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Excessive dietary lipid intake provokes an acquired form of lysosomal lipid storage disease in the kidney

Running title: Diet-induced renal phospholipidosis and associated damage

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Conflict of interest

The authors declare that they have no conflict of interest.
ABSTRACT

Obesity and dyslipidaemia are features of the metabolic syndrome and risk factors for chronic kidney disease. The cellular mechanisms connecting metabolic syndrome with chronic kidney disease onset and progression remain largely unclear. We show that proximal tubular epithelium is a target site for lipid deposition upon overnutrition with a cholesterol-rich Western-type diet. Affected proximal tubule epithelial cells displayed giant vacuoles of lysosomal or autophagosomal origin, harbouring oxidized lipoproteins and concentric membrane layer structures (multilamellar bodies), reminiscent of lysosomal storage diseases; lipidomic analysis revealed renal deposition of cholesterol and phospholipids, including lysosomal phospholipids. Proteomic profiles of renal multilamellar bodies were distinct from those of epidermis or lung multilamellar bodies and of cytoplasmic lipid droplets. Tubular multilamellar bodies were observed in kidney biopsies of obese hypercholesterolaemic patients, and the concentration of the phospholipidosis marker di-docosahexaenoyl (22:6) bis(monoacylglycerol) phosphate was doubled in urine from individuals with metabolic syndrome and chronic kidney disease. The enrichment of proximal tubule epithelial cells with phospholipids and multilamellar bodies was accompanied by enhanced inflammation, fibrosis, tubular damage markers and higher urinary electrolyte content. Concomitantly to the intralysosomal lipid storage, a renal transcriptional response was initiated to enhance lysosomal degradation and lipid synthesis. In cultured proximal tubule epithelial cells, inhibition of cholesterol efflux transport or oxysterol treatment induced effects very similar to the in vivo situation, such as multilamellar body and phospholipid amassing, and induction of damage, inflammatory, fibrotic and lipogenic molecules. Onset of phospholipidosis onset in proximal tubule epithelial cells is a novel pathological trait in metabolic syndrome-related chronic kidney disease, and emphasizes the importance of healthy lysosomes and nutrition for kidney wellbeing.
Keywords: diet, lipids, kidney, chronic kidney disease, multilamellar bodies, lysosomal storage disease, lysosome, metabolism, inflammation, fibrosis
Introduction

In the current obesity epidemic, increasing attention has been given to the obesity-related diseases, atherosclerosis, diabetes and metabolic syndrome (MetS). Emerging evidence indicates that MetS individuals are prone to chronic kidney diseases (CKD) and end-stage renal diseases (ESRD). A systematic meta-analysis based on 25 general population cohort studies demonstrates that obesity confers a significantly higher risk of kidney disease compared to normal-weight individuals [1]. In addition, two large cohort studies in Japan and in the United States linked an increased risk for ESRD with high body-mass index (BMI) [2, 3] independently of hypertension and diabetes [2-6]. Remarkably, visceral obesity also puts lean subjects at risk of CKD [7, 8]. The association between MetS and CKD becomes particularly relevant in renal transplantation, as MetS appearance adversely affects allograft survival and long-term function [9].

Despite such associations between MetS and CKD, the etiological role of MetS and obesity in CKD development and progression remains unclear, due to the complexity and bidirectionality of the relationship between the two medical conditions [10]. Nevertheless, the description of ectopic fat deposition in kidneys and the strong interrelation between obesity (particularly BMI) and ESRD suggest a role for fat accumulation per se in the initiation of renal dysfunction or in the progression of existing renal disease [11, 12]. Several studies indeed hint at a causal relationship: (a) weight loss ameliorates proteinuria in overweight patients affected by chronic proteinuric nephropathies; (b) the rates of allograft dysfunction and acute rejection episodes are higher for obese living kidney donors; (c) inhibition of 3-hydroxy-3-methylglutaryl-coenzyme A reductase (HMGCR), the rate-limiting enzyme in cholesterol biosynthesis, improves proteinuria and reduces renal function loss; (d) low-carbohydrate and low-fat diets exert beneficial effects on renal function in obese individuals [13-17]. Altogether, these reports strongly indicate a lipid-mediated effect in initiating renal dysfunction or
injury. In this regard, most of the lipotoxic effect of renal lipid overload linked to obesity and MetS and kidney dysfunction has been attributed to renal steatosis through free-fatty acid (FFA) and triglyceride (neutral lipids) accumulation within cytoplasmic lipid droplets (LDs) [12, 18, 19]. Here, we report the accumulation of free cholesterol (FC) and phospholipids (polar lipids) within multilamellar organelles of lysosomal origin (phospholipidosis) inside renal proximal tubular epithelial cells (PTEC) following Western-diet (WD) feeding. A similar excessive storage of complex polar lipids has been disclosed in PTEC in lysosomal storage diseases, caused by defects in lysosomal lipases or lysosomal lipid transport proteins [20].
Material and Methods

Animal studies

Male wild-type C57BL/6 mice (Charles River) were subjected to control diet (4021.84, AB Diets), Western diet (4021.83, AB Diets) or water enriched with 15% (w/v) fructose (Sigma Aldrich) ad libitum for 16 weeks [21]. GFP-LC3 transgenic mice [22, 23] were fed Western diet. Kidneys from Npc1\textsuperscript{nh/\textit{null}}, Gla\textsuperscript{m1Kul}/\textit{Y}, ob/ob mice and 88-week-old mice were used for transmission electron microscopic (TEM) imaging [24-27]. All procedures were approved by the Animal Care and Use Committee of the Academic Medical Center Amsterdam.

