Differentiating Staphylococcus infection-associated glomerulonephritis and primary IgA nephropathy: a mass spectrometry-based exploratory study

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Staphylococcus infection-associated glomerulonephritis (SAGN) and primary IgA nephropathy (IgAN) are separate disease entities requiring different treatment approaches. However, overlapping histologic features may cause a diagnostic dilemma. An exploratory proteomic study to identify potential distinguishing biomarkers was performed on formalin fixed paraffin embedded kidney biopsy tissue, using mass spectrometry (HPLC–MS/MS) (n = 27) and immunohistochemistry (IHC) (n = 64), on four main diagnostic groups—SAGN, primary IgAN, acute tubular necrosis (ATN) and normal kidney (baseline transplant biopsies). Spectral counts modeled as a negative binomial distribution were used for statistical comparisons and in silico pathway analysis. Analysis of variance techniques were used to compare groups and the ROC curve to evaluate classification algorithms. The glomerular proteomes of SAGN and IgAN showed remarkable similarities, except for significantly higher levels of monocyte/macrophage proteins in SAGN—mainly lysozyme and S100A9. This finding was confirmed by IHC. In contrast, the tubulointerstitial proteomes were markedly different in IgAN and SAGN, with a lower abundance of metabolic pathway proteins and a higher abundance of extracellular matrix proteins in SAGN. The stress protein transglutaminase-2 (TGM2) was also significantly higher in SAGN. IHC of differentially-expressed glomerular and tubulointerstitial proteins can be used to help discriminate between SAGN and IgAN in ambiguous cases.

Staphylococcal infections have emerged as the most common cause of infection-associated glomerulonephritis in the elderly population and are also encountered with increasing frequency in young adults with a history of intravenous drug abuse1–7. Kidney biopsies from patients with Staphylococcal infection-associated glomerulonephritis (SAGN) frequently show glomerular IgA and co-dominant C3 staining with predominantly mesangial and a few capillary wall immune-type deposits. The pattern of staining is very similar to that seen in primary IgA nephropathy (IgAN), presenting a potential diagnostic pitfall. IgA vasculitis (Henoch–Schönlein purpura or HSP) can also show similar biopsy features, and 15–20% of patients with SAGN have a leukocytoclastic vasculitic rash, mimicking IgA vasculitis8,9. Making the correct diagnosis is important as different treatment approaches are warranted for SAGN and IgAN. Because of active infection, patients with SAGN should be treated with...
antibiotics, and immunosuppressive therapy should be avoided, at least in early stages of the disease. In contrast and depending on kidney function, level of proteinuria, and disease activity, primary IgAN is often treated with corticosteroids and/or immunosuppressive drugs. This is especially true for IgAN cases with crescents (Oxford score C1 or C2) and endocapillary hypercellularity (Oxford score E1)\textsuperscript{10–13}. Histologically, such cases of IgAN can resemble SAGN, and while clinical presentation of SAGN such as older age, rapidity of disease onset, recent Staphylococcal infection, presence of risk factors for infection can together help in differentiating the two diseases, it can still be challenging since not all these features are consistently present in every case\textsuperscript{7–9}.

We and others have previously demonstrated the diagnostic utility of mass spectrometry-based proteomic analysis on formalin-fixed paraffin-embedded (FFPE) tissue from kidney biopsies. Studies have also shown comparable peptide capture using both frozen and FFPE tissue\textsuperscript{14–19}. In this exploratory study, we used liquid chromatography and tandem mass spectrometry (LC–MS/MS)-based proteomics on FFPE biopsy samples to identify differentially-expressed proteins between SAGN and primary IgAN in an effort to shed light on the pathogenesis and to identify potential diagnostic biomarkers to help differentiate SAGN from primary IgAN on a kidney biopsy. Laser microdissection was used to collect glomeruli and tubulointerstitial tissue separately for analysis.

## Results

### Classification of biopsies used for the study.

Biopsies were identified from our Pathology database and from our previously published cohort of SAGN\textsuperscript{3,6}. A total of 27 biopsies were used for LC–MS/MS (Tables 1, 2). These included SAGN (n = 4), IgAN (n = 7), living donor baseline transplant biopsies (n = 8) as healthy control kidney tissue, and biopsies with a primary diagnosis of acute tubular necrosis (ATN, n = 8). The ATN group was included for comparison since SAGN biopsies typically do show acute tubular injury\textsuperscript{10–13}. For the SAGN group, we included biopsies from patients that had culture-proven Staphylococcal infection, and excluded biopsies that showed changes of underlying diabetic glomerulosclerosis. Also cases with more than moderate interstitial fibrosis and tubular atrophy were excluded to focus specifically on active SAGN as opposed to chronic renal injury. The IgAN group contained biopsies with MEST Score M1 E0 S0 T0 C0 (n = 3); MEST Score M1 E0 S1 T0 C0 (n = 8); MEST Score M0 E1 S1 T0 C1 (n = 2); and cases with MEST Score M1 E1 C1, (n = 2).

As this study was done on tissue left-over after routine diagnostic processing, most biopsy samples were depleted after LC MS/MS analysis. Hence, for immunohistochemical (IHC) staining, additional biopsies were retrieved from our Pathology database, including SAGN (n = 28), IgAN (n = 36; 17 biopsies with MEST score E0 and 19 biopsies with MEST score E1) and ATN (n = 6). Among the 28 SAGN biopsies, 22 showed endocapillary

