Proteinuria in *Cynomolgus macaques* (*Macaca fascicularis*) with Spontaneously Developed Metabolic Disorder and Diabetes: Transcriptome Analysis of Biopsy Kidney

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**Abstract**

Although Type 2 Diabetes Mellitus (T2DM) is well characterized Non-Human Primate (NHP), the phenotypes of diabetic nephropathy, and its molecular mechanisms are not well studied in NHPs. Diabetic nephropathy in *Cynomolgus macaques* with spontaneous dysmetabolism (Pre-DM, n=19) and diabetes (DM, n=20) were compared with normal controls (N, n=11). There were 9 NHPs with albuminuria (>42 mg/day) in the DM (45%), only 1 in Pre-DM and none in N group. The renal function measured by estimated glomerular filtration rate (eGFR) was not significantly different among the 3 groups, indicating that these NHPs were in an early stage of renal disease. From these NHPs, 3 N, 3 Pre-DM without albuminuria and 6 DM with albuminuria were selected for transcriptome analysis of kidney biopsies. There were 95 differentially expressed genes (DEGs) detected amongst the 3 groups, of which, 75DEGs between the N and DM related to diabetic nephropathy; 66 DEGs between the N and Pre-DM related to dysmetabolism without nephropathy; 68 DEGs between N and both Pre-DM & DM related to dysmetabolism; but only 1 nephropathy specific gene (LCT lactase) between the DM with albuminuria (DM) and Pre-DM without albuminuria; only 4 DEGs between the DM with albuminuria and both N & Pre-DM without albuminuria specific to nephropathy. Signaling pathway analysis of the relevant DEGs and encoded proteins highlighted the role of a kidney failure, renal and urological diseases, and inflammatory diseases related network, in which the most pivotal gene in this network is Tumor Necrosis Factor (TNF), indicating that nephropathy is a disease closely related to inflammation and cell death. Thus, the present study was the first detailed characterization of the diabetic nephropathy phenotypes and the kidney histopathological changes in NHPs and provided molecular insights into novel mechanisms of disease progression, potential new drug targets as well as specific diagnostic biomarkers.

**Keywords:** Micro-albuminuria; Proteinuria; Glycosuria; Renal function; Kidney biopsy; Trascriptome analysis; Affymetrix genechip microarray

**Abbreviations:** DM: Diabetes Mellitus; Pre-DM: Prediabetic or Dysmetabolic; ivGTT: Intravenous Glucose Tolerance Test; Kglucose: Glucose Disappearance Rate; Dvolume: Total Drinking Volume in 24 hours (mL); Uvolume: Total Urine Volume in 24 hours (mL); Uprotein: Total Urinary Protein Excretion in 24 hours (mg); Ucreatinine: Total Urinary Creatinine Excretion in 24 hours (mg); Ualbumin: Total Urinary Albumin Excretion in 24 hours (mg); ACR: Urinary Albumin to Creatinine Ratio; GFR: Glomerular Filtration Rate; BUN: Blood Urea Nitrogen Concentration; DEG: Deferentially Expressed Gene

**Introduction**

Long standing diabetes mellitus with characterization of nephrotic syndrome is the major cause of End-Stage Renal Disease (ESRD) that leads to dialysis in 20-40% patients with kidney disease. During the present decade, 30% of the predicted $1.1 trillion medical costs of dialysis world-wide will result from diabetic nephropathy. A study in Diabetes Care revealed that Type 2 Diabetes Mellitus (T2DM) patients with micro-albuminuria or diabetic retinopathy were more likely to progress to macro-albuminuria and a faster decline in renal function in the late stage compared with those who did not have these conditions [1]. Treatment of T2DM patients can delay the development of micro-albuminuria, e.g., the Angiotensin II Converting Enzyme (ACE) inhibitor, trandolapril, or the angiotensin-receptor blocker, olmesartan, both have shown to delay the T2DM-related micro-albuminuria [2-7]. Currently this disease is not recognized early enough because of inadequate diagnostic methods, which increases the chances that early nephropathy with micro-albuminuria will progress toward ESRD. Thus, in addition to micro-albuminuria that has been commonly used as an early biomarker of diabetic nephropathy, there is still an unmet medical need to identify novel and highly sensitive biomarkers for detecting diabetic nephropathy at an early stage, and possibly also to monitor disease progression or regression after drug therapy. Some biomarkers present in blood and urine are associated with injury at specific sites in the nephron, which could help to identify and localize the injury earlier and even understand the mechanism of the pathogenesis of disease progression.