Cell culture and isolation of multilamellar bodies (MLBs)

HK2 and IMM-PTEC were cultured in DMEM/F12 supplemented with ITSe, triiodothyronine, hydrocortisone and prostaglandin E1 (Sigma-Aldrich), plus interferon-\(\gamma\) for IMM-PTEC (PROSPEC) [28]. MLSs were isolated by homogenization and sucrose gradient fractionation; MLSs were recovered at the interface between 0.4 M and 0.35 M sucrose. Urine MLSs were purified by ultracentrifugation [29].

Lipidomics and proteomics

Lipids were isolated by chloroform/methanol extraction and quantified by ESI-MS/MS [30-32]. Sphingolipids were butanol-extracted and analysed by LC-MS/MS [33-35]. GC-MS Solution Software was used for data acquisition and processing [36]. Urine di-docosahexaenoyl (22:6) bis(monoacylglycerol) phosphate (di-22:6-BMP) content was determined by HPLC-MS [37, 38]. For LC-MS proteomics, samples were denatured and digested, following by Nanoscale LC separation of tryptic peptides. MS data were acquired in triplicate in ion mobility enabled data independent analysis mode [39] using a Synapt G2-Si instrument (Waters Corporation). ISOQuant was applied for integrated quantitative analysis of data from multiple LC-MS [40, 41]. Reactome, BLAST2GO, GO...
terms and KEGG pathway annotation analyses were used for pathway/gene ontology annotation [42-44].

**Patient study**

Urine samples belong to the TransplantLines cohort (ClinicalTrials.gov Identifier: NCT02811835, University Medical Center Groningen, The Netherlands) [45-48] Control samples were obtained from transplant donors and the disease group was selected on basis of CKD clinical diagnosis and presence of clinical parameters matching MetS World Health Organization criteria (hypertension, BMI, plasma cholesterol and triglycerides). The transmission electron microscopy (TEM) images shown in supplementary material, Figure S5 were selected from screening 69 biopsies of nephrotic syndrome patients from the University Hospital of Nijmegen, The Netherlands; the images of Figure 5E are derived from obese hypercholesterolaemic patient renal biopsies collected at the Academic Medical Center of Amsterdam, The Netherlands.

**Statistics**

Statistics was performed using One-Way ANOVA and Dunnett’s tests; Mann-Whitney U test and Spearman rank correlation. Data are presented as mean and SEM; p < 0.05 was considered significant.
Results

Accumulation of polar lipids/phospholipids and free-cholesterol in murine proximal tubular epithelial cells upon Western-type diet feeding

Feeding mice for 16 weeks a Western-type diet (WD) containing 0.15% cholesterol and providing 43% energy from fat vs. a control diet (CD), in which 11% of energy comes from fat, leads to a prominent cytoplasmic vacuolization of renal tubular epithelial cells (Figure 1A,B). In all kidneys from WD-fed mice, the vacuoles were located in the proximal tubules S2/S3 segments (not in S1) and absent in distal tubular, endothelial or glomerular cells (Figure 1A, and supplementary material, Figure S1A). By staining kidney sections for the lipid droplet marker perilipin-2 [49, 50], we found numerous small LDs along the tubular cell basolateral membrane after WD-feeding; however, the large, more apically located, vacuoles were all negative for perilipin-2 (Figure 1C). Remarkably, these vacuolized tubules were enriched in phospholipids and free-cholesterol, as shown by Nile red and filipin staining, respectively (Figure 1D,E). Accordingly, quantitative lipidomics analysis of renal tissues from mice subjected to WD revealed an increase in several lipids, including free-cholesterol (FC), fatty acids (FA), several phospholipids and sphingolipids (Figure 1F).

As human studies showed that obesity increases the incidence of a distinct obesity-related glomerulopathy (ORG) [51-53], the extent of glomerular lesions (mesangial cell proliferation, mesangial matrix expansion and mesangiolysis) after WD feeding was evaluated. However, no significant alterations were observed in the glomerular compartment between CD and WD groups (supplementary material, Figure S1B). Thus, although kidneys are generally regarded as non-highly active metabolic organs, we show that tubular epithelial cells are sites of profound lipid deposition within large vacuoles during lipid overnutrition.