| Disease category | Age (years) | Ethnicity | Sex | S. cr | Urine protein | Urine RBCs/hpf | Crescents | Endocapillary proliferative lesions | IF/TA (%) | IgG | IgA | C3 | EM deposits |
|------------------|------------|-----------|-----|-------|---------------|----------------|-----------|------------------------------------|-----------|-----|-----|----|-------------|
| SAGN             | 48         | C         | M   | 2.3   | 30 mg/dl      | 10–14          | Present   | Focal                              | 30        | Trace | 1 + m | 1 + | Paramesangial |
| SAGN             | 58         | AA        | M   | 1.5   | 140 mg/dl     | 10–14          | Absent    | Diffuse                            | 20        | 1 + m, g | 1 + m | 2 + | Mesangial, few humps |
| SAGN             | 44         | C         | F   | 2     | 300 mg/dl     | > 50           | Present   | Diffuse                            | < 10      | 1 + m, g | 2 + m, g | 1 + m, g | Mesangial, intra-membranous |
| SAGN             | 51         | C         | M   | 2.1   | Trace         | 3–4            | Absent    | Focal                              | < 10      | Trace | 2 + m, g | 2 + m | Mesangial |
| IgAN             | M1 E0 S1 T0 C0 | 24         | Asian | M   | 1.6 g/d      | 15–19          | Absent    | Absent                            | Absent    | Trace | 2 + m, trace | Mesangial |
| IgAN             | M1 E0 S1 T0 C0 | 29         | C     | M   | 0.8         | 0.5 g/d       | 5–9       | Absent                            | Absent    | 25    | Trace m | 2 + m | 2 + m | Mesangial |
| IgAN             | M1 E0 S1 T0 C0 | 19         | C     | M   | 1.1         | 0.4 g/d       | 3+        | Absent                            | Absent    | 10–15 | 1 + m | 3 + m, g | Mesangial, rare subendothelial |
| IgAN             | M1 E1 S0 T0 C1 | 70         | C     | F   | 1.2         | 4 g/d         | 3+        | Present                           | Focal     | < 10   | Absent | 1 + m | 2 + m | Mesangial |
| IgAN             | M1 E1 S1 T0 C1 | 58         | C     | M   | 1          | 300 mg/dl     | 15–19     | Present                           | Focal     | < 10   | Trace | 2 + m | Trace | Mesangial |
| IgAN             | M1 E0 S1 T1 C1 | 42         | Hispanic | C   | 1.7 g/d     | 10–14          | Present   | Absent                           | Absent    | 25    | Trace | 3 + m, g | Mesangial |
| IgAN             | M1 E0 S0 T1 C1 | 26         | C     | M   | 1.3         | 100 mg/dl     | > 50       | Present                           | Absent    | < 10   | 1 + | 3 + m, g | Mesangial |

Table 1. SAGN and IgAN cases for mass spectrometry analysis. SAGN Staphylococcus infection associated glomerulonephritis, S. cr serum creatinine, EM electron microscopic, m mesangial, g glomerular capillary wall, IF/TA interstitial fibrosis and tubular atrophy, HSP Henoch–Schönlein purpura, IgG IgA, C3 staining by immunofluorescence, C Caucasian, AA African American.
| Disease category | Age (years) | Ethnicity | Sex | Etiology of AKI | Antibiotics | S. Cr | Urine protein | IF/TA |
|------------------|-------------|-----------|-----|----------------|-------------|------|--------------|-------|
| Vanc ATN         | 26          | C         | F   | Cellulitis     | Vancomycin, Keflex, Zosyn, Levofloxacin | 8    | 2 g/d        | Absent |
| Vanc ATN         | 31          | C         | M   | Pneumonia, empyema | Vancomycin | 4.5  | 1.6 g/d      | Absent |
| Vanc ATN         | 60          | AA        | M   | Osteomyelitis   | Vancomycin, Zosyn | 8.4  | 30 mg/dl     | 30%   |
| Vanc ATN         | 63          | C         | M   | Infected gangrenous toe | Vancomycin | 4.5  | 30%          | Absent |
| ATN              | 58          | Hispanic C| M   | Cisplatin nephrotoxicity, dehydration | N/A | 3.7  | 10%          | Absent |
| ATN              | 72          | AA        | F   | Gemzar, Cisplatin, intravenous contrast | N/A | 3.7  | 10%          | Absent |
| ATN              | 59          | C         | F   | Hypotensive shock | N/A | 3.7  | 10%          | Absent |
| ATN              | 45          | AA        | F   | NSAIDs, antibiotics (Ciprofloxacin, Zosyn, Gentamycin) | N/A | 9.9  | 10%          | Absent |
| Baseline Tx biopsy | 58         | C         | F   | LUD | N/A | N/A  | N/A          | N/A   |
| Baseline Tx biopsy | 63         | C         | F   | LUD | N/A | N/A  | N/A          | N/A   |
| Baseline Tx biopsy | 60         | C         | M   | LUD | N/A | N/A  | N/A          | N/A   |
| Baseline Tx biopsy | 45         | C         | M   | LRD | N/A | N/A  | N/A          | N/A   |
| Baseline Tx biopsy | 51         | A         | M   | LUD | N/A | N/A  | N/A          | N/A   |
| Baseline Tx biopsy | 62         | C         | F   | LUD | N/A | N/A  | N/A          | N/A   |
| Baseline Tx biopsy | 63         | AA        | F   | LUD | N/A | N/A  | N/A          | N/A   |
| Baseline Tx biopsy | 57         | C         | F   | LUD | N/A | N/A  | N/A          | N/A   |

Table 2. Cases of vancomycin-induced ATN and ATN of other causes for mass spectrometry analysis. Baseline transplant biopsies are also shown. Vanc ATN- Vancomycin-induced acute tubular necrosis; S.Cr. = serum creatinine; IF/TA = interstitial fibrsis and tubular atrophy; AKI = acute kidney injury, C = Caucasian; AA = African American. C = Caucasian, AA = African American; LUD = living unrelated donor; LRD = living related donor.

proliferative lesions, 11 had focal crescents. Among the 19 biopsies with IgAN MEST score E1, 14 had focal glomerular crescents.