Unlike conventional experimental rodent models of diabetic nephropathy, of which, the relevance to human pathophysiology and metabolic kinetics remains unclear, it has been well-recognized that...
the Non-Human Primate (NHP) model of diabetes and metabolic disorders is the most clinically translatable animal model that resembles all the disease characteristics in patients at different stages of metabolic disorder and diabetes, including the progressive development of the known complications, such as diabetic nephropathy, thus providing an ideal animal model for investigating the mechanisms underlying human T2DM and diabetic nephropathy as well as for pharmaceutical discovery, development and evaluation of novel therapeutic agents [8]. Although there are some reports including the present laboratory with detailed characterization of the NHP model of diabetes and metabolic disorders, the diabetic complications, especially diabetic nephropathy in NHPs and its underline molecular mechanisms have not yet been well characterized [8-11].

Therefore, the aims of the present study were to
1. Characterize 50 cynomolgus monkeys at different stages of spontaneously developed metabolic disorders and diabetes for nephropathy phenotypes, such as urinary albumin, protein, glucose and creatinine excretion, kidney functions, etc.;
2. Examine the histopathological changes in the diabetic nephropathy kidneys;
3. Select 12 representative monkeys under 3 categories, normal control (N), pre-diabetic without albuminuria (Pre-DM) and diabetic with albuminuria (DM) for transcriptome analysis of kidney biopsy specimen.

The present study was the first detailed characterization of the diabetic nephropathy phenotypes and the kidney histopathological changes in NHPs and provided molecular insights into novel mechanisms of disease progression, potential new drug targets as well as specific diagnostic biomarkers.

Research Design and Methods

Animals

Fifty Cynomolgus macaques (Macaca fascicularis) of both genders were individually housed at Crown Bioscience Inc., Taicang, China in species appropriate cages in temperature-controlled rooms maintained at 23 ± 3°C on a 12-hour light-dark cycle. They were fed normal primate chow containing 19% protein, 5% fat and 3.6% fiber (Shanghai Shilin Biotechnology Inc., Shanghai, China) twice daily and had free access to tap water. All animals were weight stable prior to study initiation and supplemented to maintain sedation during the entire experiment period. The monkeys were divided into 3 distinct groups: metabolically normal (N), prediabetic or dysmetabolic (Pre-DM), and overtly diabetic (DM).

Renal functional measurements

The 24 hour urine collection was carried out with the monkeys kept individually in a stainless steel metabolic cage with a separation net being placed between the cage and the urine collection tray (Suzhou Houhuang Animal Laboratory Equipment Technology Co. Ltd, China) to prevent the feces and residual food from contaminating the urine. An intra-venous blood sample was taken on the next day after urine collection for the biochemical measurements by a clinical analyzer (Roche, Cobas 601). Renal function was evaluated by the clinically relevant surrogate markers, Blood Urea Nitrogen (BUN) and serum creatinine (Screatinine) concentrations, as well as Estimated Glomerular Filtration Rate (eGFR) based on creatinine clearance with the equation of Ucreatinine/Screatinine × 1.73 / body weight2/3 × 0.118, which is commonly used in the clinic as a “first line” test of kidney function [12-14].