Features of acquired lysosomal storage disease in kidneys exposed to lipid overloading
Transmission electron microscopy revealed that these tubular vesicles contained lots of concentric thin electron-dense lamellae surrounded by a limiting membrane (Figure 2A). These structures reached a maximum diameter of 8-9 µm and ultrastructurally resembled the multilamellar bodies (MLBs) that we and others found in kidneys from mice with lysosomal storage diseases such as Fabry and Niemann-Pick type C [24, 54] (supplementary material, Figure S2A). In line with studies associating aging with lysosomal impairment [55], aged mice also presented MLBs. As after WD feeding, subjecting mice to high-fat diet (HFD) resulted in MLB formation in PTEC (supplementary material, Figure S2A). In contrast, leptin-deficient (ob/ob) mice on a regular diet and Wt mice given ad libitum water enriched with 15% fructose for 16 weeks have hardly any renal MLBs (supplementary material, Figure S2B, S2C). This implies that the massive expansion of MLBs in PTEC is a specific reaction to excessive dietary lipid intake, and not generally to any kind of overnutrition.

Next, we discovered that lysosomal and autophagy markers, LIMP-2, CD63, p62 and LC3 [56], were present in the vacuole delimiting zone in WD-PTEC (Figure 2A,B). To directly monitor autophagy, GFP-LC3 transgenic mice were fed a WD for 16 weeks; we found that both LIMP-2 and GFP-LC3 co-localized at the vacuole limiting membrane and Western diet dramatically increased renal GFP-LC3 puncta formation (Figure 2C). This suggests a lysosomal/autophagic origin (autolysosomes) [57] of renal diet-induced MLBs, similarly to what is described in lysosomal storage diseases [20].

We next used laser-capture microdissection (LCM) technology to obtain a histologically pure cell population enriched in vacuolized PTEC for RNA transcript profiling. The expression of lysosome- and lipid metabolism-associated genes was enhanced in response to high-fat/cholesterol feeding in vacuolized proximal tubules (Figure 2D). Indeed, the transcription of genes encoding LIMP-2 and the master regulator of lysosomal function transcription factor EB (TFEB) showed a trend towards increased expression upon overnutrition (p = 0.057) and the gene expression of the lysosomal
enzymes acid ceramidase Asah1 (N-acylsphingosine amidohydrolase 1) and acid sphingomyelinase Spmd1 (sphingomyelin phosphodiesterase 1) was significantly upregulated in WD-vacuolized tubules [58, 59]. Furthermore, the lipogenic transcription factors sterol regulatory element-binding protein 2 and 1c (Srebp2, Srebp1c) and their target genes Ldlr (low-density lipoprotein receptor) and Fas (fatty acid synthase), respectively, were strongly induced in lipid-enriched tubules [60, 61] (Figure 2D).

Altogether, our findings disclose the importance of tubular lysosomes in the renal response to WD feeding and the acquisition of a “lysosomal storage disease phenotype” in PTEC upon diet-derived lipid overload.

Finally, we found that renal MLB formation is likely to be driven by lipoprotein uptake, since WD-induced renal vacuoles harbour lipoproteins, including oxidized lipoproteins, and LDLR (Figure 2E).

**Renal and urinary MLBs have a unique proteomic footprint**

As MLBs from other organs are known to be released into the extracellular space [54], we wondered whether MLBs could be secreted by PTEC. MLBs can be detected by TEM in urine of mice fed a WD (Figure 3A). In addition, immortalized proximal tubular epithelial cells (IMM-PTEC) secreted extracellular vesicles in a lysosomal storage disease-like state, as proven by the detection of the flotillin-2 extracellular vesicle marker [62] in the supernatant of IMM-PTEC treated with U18666A, an inhibitor of cholesterol trafficking out of lysosomes, largely used to reproduce in vitro the NPC1 disease [63-65] (Figure 3B).

To better characterize MLBs, proteomic analysis was performed on MLBs isolated from kidneys and urine of WD-fed mice or U18666A-stimulated IMM-PTEC (supplementary material, Figure S3A-C). We identified 526 proteins (supplementary material, Excel spreadsheet S1); Figure 3C shows the most abundant proteins detected in kidneys, urine and IMM-PTEC upon metabolic overloading. Many of these proteins have been identified as located within extracellular exosomes, and some, such as
GGT1 (gamma-glutamyltransferase 1), ATPA1 (ATPase, Na+/K+ transporting, alpha 1 polypeptide), and ALDOB (aldolase B, fructose-bisphosphate) have high (though not exclusive) renal expression, (www.ncbi.nlm.nih.gov/gene). In addition, kidney MLBs express proteins involved in lipid cellular uptake and trafficking, such as megalin [66, 67], sterol carrier protein 2 (SCP2) [68, 69], Hsp90 [70], legumain (LGMN) [71], Na⁺/H⁺ exchange regulatory cofactor 3 (NHRF3) [72], and fatty acid-binding proteins (FABPs), albumin and apolipoprotein E [73].