**LC–MS/MS. Glomerular compartment.** A total of 1063 proteins were identified across all samples in the glomerular compartment based on a 2-peptide hit criterion filtering the peptide identifications for 2 ppm mass accuracy and a peptide FDR less than 1%. Protein lists were filtered using a spectral counting rule to remove infrequently observed and low abundant proteins with four or less (≤ 4) spectral counts thus curating the list down to 320 proteins. Several proteins were similarly expressed in SAGN and IgAN glomeruli compared to normal kidney controls (Fig. 1A,B). Fibrinogen (alpha, beta and gamma chains), complement C3, IgA heavy chain, kappa and lambda light chains, and chymotrypsin (SERPINA3) showed high expression in both SAGN and IgAN. LC MS/MS is a semiquantitative method, looking for relative differences in protein expression based on spectral counts. It does not perform absolute protein quantification (Supplemental file 2). Therefore mild quantitative differences in protein molecules (such as IgA heavy chains, complement C3) between SAGN and IgAN may not be detectable by this method, even though C3 tends to show stronger staining in SAGN by immunofluorescence microscopy. Under-expressed proteins in diseased compared to normal glomeruli were mainly hydrolytic enzymes, including aminopeptidase, fractucose-1,6-biphosphatase 1, cytoplasmic aconitase hydratase, arginosuccinate synthase, carbonic anhydrase, and creatine kinase B. Many of these play important roles in energy pathways such as tricarboxylic acid cycle (Kreb’s cycle) in carbohydrate metabolism, acid base balance, and intracellular synthetic functions.

Only a few glomerular proteins showed differential expression between SAGN and primary IgAN (Table 3). Four were monocyte/macrophage-related proteins and were significantly higher in SAGN glomeruli compared to normal and IgAN glomeruli. Conversely, several cytoskeletal and adhesion proteins were down-regulated in SAGN compared to IgAN including alpha-actinin-4, laminin subunit alpha-5, isoform 6 of agrin, and basal cell junction proteins. Glutamine gamma-glutamyltransferase (TGM2), a tubular epithelial stress protein was one of the most highly expressed protein in SAGN. Compared to IgAN, SAGN showed marked downregulation of mitochondrial enzymes, enzymes involved in fatty acid oxidation and carbohydrate metabolism, and proteins involved in amino acid degradation. The differences in tubulointerstitial protein expression were generally more

**Tubulointerstitial compartment.** A total of 1317 proteins were identified across all samples in the tubulointerstitial compartment and after filtering using the same spectral counting rule as for the glomerular compartment, we curated the list down to 622 proteins. In contrast to the glomerular compartment, the tubulointerstitial proteome of SAGN and primary IgAN showed many differentially-expressed proteins (Fig. 2A,B, Table 4). The top upregulated proteins in SAGN compared to IgAN included mainly extracellular matrix and epithelial cell junction proteins. Glutamine gamma-glutamyltransferase (TGM2), a tubular epithelial stress protein was one of the most highly expressed protein in SAGN. Compared to IgAN, SAGN showed marked downregulation of mitochondrial enzymes, enzymes involved in fatty acid oxidation and carbohydrate metabolism, and proteins involved in amino acid degradation. The differences in tubulointerstitial protein expression were generally more
pronounced between SAGN and IgAN (E0, C0) than between SAGN and IgAN (E1, C1). Importantly, changes in the SAGN tubulointerstitium were similar to those in biopsies with ATN of other causes (Fig. 2B).

Pathway analysis also predicted inhibition of metabolic and amino acid degradation pathways in SAGN compared to primary IgAN subgroups, primarily the Kreb’s cycle, fatty acid β-oxidation, glycine betaine degradation, isoleucine degradation, and valine degradation (Fig. 2C,D). In terms of activated and inhibited pathways, the IgAN E1 C1 subgroup appeared intermediate between SAGN and the IgAN E0 C0 subgroup.

Table 3. Proteins with statistically significant differential expression between SAGN and IgAN in the glomerular compartment. Mean spectral counts, fold change and p values are shown.

| Protein identification                              | avg.IgAN | avg.NL.Tx | avg.SAGN | SAGN vs IgAN | Fold change | p value |
|-----------------------------------------------------|----------|-----------|----------|--------------|-------------|---------|
| Myeloid cell nuclear differentiation antigen         | 0.20     | 0.35      | 0.81     | 3.70         | 0.0120      |         |
| Thymidine phosphorylase                              | 0.39     | 0.56      | 1.69     | 3.97         | 0.0104      |         |
| Protein S100-A9                                      | 0.79     | 1.98      | 3.43     | 4.02         | 0.0055      |         |
| Isoform 3 of Myoferlin                               | 0.33     | 0.70      | 1.42     | 4.03         | 0.0095      |         |
| Coronin-1A                                           | 0.26     | 0.51      | 1.22     | 4.40         | 0.0056      |         |
| Lysosome C                                           | 0.53     | 0.94      | 2.71     | 4.74         | 0.0027      |         |
| Alpha-1-antichymotrypsin                            | 0.40     | 0.59      | 2.18     | 5.00         | 0.0024      |         |
| Basal cell adhesion molecule                         | 12.17    | 12.90     | 6.47     | −2.03        | 0.0080      |         |
| Alpha-actinin-4                                      | 89.96    | 93.46     | 66.15    | −1.34        | 0.0098      |         |
| Laminin subunit alpha-5                              | 55.83    | 56.31     | 39.14    | −1.42        | 0.008       |         |
Vancomycin-induced ATN and ATN of other causes showed changes similar to those in SAGN. As in the glomerular compartment, LXR/RXR pathways were activated in primary IgAN, (especially the E0 C0 subgroup of IgAN) but not in SAGN. RhoA signaling, acute phase response signaling, remodeling of epithelial adherens junctions, BAG2 (Bcl2 associated athanogene 2) signaling were upregulated in SAGN and ATN subgroups as compared to the IgAN E0 C0 group (Fig. 2C). In contrast to the glomerular proteome, the Hippo pathway was inhibited in the tubulointerstitial compartment of all the diseased kidneys compared to normal kidney.