Histopathology

The kidneys from representative normal and diabetic nephropathy monkeys sacrificed for other IACUC approved studies with no kidney toxicity were dissected through laparotomy, cut open along coronary axis, fixed in 10% formalin (Sinopharmard), paraffin embedded (Leica EG11150G), cut by a microtome (Leica RM2245) at 5 µm thick, H&E and PAS stained, examined under a microscope (Nikon, TI-DH) and the image was taken with a cool CCD camera.

Percutaneous kidney biopsy

Twelve monkeys, 3 N, 3Pre-DM without albuminuria and 6DM with albuminuria, were selected for kidney biopsy under ketamine anesthesia after an overnight fast. A 18 G semi-automotive biopsy needle was used to penetrate the inferior pole of the kidney under sonographic localization and guidance. Each biopsy session would get 4 core samples to obtain a total of ~10 mg renal specimen. At the end of the procedure, buprenorphine (5~10 µg/kg) was given subcutaneously for pain relief. The collected renal biopsy specimen was put into RNA Stabilization Reagent, RNA later (Qiagen, USA) immediately, from which total messenger RNA was extracted and hybridized on AffymetrixGeneChip® Rhesus Macaque Genome Array. It has been reported that the genome divergence between rhesus and cynomolgus macaque is only 0.34-0.40% [15].

Transcriptome analysis of renal biopsy specimen

A total of 14764 genes were profiled on the AffymetrixGeneChip®
Rhesus Macaque Genome Arrays, of which 11074 genes could be assigned unique gene symbols and were selected for further analysis. Gene centric expression values were obtained using the custom Cumulative Distribution Function (CDF) file Rhesus_MAmu_ENTREZG (version 16.0.0) from Brain array [16]. The background correction was performed using Robust Multichip Average (RMA) algorithm and quantile normalization. The expression values of mRNA were log₂-transformed. The multiple pair wise comparisons were carried out to detect the Differentially Expressed Genes (DEGs) using Significance Analysis of Microarrays (SAM) software. The threshold was set to 4 fold change of mRNA expression between the 2 categories. As illustrated in Figure 1, the below 5 pairs were compared to delineate respective effects of dysmetabolism, diabetes and/or nephropathy contributing to the changes of gene expression in the renal biopsy specimen.

1. N vs. DM for the contribution of diabetic nephropathy;
2. N vs. Pre-DM for the contribution of dysmetabolism without nephropathy;
3. N vs. Pre-DM & DM for the contribution of dysmetabolism with and without nephropathy;
4. Pre-DM vs. DM for the contribution of nephropathy in the presence of dysmetabolism;
5. DM vs. N & Pre-DM for the contribution of nephropathy only.

Gene Set Enrichment Analysis (GSEA) by IPA software (Ingenuity) was used to draw biological interpretation of DEGs including their relationship to biological networks, diseases, and canonical pathways.

Data and statistical analyses

The diabetic nephropathy phenotype data are presented as mean and standard error of the mean. Group means among different groups were compared using ANOVA with Fisher LSD pairwise post hoc comparisons. Statistical significance was set at p<0.05.

Results

Basic classification of the NHP colonies

The monkeys were classified as 3 categories, N, Pre-DM, and DM based mainly on general characterization for the fasting blood glucose, HbA1c, and ivGTT parameters. As shown in Table 1, the normal monkeys were obtained from a young colony with lighter body weight compared to the pre-DM and DM ones, which could only be obtained from an older population. Both the average fasting blood glucose concentrations and HbA1c were similar between the N and pre-DM, but significantly higher in DM monkeys. The calculated glucose clearance rate (Kglnsucrose) via ivGTT was significant lower in the pre-DM and the lowest in DM compared to N.