Proteomics also revealed the presence of lysosomal proteins and proteins normally located in other organelles and cellular compartments, such as cytoskeleton, mitochondria and plasma membrane (Figure 3D and supplementary material, Figure S4). Reactome pathway-based analysis show that metabolism-related pathways are among the most enriched pathways (supplementary material, Figure S3D, highlighted in yellow) associated with the MLB proteome. This is in line with the known lysosomal functions in cellular clearance and nutrient/energy sensing [74, 75] and the above described data indicating that lysosome-derived MLBs are part of a metabolic cellular adaptation to lipid overload.

It is to be noted that we used a common organelle isolation procedure based on gradient-density centrifugation, as employed by others for MLB isolation [76, 77]. Although this method is not completely contamination-free, it is the only one suitable for organelle isolation from tissues, in contrast to isolation from cell lines, which can be engineered to stably express a tagged organelle-specific marker. Comparison of the renal MLB proteome with published data from proteomics of lysosome-related organelles (LRO), lung/epidermis-derived MLBs and lipid droplets from enterocytes, adipocytes or hepatocytes [76-81] revealed that renal MLBs have a unique protein profile, as shown by frequency/hierarchical clustering analysis (Figure 4). Indeed, most proteins were identified solely in renal MLBs, but not in lung/skin MLBs or LDs (supplementary material, Excel spreadsheet S2 and
Table S1), indicating that they are distinct organelles. However, comparing GO annotation and pathways analyses of all seven sets of proteomes revealed that most of the detected proteins share similarities in general function such as catalytic activity and binding and participation in metabolic processes (supplementary material, Figure S5).

**Oxysterols and impaired cholesterol trafficking cause MLB formation, disruption of lysosome homeostasis and induction of cholesterogenesis genes**

By means of *in vitro* assays with IMM-PTEC, we demonstrate that the same multilamellar structures observed *in vivo* upon nutrient overload appear in PTECs after exposure to U18666A, oxidized LDL (oxLDL) and 7-ketocholesterol (7KC), which is the major oxidation product of cholesterol and among the most common oxysterols found in food [82] (Figure 5A). Accordingly, IMM-PTEC treatment with U18666A, oxLDL or 7KC elevates the intralysosomal phospholipid content and the expression of LIMP-2, as well as their colocalization (Figure 5B). Furthermore, oxysterols increase expression of both LIMP-2 and the autophagy adaptor p62 [56, 83] (Fig S2D). Altogether, the *in vitro* data validate the lysosomal origin of the MLB/vacuole organelles seen *in vivo* and suggest an etiological role of cholesterol overload in the disruption of lysosomal homeostasis.

Since the maintenance of a proper acidic pH is fundamental for the digestion of lysosomal cargo, including lipids [84], we studied the lysosomal acidification rate and found that U18666A, oxLDL and 7KC all increased lysosomal pH. Chloroquine was used as a positive control to inhibit lysosome acidification [85] (Figure 5C).

Lastly, tubular phospholipidosis was accompanied by internalization of cell surface free cholesterol and an increase in intracellular bismonoacylglycerolphosphate (BMP), an atypical phospholipid specific for the (endo)lysosomal compartment involved in intracellular cholesterol transport [86] (Figure 5D). Besides, BMP expression overlapped with the sites of phospholipidosis in U18666A-
treated human PTECs (Figure 5E). Finally, the gene expression of Srebp2, Ldlr and Hmgcr, the rate-controlling enzyme in cholesterol synthesis [60], were all significantly upregulated by U18666A in IMM-PTEC, whereas the gene expression of the lysosomal phospholipase A2 was downregulated (Figure 5F).

Altogether, these data indicate that metabolic overloading of PTECs disrupts lysosomal homeostasis and reprograms cells towards the onset of anabolic pathways.

**Diet-mediated lipid overload of tubular cells instigates tubular damage, inflammation and collagen deposition in mice, and promotes formation of lipid-rich MLBs in human TECs**

To expand our view of the effects of overfeeding on kidney health and function, changes in damage, inflammatory and fibrotic parameters in kidneys were examined. Strikingly, after high-fat/cholesterol feeding, the expression of the sodium-glucose co-transporter 2 (SGLT2) at the apical membrane of proximal tubules almost vanished and concomitantly the proximal tubule injury marker KIM-1 (kidney injury molecule-1) [87] was induced in PTECs. The kidneys of WD-fed mice also show macrophage infiltration and collagen deposition (Figure 6A). The gene expression of Sglt2 was significantly diminished in WD kidneys, whereas the transcription of genes encoding MCP-1 (monocyte chemoattractant protein-1), TGF-β (transforming growth factor-β) and CTGF (connective tissue growth factor) was augmented (Figure 6B). Accordingly, *in vitro* stimulation of IMM-PTEC with U18666A or 7KC provided similar results, with a reduced gene expression of the transporters Sglt2 and Atp1a1 (ATPase Na+/K+ transporting subunit α 1) and an increased expression of Mcp1, Kim1, Tgfβ1, Ctgf (Figure 6C). In addition, WD feeding caused higher urinary concentrations of electrolytes that are normally reabsorbed from the tubular lumen after glomerular filtration (Figure 6D). These data strongly indicate that lipid-rich diets can provoke kidney disease as tubular injury and
malfunction, inflammation and collagen deposition are all key features of renal dysfunction and pathology [88].