**Immunohistochemical Staining (IHC).**  
**Glomerular compartment.** IHC staining for lysozyme (Fig. 3A−C), CD68 and S100A9 was performed on a total of 67 biopsies including subgroups SAGN, IgAN MEST E0 and IgAN MEST E1. Additionally 5 biopsies each of ATN and normal kidney were also stained. The number of cells that were positively stained for each marker was significantly higher in SAGN than IgAN E0 and E1, ATN

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**Figure 2.** The tubulointerstitial proteome of SAGN and IgAN. (A) Mean protein expression using dispersion plots—proteins that are differentially expressed are in blue while proteins that are not are in yellow. Fold-change (in log2 scale) is plotted against the mean of normalized protein expression. Each dot represents an individual protein. A fold-change cut-off of ≥2 or ≤-2 in combination with a p value cutoff of ≤0.01 was used for detecting differential expression. Many proteins were differentially expressed. (B) Heat map with unsupervised hierarchical clustering analysis of the top differentially expressed tubulointerstitial proteins between SAGN and IgAN for greater fold difference, p < 0.01. Each column represents an individual biopsy sample, and each row represents an individual protein. Blue represents higher expression and yellow represents lower expression. The degrees of similarity in expression levels are presented in the dendrogram. R version 3.6.0. (C) Differentially-expressed tubulointerstitial signaling pathways between SAGN (heat map column 1), vancomycin-induced ATN (heat map column 2), ATN of other causes (heat map column 3), primary IgAN-Oxford Class E1 (heat map column 4), and primary IgAN-Oxford Class E0 (heat map column 5). Orange indicates pathways predicted to be activated while blue indicates pathways predicted to be inhibited. (D) Glycine betaine degradation pathway in SAGN as seen by IPA (Qiagen Bioinformatics). Sarcosine dehydrogenase is downregulated (shown in green and labeled as sarcosine oxidase). This correlates with the significantly low spectral counts for sarcosine dehydrogenase in SAGN biopsies by mass spectrometry.
| Identified proteins | Spectral counts (group means) | SAGN vs IgAN E0 | SAGN vs IgAN E1 | SAGN vs ATN | SAGN vs Vanc ATN |
|--------------------|-------------------------------|------------------|----------------|-------------|-----------------|
| Normal Kidney     | IgAN E0 | IgAN E1 | SAGN | ATN | Vanc ATN | Fold change | p value | Fold change | p value | Fold change | p value | Fold change | p value |
| Alpha-1-antichymotrypsin | 0.37 | 0 | 5.21 | 6.43 | 5.22 | 6.44 | 1.00E−04 | 1.21 | 0.6274 | 1.17 | 0.6855 | 1.04 | 0.9225 |
| Isoform 17 of Fibronectin | 1.84 | 1.72 | 7.06 | 21.42 | 25.09 | 17.75 | 6.09 | 1.00E−04 | 2.54 | 0.0219 | −1.2 | 0.6503 | −1.14 | 0.7487 |
| PDZ and LIM domain protein 5 | 0.48 | 0 | 1.54 | 6.27 | 3.19 | 1.38 | 6 | 4.00E−04 | 2.63 | 0.0523 | 1.56 | 0.3659 | −2.95 | 0.0306 |
| Ras GTPase-activating-like protein IQGAP1 | 2.08 | 0.37 | 3.54 | 7.56 | 14.34 | 9.23 | 5.71 | 1.00E−04 | 1.95 | 0.0657 | −1.89 | 0.0497 | 1.21 | 0.5642 |
| Protein-glutamine gamma-glutamyltransferase 2 | 0 | 1.64 | 5.28 | 12.95 | 7.21 | 7.29 | 5.29 | 1.00E−04 | 2.19 | 0.0238 | 1.65 | 0.1425 | −1.64 | 0.1442 |
| Fibrinogen gamma chain | 0.36 | 0 | 1.39 | 5.23 | 7.14 | 10.75 | 5.13 | 9.00E−04 | 2.44 | 0.0544 | −1.37 | 0.4658 | 2 | 0.1063 |
| Plastin-2 | 0.22 | 0 | 0.91 | 4.56 | 3.74 | 5.06 | 4.52 | 0.0033 | 2.78 | 0.0436 | 1.09 | 0.8611 | 1.14 | 0.7942 |
| Cytoskeleton-associated protein 4 | 0 | 0.7 | 0.44 | 7.65 | 4.18 | 2.25 | 4.4 | 0.0039 | 5.42 | 0.001 | 1.46 | 0.4435 | −2.35 | 0.089 |
| Isoform 2 of Septin-2 | 1.46 | 0.95 | 3.54 | 6.51 | 6.98 | 5.13 | 3.68 | 0.0038 | 1.68 | 0.1904 | −1.1 | 0.8045 | −1.22 | 0.6131 |
| Zyxin | 0.12 | 0.7 | 1.85 | 5.01 | 3.58 | 1.99 | 3.55 | 0.0114 | 2.09 | 0.1225 | 1.24 | 0.6403 | −2.09 | 0.1238 |
| Fibrinogen beta chain | 0.36 | 0.75 | 1.91 | 5.52 | 6.36 | 13.01 | 3.46 | 0.0106 | 2.35 | 0.0633 | −1.17 | 0.722 | 2.22 | 0.0613 |
| Isoform 3 of Plectin | 2.74 | 5.48 | 9.15 | 23.52 | 23.22 | 13.98 | 3.32 | 0.0029 | 2.23 | 0.0325 | −1.01 | 0.9682 | −1.54 | 0.2402 |