Characterization of diabetic nephropathy phenotypes in the NHPs

Both the Dvolume (Figure 2A) and Uvolume (Figure 2B) were significantly lower in the pre-DM than both N and DM monkeys with the average Dvolume being ~14% higher than Uvolume measured simultaneously on the same day in all 3 groups. The Dvolume and Uvolume were highly correlated with each other for the individuals (Figure 2D). The Ualbumin was significantly higher in the DM than both N and pre-DM groups (Figure 2C). Since the Uvolume and Ualbumin could be variable in different days, 2 repeated experiments were carried out by collecting 24 hr urine at ~6 weeks apart. The mean values of both the Uvolume (Figure 2B) and Ualbumin (Figure 2C) in each group were not significantly different, and the values from each individual monkey were highly correlated between the 2 separate measurements (Figure 2E and 2F). These data demonstrated that the present method for measurements of total daily drinking and urine volumes as well as urinary albumin excretion were highly reproducible.

The Uglucose (Figure 3A) and Uprotein (Figure 3B), but not Ucreatinine (Figure 3C), were significantly elevated only in the DM monkeys. However, the urinary albumin index ACR was significantly lower in the pre-DM than both N and DM monkeys. The calculated Dvolume and Uvolume were highly correlated with each other for the individuals (Figure 2D). The Dvolume and Uvolume were highly reproducible.

<table>
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<th>FDEG</th>
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<td>Diabetic nephropathy</td>
<td>75</td>
</tr>
<tr>
<td>2</td>
<td>N vs. Pre-DM</td>
<td>Dysmetabolic without nephropathy</td>
<td>66</td>
</tr>
<tr>
<td>3</td>
<td>N vs. Pre-DM &amp; DM</td>
<td>Dysmetabolic with without nephropathy</td>
<td>88</td>
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<tr>
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<td>DM vs. N &amp; Pre-DM</td>
<td>Nephropathy only</td>
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Figure 1: Comparative study design (top); the number of Differentially Expressed Genes (DEGs) detected for each corresponding comparison (middle) and overlap map for the DEGs (bottom).

Table 1: Descriptive characteristics (mean ± standard error).
progressively elevated in the pre-DM and DM compared to N monkeys (Figure 3D). There were no significant changes in the renal functional indexes: BUN (Figure 4A) and Scratinine (Figure 4B) between the N and pre-DM, however, both were slightly but significantly reduced in DM group. The calculated eGFR was not significantly different among the 3 groups (Figure 4C).

The correlation analysis was used to examine the relationship between several parameters. The Ualbumin was highly correlated with both Uprotein (Figure 5A) and ACR (Figure 5B) in all groups. When the FBG was <150 mg/dL in most of the N and pre-DM monkeys, there was a minimal Uglucose (<110 mg, Figure 5D) and ACR (<5mg/mmol) (Figure 5E). However, when the FBG was >150 mg/dL threshold in most of the DM monkeys, it was positively correlated with both the Uglucose (Figure 5D) and ACR (Figure 5E). The Uglucose (Figure 5F), but not eGFR (Figure 5C) was modestly correlated with ACR.

**Histopathology of the kidney**

Histology of the kidneys from the representative DM monkeys with albuminuria was analyzed in comparison with a normal control one (Figure 6). The size of glomeruli was enlarged in the DM (Figures 6D and 6E) compared with N kidney (Figures 6A and 6B). Hemorrhage was frequently observed in the glomerulus of the DM kidney (Figure 6E). Typical vacuolar degeneration of the tubules was shown in the DM kidney (Figure 6F). PAS staining revealed basement membrane thickening of the glomerular epithelial cells, mesangial matrix expansion and glomerulosclerosis in the DM (Figure 6D) in comparison with N kidney (Figure 6A). These pathological characteristics are highly representative of the human diabetic nephropathy phenotype.