To translate our in vivo and vitro findings to human obesity-related nephropathology, kidney biopsies and urine samples from obese and lean individuals were used. In the presence of obesity and hypercholesterolaemia, human PTEC display LIMP-2-positive vacuoles, BMP accumulation and MLBs (Figure 6E). In addition, patients affected by nephrotic syndrome, characterized by elevated serum levels of total cholesterol, LDL cholesterol, and lipoprotein(a) [89], present MLBs in renal tubules (supplementary material, Figure S6A), emphasizing the potential role of hyperlipidaemia in kidney MLB genesis. MLBs were also present at the brush border of tubular cells (supplementary material, Figure S6B) and in the tubular lumen (supplementary material, Figure S6C), which together with the detection of MLBs in urine (Figure 3A), strongly suggest that MLBs are excreted by TEC into the urinary space. Interestingly, we also found that the urinary concentration of di-22:6-BMP, a reliable marker for tissue-phospholipidosis [90], is much higher in the urine of a small cohort of individuals affected by CKD and MetS (Figure 6F) and shows a moderate positive correlation with the total albumin urinary content (Figure 6G). These findings imply that renal phospholipidosis also takes place in humans with metabolic/lipid disorders.
Discussion

Although a number of investigators have outlined the “renal lipotoxicity” hypothesis, the role of lipids *per se* as promoters of renal injury remains largely elusive and hard to prove, particularly in humans [91]. Altogether, our data indicate that excessive dietary lipid intake (and not overnutrition *per se*) cause the development of an “acquired lysosomal storage disease” in the kidney epithelium. Cholesterol/fat-rich Western diet induces large vacuoles enriched in free cholesterol, fatty acids and phospholipids and a robust MLB biogenesis in S2/S3 proximal tubules. Intratubular MLBs display lysosomal or autophagolysosomal origin and express the lysosome-bound BMP glycerolipid, resembling the features of lysosomal storage disease and oxLDL-loaded macrophages [54, 86]. Besides this, we detected MLBs in urine and found a striking excretion of the lysosomal di-22:6-BMP in urine of individuals affected by MetS and CKD. The lack of vacuoles, thus MLBs, in PTEC of S1 segment can be explained by their well-developed endo-lysosomal apparatus [92] and greater lysosomal volume and enzymatic activity compared to S2/S3 PTEC [93-96]. Although we cannot completely exclude MLB leakage from damaged TEC into urine, TEC-mediated MLB secretion seems more likely, in light of the known MLB secretion in other organs [54] and of our TEM-based observations of intact MLBs within TEC brush border. Accordingly, urine and kidney MLBs contain identical proteins, many of which are reported to be present in exosomes, and some of the MLB proteins (i.e. GGT1, ATPA1, ALDOB) with the highest relative abundance are also highly expressed in kidneys. In addition, the identification of proteins known to participate in lipid uptake and lipid intracellular trafficking within kidney-derived MLB strengthen our hypothesis of cholesterol/lipid-driven MLB formation, e.g. megalin (receptor for albumin-FA/LDL), SCP2 (intracellular cholesterol movement), Hsp90 (stabilization of NCP1); LGMN (lysosomal protein degradation), NHRF3 (maintenance of expression and function of the HDL receptor SR-BI),
apolipoprotein E (major cholesterol carrier), albumin and FABP suggest their uptake and incorporation in MLBs [66-73]. Thus, renal MLBs appear to be active organelles involved in storage of excessive lipids and secretion as urinary products; in kidneys, MLBs seems to act as a defence mechanism against lipid nephrotoxicity by sequestering and excreting surplus lipids.

Treatment of IMM-PTEC with 7KC hydroxysterol, oxidized-LDL and cholesterol trafficking inhibitor U18666A provoked tubular alterations very similar to the in vivo tubular changes, suggesting an etiological role of oxLDL/oxysterols in the obesity-related renal disease. Indeed, WD-induced tubular vacuoles contained LDLR and oxidized lipoproteins, which are more abundant in obesity [97].

Apolipoprotein E, FABP, albumin and megalin were detected in MLBs by proteomics; thus several lipotoxic molecules may promote lysosomal lipid storage development in our model, besides oxysterols. Besides oxysterols and lipoproteins, other lipotoxic molecules may account for lysosomal dysfunction and lipid-rich vacuole generation in PTEC. In agreement, Kuwahara et al. attributed the appearance of cytosolic vacuoles in PTEC following HFD-feeding to megalin-mediated FA endocytosis [67]. Nevertheless, autophagy is very likely implicated in “renal lysosomal storage disease” as we showed with GFP-LC3 WD-fed mice; consistent with this, autophagic flux contributes to the physiological production of MLBs in lung type II alveolar cells [98]. Although autophagy is regarded as a protective mechanism that contributes to lipid catabolism by delivering LD content to lysosomes (lipophagy) [99, 100], the rise in lysosomal pH observed in our in vitro model may block lipophagy through inhibition of pH-sensitive lysosomal hydrolases.

