human blood group</td>
<td>/***</td>
<td>/***</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>AAL</td>
<td>α-Fuc, Fuc(1-6) GlcNac (core fucose)</td>
<td>/**</td>
<td>0.65</td>
<td>—</td>
<td>0*</td>
</tr>
<tr>
<td>DBA</td>
<td>αGalNAc, Tn antigen, GalNAcα-1-3(Fucα1-2)Gal (blood group A antigen)</td>
<td>/***</td>
<td>/***</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>RCA120</td>
<td>β-Gal, Galβ1-4GlcNac (type II), Galβ1-3GlcNac (type I)</td>
<td>0.22***</td>
<td>1.74*</td>
<td>0.24***</td>
<td>1.95***</td>
</tr>
<tr>
<td>STL</td>
<td>(GlcNAc)2-4 core (GlcNAc) of N-glycan</td>
<td>0***</td>
<td>0***</td>
<td>0***</td>
<td>0***</td>
</tr>
<tr>
<td>BS-I</td>
<td>α-Gal/GalNAc, Galα-1,3Gal</td>
<td>0.41**</td>
<td>/**</td>
<td>0.22***</td>
<td>/***</td>
</tr>
<tr>
<td>ConA</td>
<td>High-Mannose, Manα1-6(Manα1-3)Man, terminal GlcNac</td>
<td>—</td>
<td>—</td>
<td>/***</td>
<td>/***</td>
</tr>
<tr>
<td>PTL-II</td>
<td>Gal, blood group H, T-antigen</td>
<td>/***</td>
<td>/***</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>DSA</td>
<td>(α-1,4)-linked GlcNac</td>
<td>—</td>
<td>3.28***</td>
<td>—</td>
<td>3.28***</td>
</tr>
<tr>
<td>SBA</td>
<td>α- or β-linked terminal GalNAc, (GalNAc)α, GalNAcα-1-3Gal, blood-group A</td>
<td>/**</td>
<td>4.64*</td>
<td>—</td>
<td>0***</td>
</tr>
<tr>
<td>PSA</td>
<td>Fucα-1,6GlcNac, α-D-Man, α-D-Glc</td>
<td>0.46*</td>
<td>/**</td>
<td>/**</td>
<td>—</td>
</tr>
<tr>
<td>UEA-I</td>
<td>Fucα1-2Galβ1-4Glc(Nac)</td>
<td>0***</td>
<td>—</td>
<td>0***</td>
<td>—</td>
</tr>
<tr>
<td>PWM</td>
<td>(GlcNAc)α2-branched (LacNAc)α</td>
<td>0.50**</td>
<td>1.20</td>
<td>0.26***</td>
<td>0.62***</td>
</tr>
<tr>
<td>GNA</td>
<td>Manα1-3Man</td>
<td>1.49</td>
<td>3.80**</td>
<td>0.73</td>
<td>1.85**</td>
</tr>
<tr>
<td>PHA-E+L</td>
<td>Bisecting GlcNAc, bi-antennary N-glycans, tri- and tetra-antennary complex-type N-glycan</td>
<td>/**</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>SNA</td>
<td>Sia2-6Gal/GalNAc</td>
<td>1.00</td>
<td>1.80</td>
<td>2.56***</td>
<td>4.57***</td>
</tr>
</tbody>
</table>

*Fold changes of lectins bound to DN groups compared to NDRD calculated with mean value of normalized fluorescence intensities of lectins obtained from 3 biological reduplicates. —, no significant difference; /, the fold-change was infinite; 0, the numerator of the fold-change was zero. * versus p < 0.05; ** versus p < 0.01; *** versus p < 0.001, p value was calculate by Student’s t-test.
Figure 3: The alterations of glycopatterns between DN and NDRD. (a) Hierarchical clustering analysis of the 19 lectins with three biological replicates. DN groups and NDRD were listed in columns, and the lectins were listed in rows. (b) Ten lectins revealed significant differences between DN and NDRD. Significant differences between groups were analyzed according to Student's \( t \)-test, respectively. \( ^* p < 0.05 \), \( ^{**} p < 0.01 \), and \( ^{***} p \leq 0.001 \). The data were the averaged NFI ± SD of three biological replicates.
the Manα1-3Man binder GNA associated with increased NFIs in both DN groups compared with IgAN (fold change ≥ 1.85, \( p < 0.01 \)). And the β-Gal/GalNAc binder RCA120 and α-Gal/GalNAc binder BS-I exhibited increased NFIs in both DN groups compared with patients with IgAN (fold change ≥ 1.74, \( p < 0.05 \)) but showed decreased NFIs compared with MN (fold change ≤ 0.41, \( p < 0.05 \)). However, the (GlcNAc), and branched (LacNAc), binder PWM exhibited significantly decreased NFIs (all fold change ≤ 0.50, \( p < 0.01 \)) in both DN groups compared with MN (Figure 3(b)).