**Trascriptome analysis of the renal biopsy specimen**

Among the 11074 genes profiled and analyzed in the N, Pre-DM without albuminuria and DM with albuminuria NHPs selected for kidney biopsy, there are 95 unique DEGs detected with varying up- and down-regulation by a threshold of >4 folds between the

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**Figure 2:** A. 24 hour total urine volume (Uvolume) measured repeatedly in 2 separate experiments; B. the correlation of the Uvolume between the 2 measurements; C. 24 hour total urine albumin excretion (Ualbumin) measured repeatedly in 2 experiments; D. the correlation of the Ualbumin between the 2 measurements; E. 24 hour total urine volume (Uvolume) measured simultaneously with the total drinking volume (Dvolume) in the same day; and F. the correlation between the urine and drinking volume.

**Figure 3:** A. 24 hour total urinary glucose (Uglucose); B. creatinine (Ucreatinine); C. protein (Uprotein) excretion; and D. urinary Albumin/Creatinine Ratio (ACR).

**Figure 4:** Renal functional surrogate markers measured by A. Blood Urea Nitrogen (BUN); B. serum creatinine concentrations; C. Glomerular Filtration Rate (GFR) calculated by creatinine clearance rate.
diabetic nephropathy DEGs (N&DM pair) and 66 dysmetabolic without albuminuria DEGs (N& Pre-DM pair), there are 46 DEGs overlapping, thus, the 29 of 75 DEGs are unique only to N& DM comparison, which may be more likely related to advanced disease stages including diabetes and nephropathy, while 3 of the 4 nephropathy DEGs (LCT, MMP7, and SPP1) found previously are members of the 29 DEGs, and HAVCR1 is 1 of the 46 overlapping common DEGs.

Using GSEA to analyze signaling pathways for the above 4 DEG encoded proteins, together with other 31 related proteins and protein complexes; it constructs a kidney failure, as well as renal, urological and inflammatory diseases related network (Figure 8). The most pivotal gene in this network is Tumor Necrosis Factor (TNF), which is a group of apoptosis-inducing cytokines, indicating that nephropathy is a disease closely related to inflammation and cell death.

**Discussion**

**Characterization of diabetic nephropathy phenotypes in NHPs**

Similar to early stages of diabetic nephropathy patients, the present study observed for the first time in cynomolgus monkeys that there is a progressive albuminuria and elevated ACR in those with severe diabetes, accompanied by aglycosuria, polydipsia and polyuria, with no significant changes in the Ucreatinine and Screatinin, BUN and eGFR, indicating that these monkeys may have impaired or leaky tubules but the kidney function is still largely preserved.

The great correlation between the Ualbumin and Uprotein in the present study indicates that these 2 parameters are parallel, both of which are the consequence of the leaky tubules allowing filtration of large molecules. However, as in many clinical reports, the Ualbumin is an earlier and more sensitive marker for tubular injury, which proceeds the detection of other Uprotein, thus has been commonly used in...

The present data further demonstrated that in the DM monkeys, the level of hyperglycemia is correlated with the severity of glucosuria (Figure 5D) and ACR (Figure 5E), indicating that hyperglycemia could contribute to the pathogenesis of nephropathy. There is no correlation between eGFR and ACR (Figure 5C) because these monkeys are still in the early stage of diabetic nephropathy with the renal function not significantly compromised yet. The increased Uvolume is caused by the osmotic effect of glucosuria in the DM compared to age-matched Pre-DM group. The relatively high Uvolume in the N monkeys is probably due to the much greater physical activities in the young population that is not compatible with the Pre-DM and DM population whose age is much older with much less physical activities, which may account for the less demand for drinking water, thus, less Uvolume in the Pre-DM group.