| Lower expression in SAGN compared to IgAN |
|-------------------------------------------|
| Sarcosine dehydrogenase mitochondrial | 9.27 | 8.79 | 3.08 | 0 | 1.85 | 1.91 | −8.53 | 0 | −3.49 | 0.0072 | −2.28 | 0.0884 | 2.58 | 0.0473 |
| Bifunctional ATP-dependent dihydroxyacetone kinase/FAD-AMP lyase (cyclizing) | 18.28 | 16.78 | 18.89 | 1.65 | 9.41 | 9.6 | −5.66 | 0 | −6.47 | 0 | −3.43 | 0.001 | 3.45 | 0.001 |
| Formimidoyltransferase-cyclodeaminase | 9.56 | 13.76 | 7.99 | 1.34 | 5.41 | 8.24 | −5.36 | 1.00E−04 | −3.38 | 0.0029 | −2.36 | 0.0409 | 3.48 | 0.0024 |
| L-xylulose reductase | 9.94 | 11.07 | 9.6 | 1.27 | 6 | 7.63 | −4.7 | 1.00E−04 | −4.24 | 1.00E−04 | −2.74 | 0.0105 | 3.42 | 0.0015 |
| Isoform B of Band 4.1-like protein 3 | 11.27 | 9.33 | 5.62 | 1.27 | 2.76 | 2.97 | −4.31 | 3.00E−04 | −2.66 | 0.0146 | −1.5 | 0.3378 | 1.65 | 0.2337 |
| Isoform 2 of 4-hydroxyphenylpyruvate dioxygenase | 6.12 | 5.96 | 4 | 0 | 1.7 | 2.18 | −5.33 | 6.00E−04 | −4.08 | 0.0035 | −2.15 | 0.1232 | 2.62 | 0.0503 |

Continued
and normal kidney (Fig. 3D–F, Table 5). The differences between IgAN E1 and IgAN E0 were not found to be statistically significant, except for lysozyme (p = 0.06 by non-parametric Dunn test for all pairwise comparisons; and p = 0.0015 using Tukey’s HSD parametric test applied to log-scale data, details not shown). Double immuno-fluorescence staining with antibody pairs performed on SAGN biopsies showed a 65% co-localization between lysozyme and CD68, but only a 31% co-localization between S100A9 and CD68 (Fig. 4A–G).

To determine whether glomerular staining for lysozyme, CD68 or S100A9 could be used to distinguish between SAGN, IgAN E0 and IgAN E1, ROC analysis was done (Fig. 3G–I). S100A9 was the most robust marker with an area under the ROC curve (AUC) of 0.98 for IgAN E0 versus SAGN and 0.94 for IgAN E1 versus SAGN (Table 5). Using logarithms of lysozyme, S100A9 and CD68-positive cells, two variable-predictors in the logistic regression model were also evaluated, but the single predictor based on S100A9 gave the best results.

These histologic studies were validated in an independent set of 30 kidney biopsies. The achieved sum of sensitivity and specificity for the validation set was comparable to what was achieved for the training set (Table 5).

Tubulointerstitial compartment IHC. IHC staining for two of the differentially expressed proteins—transglutaminase 2 (TGM2) and l-xylulose reductase (DCXR) was performed on 9 SAGN biopsies, 6 IgAN E0 biopsies, 6 IgAN E1 biopsies, and 6 ATN biopsies (etiologies being vancomycin toxicity, thrombotic microangiopathy, rhabdomyolysis, oxalate nephropathy, and minimal change disease), Fig. 5A–H. Both TGM2 and DCXR were found to be constitutively expressed in endothelial cells of the glomerular and peritubular capillaries, but not in the tubules or interstitium of normal kidney (Fig. 5A, E). Quantification was performed using image analysis software, calculating the ratio of brown pixel area to total pixel area within the biopsy sample (Aperio Imagescope ver. 12.3.0.5056 (Leica Biosystems Inc., Buffalo Grove, IL). These ratios were expressed as percentage. Distribution of the average percentage by group are shown in Fig. 5I, J for TGM2 and DCXR respectively. For TGM2 the highest mean ratio was seen for SAGN (74 ± 15.2), followed by ATN (52.5 ± 13), IgAN E1 (27.4 ± 17),

| Identified proteins | Normal Kidney | IgAN E0 | IgAN E1 | SAGN | ATN | Vanc ATN | Fold change | p value | Fold change | p value | Fold change | p value | Fold change | p value |
|---------------------|---------------|---------|---------|------|-----|---------|------------|--------|------------|--------|------------|--------|------------|--------|
| Dimethylglycine dehydrogenase, mitochondrial | 17.8 | 15.06 | 12.64 | 3.17 | 6.84 | 9.38 | − 3.42 | 6.00E−04 | − 2.99 | 0.0015 | − 1.69 | 0.1444 | 2.26 | 0.0207 |
| Choline dehydrogenase, mitochondrial | 12.25 | 13.67 | 10.42 | 3.65 | 5.27 | 6.8 | − 3.16 | 7.00E−04 | − 2.5 | 0.0056 | − 1.36 | 0.3832 | 1.66 | 0.1423 |
| Glutaryl-CoA dehydrogenase, mitochondrial | 5.01 | 6.92 | 2.96 | 0.66 | 3.39 | 2.78 | − 4.49 | 8.00E−04 | − 2.29 | 0.0058 | − 2.43 | 0.0501 | 2.16 | 0.0899 |
| Ribosyl-dihydronicotinamide dehydrogenase (quinone) | 6.33 | 5.38 | 2.28 | 0.31 | 1.01 | 2.17 | − 4.75 | 9.00E−04 | − 2.29 | 0.0803 | − 1.34 | 0.5507 | 2.24 | 0.0906 |
| Isoform L-type of Pyruvate kinase PKLR | 7.58 | 9.48 | 6.73 | 1.69 | 1.54 | 3.42 | − 3.81 | 9.00E−04 | − 2.86 | 0.0008 | 1.15 | 0.7569 | 1.57 | 0.2821 |
| Glycine dehydrogenase (decarboxylating), mitochondrial | 6.97 | 7.12 | 5.48 | 0.72 | 3.12 | 2 | − 4.35 | 0.0012 | − 3.59 | 0.004 | − 2.18 | 0.0911 | 1.55 | 0.3473 |
| Isoform 2 of Amiloride-sensitive amine oxidase [copper-containing] | 7.6 | 5.49 | 2.21 | 0.66 | 0 | 0 | − 4.07 | 0.0036 | − 1.97 | 0.1617 | 1.86 | 0.2316 | − 1.89 | 0.2185 |