### 3.4. Assessment of Urinary Glycopatterns as Potential Biomarkers for Diagnosis of DN. The most reliable way of biomarker discovery is to test the candidates with a large cohort of samples that have clear and sufficient clinical records. Here, a urinary protein microarray is produced by spotting 48 individual urine samples (7 HVs, 9 T2DM patients, 15 patients with DN (7 DN group I, 8 DN group II), and 17 patients with NDRD (9 MN and 8 IgAN)) on the surface of the epoxy slide. Each urine sample was spotted in triplicate, and the layout of the urinary protein microarrays was shown in Figure 4(a). SNA and STL that revealed significant differences (\( p < 0.05 \)) in urinary glycopatterns according to the results of the lectin microarrays were selected to validate the alteration of the targeted glycopatterns in individual sample and evaluate the reliability of urinary glycopatterns as potential biomarkers for diagnosis of DN.

As a result, SNA staining showed increased signals in both DN groups compared with HVs and patients with T2DM and significant difference between DN group I and group II (\( p < 0.05 \)). Interestingly, the signals of SNA staining showed an increasing trend with development of DN (Figure 4(b)). The result of correlation analysis indicated that the abundance of Siaα2-6Gal/GalNAc was positive correlation with the progression of DN (rs = 0.888). On the contrary, the signals of STL staining were significantly lower in both DN groups than that in NDRD (\( p < 0.001 \)). The result of ROC analysis showed that the ROC integral was 0.969 and \( p < 0.001 \), which indicated the glycopattern of (GlcNAc)2-4 recognized by STL in urine glycoprotein could be a potential diagnostic indicator to differentiate DN and NDRD (Figure 4(c)). These results were consistent with lectin microarrays.

### 3.5. Lectin Blotting Analysis. In order to investigate precisely alterations of glycopatterns associated with DN, SDS-PAGE and lectin blotting analysis were performed with SNA and STL staining (Figure 4(d)). SNA staining showed obvious binding patterns with six distinct protein bands of approximately 120, 90, 70, 50, 40, and 25kDa marked as b1–b6, respectively. The band of b4 showed strong staining signals in all samples except HVs. There were four bands (b2, b3, b5, and b6) to show increased signals in both DN groups compared with T2DM. Moreover, the glycoprotein bands (b2, b3, and b5) at about 90 kDa, 70 kDa, and 40 kDa exhibited increased staining signals in DN group II compared with group I (Figure 4(e)). Therefore, the abundance of Siaα2-6Gal exhibited general increase in urine with development of DN. Comparing DN and NDRD, the band of b4 (50 kDa) showed strong staining signals in both DN and NDRD groups but did not exhibit significant difference, whereas the bands of b3 (70 kDa) and b5 (40 kDa) revealed stronger staining intensities in DN groups than those in MN and IgAN.

STL staining showed obvious binding patterns with one distinct urinary protein band (b4) in both DN and NDRD at about 50 kDa. The binding signals of this glycoprotein band were significant higher in MN and IgAN than that in DN groups but have no difference between DN groups.