Transcriptome analysis of the gene expressions of the kidney biopsy specimen

The transcriptome analysis for the kidney biopsy tissues from 3 distinct groups in the present study detected 95 DEGs (Table S1), in which, at least 16 genes are found to be related to kidney diseases or diabetes (Figure 7). As showed in Table 2, specifically, HAVCR1 (KIM1) is involved in acute tubular necrosis and kidney injury, adriamycin-induced nephropathy, mesangial proliferative glomerulonephritis, polycystic kidney disease, anti-thymocyte serum nephritis, chromophobe renal cancer, protein overload nephropathy, renal ischemia, glomerulonephritis, proteinuria, etc. [27,28]. SPP1 is involved in acute tubular necrosis, acute kidney injury, polycystic kidney disease, systemic lupus erythematosus with renal impairment, proteinuria, etc. [28,29]. MMP7 is involved in hydropnephrosis [30]. ACE2 is related to the thickness of basement membrane, which is a typical change in diabetic nephropathy patients, and SLC01A2, CA3, SERPINA3 and TPM1 are related to nephropathy [31]. SLC01A2 is associated with proximal tubular toxicity in male rat [32]. CA3 is associated with hypercalciuria, and an inhibitor of human CA3 protein (Chlorthalidone) has been approved for the treatment of hypercalciuria [33]. SERPINA3 in kidney is associated with acute renal allograft rejection in human [27]. ATP1A2, IFI1H, TNFRSF11B, ADIPOQ, SPP1, CA3, CCL2, SERPINA3, and CAPSL are involved in diabetes mellitus [34]. Clustering the mRNA expression values of these 16 DEGs separates diabetic nephropathy and normal monkeys perfectly, while Pre-DM monkeys spread in between (Figure 9), which is probably due to the much greater physical activities in the young population that is not compatible with the Pre-DM and DM population whose age is much older with much less physical activities, which may account for the less demand for drinking water, thus, less Uvolume in the Pre-DM group.

The progressive nature of albuminuria is usually considered the result of escalating capillary endothelial dysfunction in the glomerular tuft while the basement membrane and underlying epithelial cells also play important roles [23]. The earliest detectable histopathological change in the course of diabetic nephropathy is a glomerular hypertrophy (thickening in the glomerular basement membrane and capillary walls), and mild mesangial matrix expansion as shown in the present study in diabetic nephropathy cynomolgus as well as reported by Cusumano et al. in diabetic rhesus monkeys [24]. At this stage, the leaky tubules may enable the large amounts of serum albumin traverse the glomerular endothelium and associated structures, thus filtrate into the urine. But the 2 separate functions, impaired tubular uptake and compromised degradation may also independently prevent filtered albumin from reabsorption, thus, causing micro- and macro-albuminuria [25]. The compromised degradation may be the dysfunctional pathway for micro-albuminuria before the global GFR is significantly reduced [17]. Instead, hyperglycemia may even result in kidney hemodynamic changes, leading to glomerular hyper-filtration in early stage of diabetes, which is consistent with the current observation that a slight, but not significant increase in eGFR, which in turn leading to a slight, but significant lower in both BUN and Scratinine in the DM monkeys (Figure 4) [26]. As diabetic nephropathy progresses, increasing numbers of glomeruli are damaged with progressive nodular glomerulosclerosis (formation of nodules in the glomerular tuft and angiopathy of capillaries in the glomeruli), increase in mesangial cellularity and matrix, etc., which generally showed in a clinical kidney biopsy as well as demonstrated in both rhesus and cynomolgus monkeys with diabetic nephropathy in the present study [24].