Interestingly and controversially, WD feeding initiated a transcriptional response in lipid-enriched vacuolated tubules, characterized by the up-regulation of genes accountable for lysosome biogenesis and function and concomitantly of genes responsible for cholesterol and FA synthesis. The enhanced gene expression of Tfeb itself together with Asah1 attests to TFEB activation and nuclear relocation,
which leads to enhance autophagic and lysosomal lipolytic processes [58, 59]. This regulation is probably an adaption of cells to prevent lipotoxicity; indeed, overexpression of TFEB in liver protects against diet-induced obesity and MetS [101].

The increased expression of SREBP-1/2 by dietary lipid challenge is likely to sustain intratubular lipid accumulation during persistent overnutrition, since they upregulate genes for FA and cholesterol synthesis, respectively [60]. Upon lipid overload, phospholipids and cholesterol remain engulfed in lysosomes; therefore, less cholesterol is likely to be transported to the endoplasmic reticulum, where SREBPs reside and are suppressed by the cholesterol content, as cholesterol-induced suppression of SREBP activation vanishes and SREBP target genes are upregulated [60, 102]. Accordingly, we showed that inhibition of cholesterol transport with U18666A in TEC resulted in increased transcription of \textit{Srebp2} and of its target genes \textit{Hmgcr} and \textit{Ldlr} [60, 61, 102].

The increased urinary electrolytes and the appearance of damage, inflammatory and fibrotic markers in WD-kidneys and in U18666A/7KC-treated IMM-PTEC provide an important causal link between kidney pathology and overnutrition. In fact, even though several epidemiological studies have proven a significant association of kidney dysfunction with obesity, dyslipidaemia and high BMI [2, 6, 103, 104], a direct detrimental impact of overnutrition and excessive metabolic loading of kidney cells on kidney health and function has been difficult to prove and is not yet established by scientific evidence.

Bis(monoacylglycerol)phosphate (BMP) is an atypical phospholipid localized mainly at the inner membrane of lysosomes, where its negative charge (due to the acidic pH) helps adherence of positively charged polycationic enzymes and activator proteins, thereby promoting hydrolysis/degradation [105]. Importantly for a translational aspect, the tissue phospholipidosis marker di-22:6-BMP [90, 106] was increased in urine specimens of individuals affected by CKD and
MetS, and urinary BMP levels positively correlated with albuminuria rates within the group of patients with obesity, dyslipidemia and CKD. Furthermore, BMP was shown to colocalize with intracellular phospholipidosis and vacuolization sites and lysosomes in murine/human PTEC. Future studies with larger cohorts would aid assessment of the potential of urinary di-22:6-BMP as a prognostic marker for MetS/obesity-mediated renal disease.

Inflammation and fibrosis are hallmarks of progressive renal disease [107]. In kidneys, MCP-1 was found to be increased as early as one week after initiation of high fat diet feeding [108]. This together with our in vitro findings (48h-treatment with U18666A, oxLDL, or 7-KC) suggests that proximal TEC can produce this chemokine at a very early stage upon metabolic challenge, possibly contributing to the macrophage infiltration observed in WD-fed mice. The induction of fibrotic markers in the kidney in the context of hypercaloric diet was previously reported [108-111]. Interestingly, the upregulation of SREBPs has a role in kidney fibrosis development, as shown by studies with SREBP-1c-deficient mice and SREBP-1a transgenic mice [12, 112]. Importantly, a recent report highlights the role of lysosomes and autophagy in renal lipotoxicity [113], while another reports enlarged vacuoles and multilamellar inclusions within PTEC following high fat diet [110]. We show that a high-cholesterol WD provokes equal pathological alterations in proximal tubules.

Among the lipids found accumulated in kidneys after WD feeding, the sphingolipid ceramide and its metabolites, such as ceramide-1-phosphate (C1P) and sphingosine, warrant particular attention, due to their pro-inflammatory properties, which may favour renal inflammation [114, 115].

Altogether, our study shows that phospholipidosis in kidneys is a novel pathological pathway in obesity-related kidney disease and brings attention to the lysosomal compartment of renal tubules as potential target site for intervention/preventive approaches against MetS-induced nephropathy. It provides a rationale for screening individuals with obesity or metabolic syndrome, for kidney...
dysfunction and phospholipidosis, e.g. with proton magnetic resonance spectroscopy [116], to identify early-stage CKD patients, who can still profit from changes in diet, lifestyle and lipid-lowering therapies.
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Author contributions