### 4. Discussion

Glycosylation is an important co- and posttranslational modification of glycoprotein, involved in many biological processes such as embryogenesis and neoplasm metastasis. The altered oligosaccharide moiety of glycoprotein was considered to be associated with the occurrence and development of diseases [26–29]. Abnormal glycosylation of certain glycoprotein in urine could be a more sensitive biomarker for disease diagnosis and monitoring. It is reported that aberrantly excreted urinary O-glycosylated proteins (clusterin, leucine-rich alpha-2-glycoprotein, and kininogen) could serve as potential biomarkers for the early detection of early stage ovarian cancer [30]. As a powerful tool, lectin microarray was used to investigate the glycopatterns of protein for decades. With the development of mass spectrometry, the glycan profile could be analysis by MS. However, as a rapid analysis approach, lectin microarray owned some unique advantages. First of all, the proteins do not need to enzymolysis prior to analysis, and the target is simple to prepare. Secondly, the linkage information of glycopatterns could be obtained by lectin microarray; however, mass spectrometry has difficulty in discriminating between the many structural isomers presented by glycans [31]. There is no denying that MS has advantage to acquire precisely information about the component of glycan. Notwithstanding intense efforts by multiple research groups, the use of mass spectrometry as a stand-alone technique for complete characterization of glycans is still far from complete [32].

In this study, glycopatterns of urine glycoprotein in patients with DN were compared with controls (healthy volunteers and patients with T2DM) and patients with NDRD (MN and IgAN), respectively. As a result, STL showed strong binding signals in HVs and patients with T2DM and IgAN; in addition, DSA exhibited strong binding signals in patients with DN and MN. It indicated that GlcNAc and polymers of GlcNAc were predominant glycopatterns in urine of patients. In urinary glycoprotein, HexHexNAc1–N-Asn was typically N-glycan observed which is similar to our findings [33]. As a result of lectin microarrays, there were 9 and 15 lectins to show binding signals in patients with T2DM and DN group I, which indicated more glycopatterns emerged in urine of patients in middle and advanced stage of DN without renal insufficiency (DN group II) compared with patients with T2DM. However, when compared to patients with overt proteinuria and renal insufficiency (DN group II), the situation was changed, only 7 lectins (e.g., WFA, GSL-II, and PNA) exhibited effective binding signals in DN group II, and 9 lectins (e.g., WFA, GSL-II, and PNA) failed to detect binding
Figure 4: Validation of the differential expressions of the glycopatterns in the urine associated with DN. (a) Scan images of Cy3-labeled SNA and STL incubating to urinary protein microarray, respectively. The incubation images were extracted by GenePix 3.0. (b) Scatter plot analysis of the original data obtained from the urinary protein microarrays. The statistical significance of the differences between groups was calculated by Kruskal-Wallis test. (c) ROC curve analysis of STL to differentiate DN and NDRD, AUC = 0.968, and \( p < 0.0001 \). (d) SDS-PAGE and lectin blot of pooled urinary protein using SNA and STL, M, marker; lane 1, HVs; lane 2, patients with T2DM; lane 3, DN group I; lane 4, DN group II; lane 5, patients with MN; lane 6, patients with IgAN. (e) The lectin binding strength of selected protein bands is displayed, and the binding strength was counted by ImageJ.
signals compared with DN group I. Furthermore, there were 3 lectins (BS-I, PWM, and GNA) to exhibit decreased NFIs in DN group II compared with DN group I. These findings demonstrated that the glycopatterns, especially the glycopatterns modified by Gal/GalNAc, were decreased as occurred in renal insufficiency. Ionou et al. reported the global reduction of the binding to lectins in urine samples of DN at macroalbuminuria stage [30]. Our finding further clarified that glycopatterns showed global reduction when occurring in renal insufficiency in stage of overt proteinuria. In rat diabetic nephropathy model which was induced by streptozotocin injection, the staining intensities of Gal/GalNAc binders PNA and RCA showed remarkable decrease in model after being induced for 5 weeks [34]. It is known that around 70% of the urinary proteins are estimated to be derived from the kidney and the urinary tract [35]. Therefore, we suspected that the general reduction in signature glycopatterns of Gal/GalNAc in kidney may contribute to the decrease of these glycopatterns in urinary glycoprotein with development of DN; the relationship between glycopatterns in kidney and urine will be clarified in next step.