The transcriptome analysis for the kidney biopsy tissues from 3 distinct groups in the present study detected 95 DEGs (Table S1), in which, at least 16 genes are found to be related to kidney diseases or diabetes (Figure 7). As showed in Table 2, specifically, HAVCR1 (KIM1) is involved in acute tubular necrosis and kidney injury, adriamycin-induced nephropathy, mesangial proliferative glomerulonephritis, polycystic kidney disease, anti-thymocyte serum nephritis, chromophobe renal cancer, protein overload nephropathy, renal ischemia, glomerulonephritis, proteinuria, etc. [27,28]. SPP1 is involved in acute tubular necrosis, acute kidney injury, polycystic kidney disease, systemic lupus erythematosus with renal impairment, proteinuria, etc. [28,29]. MMP7 is involved in hydropnephrosis [30]. ACE2 is related to the thickness of basement membrane, which is a typical change in diabetic nephropathy patients, and SLC01A2, CA3, SERPINA3 and TPM1 are related to nephropathy [31]. SLC01A2 is associated with proximal tubular toxicity in male rat [32]. CA3 is associated with hypercalciuria, and an inhibitor of human CA3 protein (Chlorthalidone) has been approved for the treatment of hypercalciuria [33]. SERPINA3 in kidney is associated with acute renal allograft rejection in human [27]. ATP1A2, IFI1H, TNFRSF11B, ADIPOQ, SPP1, CA3, CCL2, SERPINA3, and CAPSL are involved in diabetes mellitus [34]. Clustering the mRNA expression values of these 16 DEGs separates diabetic nephropathy and normal monkeys perfectly, while Pre-DM monkeys spread in between (Figure 9), which is probably due to Pre-DM having different disease severity, spreading from the early to more advanced stages close to, develop nephropathy. Since the Pre-DM and DM are in different stages in the progress of dysmetabolism...
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and diabetes, the common genes related to the dysmetabolism are likely involved in the disease progression.

The 5 top canonical pathways significantly enriched in the 95 diabetic DEGs are the signaling pathways for calcium (ATP2A1, MYH4, MYL2, TNNC2, TNN1 and TPM1), epithelial adherents junction (ACTN2, ACTN3, MYH4 and MYL1), ILK (ACTN2, ACTN3, MYH4 and MYL2), VEGF (ACTN2, ACTN3, and EPIFAY) and Hepatic Fibrosis/Hepatic Stellate Cell Activation (CCL2, MYH4, MYL2, and TNFRSF11B). Among the biomarkers for progression and regression of diabetic nephropathy, only HAVCR1 (KIM-1) and SPP1 (Osteopontin) have shown significantly changed expression in diabetic nephropathy by our test criteria. A similar study of transcriptome analysis was reported in human diabetic kidney disease with the pathway analysis that in the glomeruli the regulation of R as homolog gene family member A, Cdc42, integrin, integrin-linked kinase, and VEGF signaling was highlighted, while in the tubule interstitial compartment, inflammation-related pathways was strong enriched [35]. The canonical complement signaling pathway was determined to be statistically differentially regulated in both glomeruli and tubuli and was associated with increased glomerulosclerosis in patients with diabetic kidney disease.

In summary, the present study demonstrated in the first time that spontaneously occurring dysmetabolic and diabetic cynomolgus monkeys developed similar diabetic nephropathy phenotypes as in human patients, such as hyperglycemia, polydipsia, polyuria, glucosuria, albuminuria, proteinuria, etc., however, with eGFR still remaining normal in the current colony. The histopathological examination also detected tubular injury without overlay glomerular destruction, which is consistent with that in patients with early stage of diabetic nephropathy. Transcriptome analysis of the kidney biopsy specimens revealed that the DEGs amongst N, pre-DM and DM monkeys are highly related to kidney diseases and/or diabetes with signaling pathways constructing a kidney failure, renal, urological, and inflammatory diseases related network, in which the most pivotal gene is Tumor Necrosis Factor (TNF), indicating that nephropathy is a disease closely related to inflammation and cell death. Therefore, the present study demonstrated that NHP model of diabetic nephropathy had the same features as that in diabetic nephropathy patients with similar molecular, cellular, biochemical and pathophysiology mechanisms, thus, can be used as the most clinically translatable model for testing novel therapeutics for both the efficacy and potential adverse responses as well as identifying novel gene signatures for mechanism based diagnostic and therapeutic targets.

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3. FLD, BDW, XLW and YPF performed the experiment. XLW also performed data analysis and manuscript writing.
4. SG and WBQ performed the transcriptome data analysis and manuscript writing.
5. YXC and XC performed histopathology analysis.
6. FMG, MS, KC, YFX and YXC involved in manuscript writing and discussion.
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