Table 4. Tubulointerstitial compartment proteins in SAGN, IgAN, Vancomycin-induced ATN and ATN of other causes. Mean peptide counts, fold changes and p values for selected proteins with highest fold changes between SAGN and IgAN are shown. Vanc ATN Vancomycin-induced acute tubular necrosis, ATN acute tubular necrosis of other causes.
IgA E0 (13.4 ± 11) and lowest was in the normal kidney samples (2.5 ± 2.1). Conversely, DCXR showed the highest ratios in normal kidney, followed by IgAN, as compared to the SAGN and ATN groups.

**Discussion**

A comparison of the glomerular and tubulointerstitial proteomes from SAGN and primary IgAN kidney biopsies was carried out to identify differences and potential biopsy-based biomarkers that could help discriminate between these conditions. Our study found that the glomerular proteomes of IgAN and SAGN were very similar, except for a significant upregulation of inflammatory protein expression in SAGN glomeruli. The
tubulointerstitial proteome of SAGN however showed marked differences from IgAN, and in fact resembled that of biopsies from patients with ATN of various causes. The tubulointerstitial proteome of IgAN was very similar to that of normal kidney.

The glomeruli from SAGN patients contained high levels of leukocyte-derived proteins known to be involved in the response to infection. Lysozyme in serum and other body fluids is antimicrobial and originates mostly from monocytes/macrophages and to a lesser extent from neutrophils. S100A9 is a calcium-binding protein highly expressed in the cytosol of neutrophils and monocytes, and participates in multiple inflammatory processes including bacterial infections, autoimmune diseases, complement activation and even aging processes.

The average age of SAGN patients in our cohort was higher than the IgAN patients (individual patient ages shown in Table 1), raising the possibility that aging contributed to inter-group differential expression of S100A9. However, even within the SAGN group, differential staining was observed (Fig. 3F), suggesting that the degree of glomerular inflammation was the main contributing factor. Myeloid nuclear differentiation antigen (higher in SAGN, shown in Table 3) is constitutively expressed by monocytes and neutrophils and can be upregulated by interferon-alpha, a cytokine increased during infections.

IHC staining confirmed that lysozyme, CD68, and S100A9-positive cells were significantly more numerous in SAGN glomeruli compared to IgAN, and not only the E0, but also the E1 subgroup of IgAN. This is important because IgAN E1 is more inflammatory than E0 and therefore more likely to be confused with SAGN histologically.

This finding is also consistent with the higher frequency and severity of glomerular endocapillary hypercellularity seen in SAGN biopsies as compared to IgAN biopsies as we have shown previously (endocapillary hypercellularity in 60% of SAGN biopsies as compared to only 10% of IgAN). Endocapillary hypercellularity is thought to be due to infiltrating macrophages and admixed neutrophils in the glomerular capillaries. Counting endocapillary neutrophils and macrophages on routine hematoxylin and eosin stained slides is practically difficult. The immunostains help to better delineate the glomerular inflammatory cells. We further showed that lysozyme and S100A9 do co-localize with the macrophage marker CD68, suggesting that infiltrating macrophages were at least partially responsible for glomerular lysozyme and S100A9 expression. Co-localization was 65% for lysozyme but only 31% for S100A9, suggesting that lysozyme and S100A9 may be staining different macrophage populations or that S100A9 may also stain neutrophils.

It is conceivable that S100A9 staining could be used to differentiate SAGN from IgAN in ambiguous clinical situations. A single predictor model based on S100A9 showed the best ability to discriminate between SAGN and IgAN. However, it should be emphasized that occasional cases of active proliferative IgAN may show high numbers of S100A9 (and lysozyme, CD68) positive cells, but such cases are rare particularly in the Western population. Interpretation of staining results however, must be done within clinical context.

Actin-binding cytoskeletal proteins and cell adhesion proteins were downregulated in SAGN glomeruli compared to IgAN and normal glomeruli, correlating well with the predicted inhibition of epithelial adherens junction remodeling pathway on IPA. These changes may be the cause or effect of increased podocyte injury in SAGN glomeruli, and may facilitate the often encountered nephrotic-range proteinuria in SAGN.
pathway, also predicted to be activated in SAGN glomeruli may result in podocyte damage\textsuperscript{27–29}, and contribute to the nephrotic range proteinuria.

The tubulointerstitial proteome of SAGN showed significantly higher expression of TGM2, compared to IgAN, also shown by IHC staining. TGM2, a marker of tubular injury\textsuperscript{30–32}, is a member of the protein glutamine γ-glutamyltransferase enzyme family that catalyzes the calcium-dependent crosslinking of protein targets in the extracellular matrix (ECM)\textsuperscript{33–35}. Secretion of TGM2 from cortical tubular epithelial cells into the surrounding interstitium has been shown to contribute to fibrotic remodeling of the kidney and the development of chronic kidney disease\textsuperscript{31,32}. Hippo signaling was predicted to be decreased and Rho signaling was predicted to be increased in the SAGN tubulointerstitium, possibly predisposing to interstitial fibrosis\textsuperscript{36–38}. In addition, the SAGN interstitium also showed lack of activation of pathways that could potentially protect against the development of chronic kidney damage, such as PPARα, LXR-RXR and eNOS signaling\textsuperscript{39,40}. The latter were more prominently expressed in IgAN in our study as well in a study by Liu et al.\textsuperscript{41}.