Elena Rampanelli and Peter Ochodnicky contributed to the design of the research, performed experiments, analyzed and interpreted data and wrote the paper. Johannes P.C. Vissers and Lee A. Gethings performed LC-MS proteomics analysis, Gerhard Liebisch performed MS lipidomics analysis. Alessia Calcagni, Lotte Kors and Gwen J. Teske conducted in vivo experiments. Loes M. Butter and Nike Claessen conducted laboratory work. Marius A. van den Bergh Weerman and Jan Aten conducted sample preparation and TEM imaging. Dave Speijer helped isolating multilamellar bodies. Bettina Jung helped in sample collection. Stephan J.L. Bakker, Martin H. de Borst, and Gerjan J. Navis collected and provided human urine samples and clinical data. Eric Steenbergen provided renal biopsies for TEM imaging and the corresponding patient clinical diagnosis. Gerd Schmitz contributed with background knowledge on lysosomal storage disease, phospholipidosis and lipid droplets. Johannes M.F.G. Aerts, Andrea Ballabio, Sandrine Florquin, Jan Aten, Stephan JL Bakker, Martin H de Borst, and Gerjan J Navis helped in interpreting and discussing data and reviewed the manuscript. Jaklien C. Leemans led the research project and contributed to the conception and design of the work. All authors discussed the results and participated in drafting or revising the article.
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Figure legends

Figure 1. Accumulation of polar lipids (phospholipids) and free cholesterol in murine proximal tubular epithelial cells upon Western-type diet feeding. (A) PAS-D-stained renal sections of mice fed a control diet (CD) or Western-type diet (WD), 20 × magnification. (B) Quantification of tubular vacuolization in PAS-D-stained kidney sections of CD- and WD-fed mice: number of vacuolized tubules per high-power field (HPF, 20 ×). (C) Perilipin-2 staining in CD and WD kidneys sections (20 × magnification, 100 × magnification far-right image). (D) Nile red and (E) Filipin staining of renal sections for detection of phospholipids and free cholesterol, respectively, in kidneys of CD- and WD-fed mice (D and E: 20 × and 40 × magnification, respectively). (F) Lipidomics analysis (ESI-MS/MS) of renal lipid content: free-cholesterol (FC), cholesterol-esters (CE), fatty acids (FA), bismonoacylglycerophosphate (BMP), phosphatidyglycerol (PG), lysophosphatidyglycerol (LPG), phosphatidic acid (PA), cardiolipin (CL), sphingomyelin (SPM), ceramide, ceramide-1-phosphate (Cer-1-P), lactosylceramide (LacCer), glucosylceramide (GluCer), sphingosine, sphinganine, phosphatidylcholine (PC), lysophosphatidylcholine (LPC), phosphatidylethanolamine (PE), phosphatidylinositol (PI), phosphatidylserine (PS), plasmagens. (B, F) Data shown as mean ± s.e.m.; *P<0.05, ***P<0.001.

Figure 2. Features of acquired lysosomal storage disease in kidneys exposed to metabolic overloading. (A) Transmission electron microscopy (TEM) images of control diet (CD) and Western-type diet (WD) murine kidneys (upper panel): L, luminal space, MLB, multilamellar bodies, N, nucleus, scale bar 10/2 μm. Staining for LIMP-2 and CD63 in CD and WD kidneys sections (20 × magnification, 40 × magnification far-right column). (B) Images of renal sections of WD-fed mice stained for p62 (40 × magnification; asterisk= positive vacuolar membrane) and of GFP-LC3/LIMP-2 double-stained kidney sections from GFP-LC3 transgenic mice upon WD feeding (100 × magnification; LIMP-2 = red;
GFP = blue, asterisk = co-localization, purple). (C) Counts of GFP-LC3 puncta per proximal tubule in CD and WD kidneys. (D) mRNA expression of genes related to lysosome function (left graph) and lipogenic pathways (right graph) in laser-dissected vacuolar tubules and respective controls from mice fed a WD and CD, respectively. Values normalized to control. (E) Positive staining for LDLR, oxidized lipoproteins (oxLP) and lipoproteins (LP) in kidneys of WD-fed mice (20 × magnification). Data shown as mean ± s.e.m.; *P<0.05, **P<0.01, ***P<0.001.

Figure 3. Proteomics analysis of MLBs isolated from murine kidneys and urine upon metabolic overloading. (A) TEM analysis of urine of mice on Western-type diet (WD); scale bars 10/2 µm. (B) Western blot for detection of exosome marker Flotillin-2 in total cell lysates and cell-derived supernatant in control and U18666A-treated immortalized murine tubular epithelial cells (IMM-PTEC). (C) Most abundant proteins found in MLBs isolated from both kidney tissue and urine of WD-fed mice and U18666A-treated IMM-PTEC. Proteins shown after selecting the 50 proteins with the highest ppm (relative LC-MS abundance) independently in the protein list derived from proteomic analysis of kidney, urine and IMMP-TEC MLBs. (D) Graph showing the frequency of protein origin among all intracellular organelles and extracellular exosomes. The proteins considered for analysis are the most abundant proteins shown in (C).