In present study, we found that the glycopatterns of Siaα2-6Gal/GalNAc showed increased tendency with development of DN, and urinary protein microarray also confirmed this result. Our finding suggested Siaα2-6Gal/GalNAc is potential indicator for monitoring of DN. Furthermore, lectin blotting analysis was employed to investigate the precise alterations of Siaα2-6Gal/GalNAc. Generally, it showed higher staining signals in DN groups than that in HVs and T2DM. The glycoprotein bands at about 90 kDa, 70 kDa, and 40 kDa showed increased staining signals with development of DN. These results indicated the sialylation of these glycoproteins was increased and it may contribute to the upregulation of expression level of Siaα2-6Gal/GalNAc in urinary glycoprotein with development of DN. It is known that the level of serum sialic acid could indicate extensive vascular damage in T2DM and predict microvascular complications occurring in diabetics [36, 37]. As a result of our finding, the expression level of Siaα2-6Gal/GalNAc in urinary glycoprotein correlated with development of DN in stage of overt proteinuria. Similarly, it is reported that a 41 kDa of sialylated glycoprotein orosomucoid which contained Siaα2-6Gal/GalNAc increased excretion in DN, and it could be a potential biomarker for DN [38–40]. In addition, it is reported that the excretion of transferrin (76.5 kDa) and alpha-1-antitrypsin (46 kDa) showed marked increase in urine of patients with DN, which could reflect the development of microalbuminuria; moreover, urinary transferrin is a sensitive indicator of glomerular damage [41, 42]. Therefore, the glycopatterns of Siaα2-6Gal/GalNAc and glycoprotein modified by Siaα2-6Gal/GalNAc (especially the glycoproteins with molecular weight about 90 kDa, 70 kDa; and 40 kDa) in urine could be valuable potential diagnosis indicators for monitoring of DN, and these glycoproteins will be characterized by MS in future.

Although certain clinical manifestations such as shorter duration of diabetes, absence of retinopathy, presence of microscopic hematuria, and active urinary sediment are markers associated with NDRD in type 2 diabetes with clinical renal disease, and the histories of diabetes mellitus, systolic blood pressure, glycated HbA1c, hematuria, diabetic retinopathy, and hemoglobin are independently related to DN, renal biopsy remains indispensable to differentiate between DN and NDRD [43–48]. In this study, we examined the difference between DN and NDRD in urinary glycopatterns; our result showed that the relative abundance of (GlcNAc)2–4 in urinary glycoprotein could effectively distinguish between DN and NDRD, and ROC curve analysis also indicated that the glycopatterns recognized by STL have the potential to serve as a clinical predictor to distinguish between DN and NDRD. It demonstrated that the patients with IgAN exhibit circulating IgA1 with reduced Gal (galactose-deficient IgA1) [49–51]. It reported that deficiency of the β1,4GalT1 glycosyltransferase in genetic remodeling mouse could cause IgAN. Because of deficiency of the β1,4GalT1, β4-galactosylation of the N-glycans on the serum IgA was completely absent, which led to the exposure of GlcNAc of N-linked glycans [52]. In addition, urinary excretion of galactose-deficient IgA1 was elevated in patients with IgAN and the urinary Gd-IgA1 levels correlated with proteinuria [53]. Hiki et al. reported that O-glycans attached to serum IgA are truncated to expose GlcNAc, which correlates with antibody glomerular deposition in IgAN [54]. Therefore, we concluded that the exposure of GlcNAc on N-/O-glycans of IgA correlated with the increase of (GlcNAc)2–4 in urinary glycoprotein from patients with IgAN.

In conclusion, we investigated the correlation of alterations of protein glycopatterns in urine samples from HVs, patients with T2DM, and patients with DN and NDRD. Our finding revealed that the elevated expression level of Siaα2-6Gal/GalNAc in urine was correlated with development of DN in stage of macroalbuminuria, and (GlcNAc)2–4 in urine could be a clinical predictor to distinguish between DN and NDRD. It should be pointed out that the sample size in this study is small, and more samples are being recruited to verify our findings and glycoprotein which associated with abnormal glycopatterns will be isolated and identified in next step. This study provided a new method to monitor the development of DN and distinguish between DN and NDRD based on the urinary glycopatterns analysis.

Competing Interests
All the authors declared no competing interests.

Authors’ Contributions
Hanyu Zhu and Moyan Liu contributed equally to this work.

Acknowledgments
This study was supported by (1) National Key Research and Development Program (2016YFC1305502); (2) the National Natural Science Foundation of China (no. 61471399; no. 61671479); (3) Innovation Nursery Fund of PLA General Hospital (no. 15KMZ04); (4) National Clinical Research Center for Kidney Disease (no. 2013BAA109B05); (5) the State Key Development Program for Basic Research of China
(no. 2013CB530800); (6) Technology Project of Beijing (no. D13110700470000).

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