This is not to say that SAGN (and ATN) does not activate any protective mechanisms. For example, the predicted activation of BAG2 signaling and inhibition of ERK5 in SAGN glomeruli could possibly exert a protective effect through inhibition of apoptosis and glomerular fibrosis\textsuperscript{42,43}. The interactions of these complex cellular pathways and their net effect likely depends on the surrounding milieu which may be different at different stages of the disease\textsuperscript{44}.

The SAGN tubulointerstitial proteome also demonstrated a marked down-regulation of enzymes involved in energy generation and metabolism, likely contributing to tubular cell dysfunction and ATN. An interesting example is the marked decrease in DCXR expression in SAGN. DCXR is a multi-functional enzymatic protein playing a role in the uronate cycle of glucose metabolism and production of organic intracellular osmolytes needed in tissue osmoregulation and water absorption in the proximal tubule\textsuperscript{45}. Loss of this enzyme in SAGN may predispose to tubular osmotic stress. However, this may be mitigated to some extent by the concomitant depression of specific amino acid degradation pathways like the glycine betaine degradation pathway. Glycine betaine is an osmoprotectant which accumulates in cells in response to osmotic stress\textsuperscript{46}. The degradation of glycine betaine proceeds by sequential enzymatic demethylation (Fig. 2D) and the last step is catalyzed by

**Figure 5.** (A–D) Immunohistochemical staining for glutamine gamma-glutamyltransferase (TGM2), and (E–H) Xylulose reductase (DCXR) staining. Tissue from normal kidney (A, E), IgAN E0 (B, F), SAGN (C, G), and vancomycin-induced ATN (D, H) was stained. Strong tubulointerstitial expression of TGM2 is seen in SAGN and ATN compared to normal and IgAN biopsies. Conversely, there is decreased expression of DCXR in SAGN and ATN compared to normal and IgAN biopsies. (I, J) Quantification of IHC staining for TGM2 and DCXR in each biopsy was performed using image analysis software, by calculating ratio of brown pixel area to total pixel area within the biopsy sample (Aperio ImageScope ver. 12.3.0.5056 (Leica Biosystems Inc., Buffalo Grove, IL, USA). These ratios were expressed as percentage. Distribution of the average percentages are shown by group for TGM2 (I) and DCXR (J).
enzyme sarcosine dehydrogenase. Sarcosine dehydrogenase was found to be markedly under-expressed in SAGN tubulointerstitial proteome.

Together these data suggest that SAGN is an aggressive form of glomerulonephritis accompanied by marked acute tubular injury as compared to primary IgAN. Interestingly, IgAN with endocapillary hypercellularity (Oxford MEST Score E1) had proteomic signatures in between SAGN and IgAN without endocapillary hypercellularity (Oxford MEST Score E0).

In summary, our study sheds light on the more inflammatory profile of SAGN compared to primary IgAN. These appear to be separate disease entities, but with plausible overlaps at some points in their downstream pathogenetic pathways such as glomerular IgA deposition. The mean spectral counts for IgA heavy chain in SAGN biopsies however were lower than in IgAN (4.2 versus 13.9) comparing well with the consistently lower intensity of IgA staining in SAGN biopsies compared to primary IgAN, and also the absence of IgA in up to 25% of SAGN biopsies. It must be acknowledged that the analytic approach used here examined proteins observed in most tissue samples and compared only proteins with high spectral counts. While this approach improves confidence in the observed results, alternative approaches, for example assessment of mass intensities, would potentially increase proteomic depth and possibly identify disease-specific biomarkers. Also it is should be noted that glomerular lesions (especially crescents) can be focal in distribution. This can result in inter-glomerular proteome heterogeneity and subsequent analytical differences. We have combined the glomeruli present in each biopsy, therefore inter-glomerular heterogeneity was minimized. The proteomic differences between SAGN and IgAN emerging from this study appear more quantitative rather than qualitative with no disease-specific biomarkers. Nonetheless, our immunostaining results with the combination of inflammatory macrophage markers (S100A9 and lysozyme) and tubular stress marker TGM2 may aid diagnosis in ambiguous cases of SAGN.

Methods
All methods were carried out in accordance with relevant guidelines and regulations. Archived tissue specimens remaining in paraffin embedded blocks after routine diagnostic work was already completed were used for this study. There was no prospective sample collection from human subjects and no direct patient interaction (approved waiver for informed consent). The study was approved by the Ohio State University Institutional Review Board (IRB Protocol number 2011H0364).

Tissue collection for LC–MS/MS. Four 10 μm sections were cut from the paraffin-embedded tissue blocks and mounted on thermoplastic (polyethylene naphthalate covered) glass slides (Carl Zeiss MicroImaging). Blades and water bath were changed before each case. Slide processing, laser capture, tissue collection by catapulting, protein retrieval and preparation for LC MS were performed in Supplementary File 1. Laser capture was used to separately collect the glomeruli and tubules.

Samples (1.5 μg) were analyzed with nanoLCMS using a 3 h 1D (C18) RP gradient and a LTQ-Orbitrap ELITE mass spectrometer collecting MS2 data by ETD/CID decision tree analysis. Details provided in Supplementary File 2. Data were annotated through ProteomeDiscoverer 1.4 using Mascot (v2.4) and Sequest HT algorithms with a merged human and Staphylococcus aureus protein databases. Results were loaded into Scaffold 4 for comparative proteomics. Protein and peptide FDR were minimized using Peptide/Protein Prophet, elimination of all decoy hits, and the 2-peptide hit rule.