Figure 4. Hierarchical cluster and frequency analysis to compare the proteomic profiles of murine kidney MLBs (our data), lysosome-related organelles (LROs) [78], lung MLBs [76], epidermis MLBs [77], lipid droplets (Lds) from enterocytes [79], adipocytes [80], and hepatocytes [81].
**Figure 5.** Interference with cholesterol trafficking and oxysterol treatment are responsible for MLB formation, disruption of lysosome homeostasis and induction of genes regulating cholesterogenesis. (A) TEM images showing the formation of MLBs after in vitro incubation of immortalized murine tubular epithelial cells (IMM-PTEC) with U18666A, oxidized low density lipoproteins (oxLDL), and oxysterol 7-ketocholesterol (7-KC). Asterisks indicate cellular MLBs localization. (B) Fluorescence microscopy (FM) images, in which phospholipidosis and LIMP-2 are detected in control and IMM-PTEC cells stimulated with U18666A/oxLDL/7KC; nuclei = blue, phospholipidosis = red, LIMP-2 = green, co-localization = yellow; 100 × magnification. (C) Increase of lysosomal pH after incubation of IMM-PTEC with U18666A, oxLDL, 7KC, and chloroquine (CQ, positive controls). (D) FM images of control and U18666A-stimulated human PTEC HK2 cells, showing cellular accumulation of phospholipids, cholesterol (filipin staining) and BMP; nuclei = blue, phospholipidosis = red, cholesterol = blue, BMP = green; 10 × magnification. (E) FM images of U18666A-stimulated HK2 cells, showing phospholipidosis and BMP distribution; nuclei = blue, phospholipidosis = red, BMP = green, co-localization = yellow; 100 × magnification. (F) Gene expression of the transcription factor Srebp2, its target genes Ldlr and Hmgcr, and lysosomal phospholipase A2 (Lpla2) in control and U18666A-stimulated IMM-PTEC. Values normalized to control. Data shown as mean ± s.e.m.; *P<0.05, **P<0.01, ***P<0.001.

**Figure 6.** Diet-mediated lipid overload of tubular cells associates with an inflammatory milieu, collagen deposition, and reduced specialized transporter expression. (A) Immunostaining of renal sections from control diet (CD)- and Western-type diet (WD)-fed mice for visualization of the apical SGLT2 transporter, tubular damage marker KIM-1, macrophage infiltration, and collagen type III deposition; 20 × magnification. (B, C) Quantitative gene expression analysis for membrane

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transporters (SGLT2, ATP1A1), chemokine MCP-1, damage marker KIM-1, fibrogenic markers TGF-β and CTGF in (B) murine renal tissues and (C) IMM-PTEC. Values normalized to control. (D) Electrolyte concentration (sodium, potassium and chloride) in urine of mice fed CD or WD; values normalized for urinary creatinine (nmol/mmol). (E) Representative images of renal biopsy from obese hypercholesterolaemic patients. Conventional staining with acidophilic dye toluidine blue (left upper image) and immunostaining for LIMP-2 revealing the presence of vacuoles within proximal tubular cells and the expression of LIMP-2 at the vacuolar membrane, respectively; 20 × magnification. Immunofluorescence for visualization of the lysosomal glycerophospholipid BMP within TEC; 100 × magnification. TEM images of human tubular cells harbouring MLBs (indicated by asterisks), scale bar 5 and 1 µm. (F) Quantitative lipidomics analysis: di-docosahexaenoyl (22:6) bis(monoacylglycerol) phosphate (di-22:6-BMP) content in human urine specimens; n = 21. (G) Spearman correlation between total albumin and di-22:6-BMP urinary concentrations; human urine specimens; n = 21. Data shown as mean ± s.e.m.; *P<0.05, **P<0.01, ***P<0.001.
SUPPLEMENTARY MATERIAL ONLINE

Supplementary materials and methods  YES

Supplementary figure legends  YES

Figure S1. PAS-D stained renal sections of wild-type mice fed a control diet (CD) or Western-type diet (WD)

Figure S2. Kidney specimens from various mouse models of acquired lysosomal storage disease, obesity and aging to compare MLB formation

Figure S3. Isolation of MLBs by sucrose gradient-centrifugation and proteomics data analysis

Figure S4. MLB proteins clustered by function and location

Figure S5. GO terms and KEGG pathway annotation of 7 distinct proteomics datasets

Figure S6. Transmission electron microscopy (TEM) images of kidney biopsies from patients with nephrotic syndrome

Table S1. Comparison among proteomes of kidney/lung/epidermis-derived MLBs, lysosome-related organelles (LROs), and lipid droplets (LDs) from enterocytes, adipocytes and hepatocytes

Table S2. Primer sequences

Excel spreadsheet file S1.

Excel spreadsheet file S2.
A. TEM images of control diet, Western diet, and Western diet with Western diet treatments. N = nucleus, L = lumen.

B. LAMP-2 and CDS1 expression levels in control diet and Western diet groups.

C. Tubular autophagy levels in control diet and Western diet groups.

D. Graph showing fold-change in expression levels of various proteins between control diet (CD) and Western diet (WD) groups.

E. LDLR and SERP1 expression levels in control diet and Western diet groups.