Immunohistochemistry (IHC). IHC was performed for representative proteins selected on the basis of their differential expression pattern between SAGN and primary IgAN and their potential applicability to diagnostic pathology. These included lysozyme, S100A9, protein-glutamine gamma-glutamyl transferase 2 (TGM2) and L-xylulose reductase (DCXR). CD68 staining was also performed, since it is a routinely used IHC macrophage maker. Appropriate secondary antibodies were used (Supplementary data file 3). Lysozyme and S100A9 are leukocyte, mainly macrophage-derived proteins and are thought to reflect inflammation.TGM2 and DCXR are expressed by renal tubular epithelial cells. All IHC staining was performed using an indirect immunoperoxidase method on FFPE tissue sections cut at 3 microns after antibody validation (Supplemental data file 3). Appropriate positive (placental tissue for TGM2 and tonsillar lymphoid tissue for the inflammatory markers) and negative controls (both rabbit non-immune serum; and omission of primary antibody) were included.

Double staining and co-localization studies were done for lysozyme, S100A9 and CD68. Sections were cut at 3 microns, deparaffinized, and subjected to HIER using Target Retrieval Solution (Agilent). The slides were then treated with Protein Block, Serum-Free (Agilent) for 10 min prior to the first primary antibody. CD68 at 1:100 for 1 h. This was followed by a 1 h incubation using DyLight 488-conjugated, AffiniPure Donkey Anti Mouse IgG (H+L) (Jackson ImmunoResearch Laboratories, INC) at a dilution of 1:50. This was followed by staining with the second primary antibodies against lysozyme (diluted 1:100) or S100A9 (diluted 1:200) for 1 h, followed by a 30 min incubation with the secondary antibody, DyLight 594-conjugated, AffiniPure Donkey Anti Rabbit IgG (H+L) at a dilution of 1:50. (Jackson ImmunoResearch Laboratories, INC). For each marker pair, images of ten different glomeruli within the biopsy sample were captured and percent co-localization was calculated using a custom image analysis workflow in R (Supplementary file 4).

Negative controls with secondary antibody alone were performed and showed no stain. Glomerular images were captured on a Carl Zeiss Axio Imager Z.1 with LSM700 confocal microscope. This microscope used a motorized stage at 20× objective with a pinhole of 0.5 Airy unit, scan speed of 8, and a resolution of 512×512 to capture images.

Statistical analysis. LC–MS/MS data. Glomerular and tubulointerstitial proteins were studied separately. To ensure a robust comparison of proteomic data without imputation of missing values for proteins having
low abundance (i.e. spectral counts ≤ 4) for at least 90% samples were excluded from further statistical analysis. Prior to statistical comparisons the protein spectral counts were normalized across samples. To compare protein abundance between groups, we applied negative binomial generalized linear models through a R package called DESeq2 (R version 3.6.0)34. DESeq2 method was used for comparing protein mean expression between groups. Dispersion estimation and fold-change estimation were then improved by applying an Empirical Bayes shrinkage method. The significance level was determined by controlling the mean number of false positives at 1 out of every 100 proteins32. Top proteins were selected by both fold-change (≥2) and the significance level (≤0.01). After excluding 3 outliers (biopsies with overall low glomerular spectral counts), hierarchical clustering on heat maps as well as principal component analysis plots (PCA) were performed.

Data files for acquired LCMS data (.RAW), for peak lists (.mgf), and scaffold search results (.sf3) files along with a sample key and sequence database were deposited in MassIVE (https://massive.ucsd.edu/) data repository (MassIVE ID: MSV000085473) with the Center for Computational Mass Spectrometry at the University of California, San Diego and shared with the ProteomeXchange (www.proteomexchange.org) (Proteome Exchange ID: PXD019422). Data with uniprot numbers is provided in Supplemental files 5 and 6.

**IHC data.** Immunostaining for lysozyme, S100A9 and CD68 was performed on individual tissue sections. Positively stained cells within the glomerular capillary tuft were manually counted using criteria described by Soares et al.35. A positive cell was defined by clearly delimited aggregates of brown-stained cytoplasmic granules with or without a juxtaposed nucleus. Counts were performed on every non-globally sclerosed glomerulus in the biopsy sample at 400× magnification. An average count for each biopsy was calculated and used for statistical comparisons. Pairwise comparisons of mean counts for IgAN E0, IgAN E1 and SAGN groups were made using both non-parametric Dunn test and Tukey’s HSD test for the log-transformed counts that were found to be normally distributed. The normal kidney and ATN cases were also stained and counted. Classification rules based on each of the three IHC markers were developed and the associated sensitivity and specificity values were computed for distinguishing IgAN E0 and IgAN E1 from SAGN. Threshold values that yielded the maximum sum of sensitivity and specificity were used to determine optimal classification rules. Receiver operating characteristic (ROC) curves were also generated. Two predictor logistic regression models using combinations of the 3 IHC markers were also developed and ROC curves were generated. These analyses were done on JMP Version 13 (SAS Institute, Cary, NC, USA). Level of significance was set at 0.05 in all analyses.

**Ingenuity pathway analysis.** Qiagen’s Ingenuity Pathway Analysis (IPA, Qiagen Redwood City www.qiagen.com/ingenuity) was used to identify canonical pathways that were enriched in the various groups and separately in the glomerular and tubulointerstitial compartments. All proteins that were used for statistical analysis were used for IPA analysis. Significant pathways were determined by Fisher’s exact test, right tailed and presented as −log( p value). The statistically significant functional pathways in each disease state, and the predicted activation (or inhibition) state of these functional pathway end-points in each group was determined by the z score algorithm calculated by IPA software.
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Author contributions
A.A.S.—the main idea, case selection, tissue procurement, laser capture experiments, cell counting, writing the manuscript, FAMEPro Grant awardee for the IHC work, data analysis/interpretation; J.S.—helped with mass spectrometry data curating and selecting antibodies for IHC; M.J.—double immunofluorescence and colocalization studies; C.B.—immunofluorescence staining; S.P.—help with laser capture work; S.B.—cell counting on IHC; L.Y. and N.H.—statistical analysis; D.W., M.M. and J.K.—mass spectrometry experiments; T.N.—critical review; B.R.—critical review, help with writing the manuscript, data analysis/interpretation, PI on U01 Grant.

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Competing interests
The authors declare no competing interests.

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