Renal UTX-PHGDH-serine axis regulates metabolic disorders in the kidney and liver

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Global obesity epidemics impacts human health and causes obesity-related illnesses, including the obesity-related kidney and liver diseases. UTX, a histone H3K27 demethylase, plays important roles in development and differentiation. Here we show that kidney-specific knockout Utx inhibits high-fat diet induced lipid accumulation in the kidney and liver via upregulating circulating serine levels. Mechanistically, UTX recruits E3 ligase RNF114 to ubiquitinate phosphoglycerate dehydrogenase, the rate limiting enzyme for de novo serine synthesis, at Lys310 and Lys330, which leads to its degradation, and thus suppresses renal and circulating serine levels. Consistently, phosphoglycerate dehydrogenase and serine levels are markedly downregulated in human subjects with diabetic kidney disease or obesity-related renal dysfunction. Notably, oral administration of serine ameliorates high-fat diet induced fatty liver and renal dysfunction, suggesting a potential approach against obesity related metabolic disorders. Together, our results reveal a metabolic homeostasis regulation mediated by a renal UTX-PHGDH-serine axis.
Obesity, which results from an imbalance between caloric intake and energy expenditure, is a severe public health problem, affecting near 2 billion population worldwide. Obesity is associated with increased risk of chronic kidney diseases, which eventually progress to end-stage renal disease (ESRD). Compared with normal weight subjects, the hazard ratios of ESRD risk are 1.87 and 3.57 for those with overweight or obesity.

Abnormalities, including glomerulomegaly, mesangial expansion, tubular atrophy, and renal interstitial fibrosis, have been observed in individuals with obesity. Moreover, ectopic lipid accumulation is found in the kidney, especially in the tubular cells, of human and mouse with obesity. Therefore, similar to accumulation is found in the kidney, especially in the tubular hematopoiesis. A crosstalk between kidney and white adipose tissue has been revealed in a mouse model of chronic kidney disease (CKD), focal segmental glomerulosclerosis, adriamycin-induced nephropathy and Alport syndrome. Moreover, increased absorption of albumin-bound fatty acids by proximal tubular cells leads to tubular insulin resistance and enhanced renal gluconeogenesis, eventually causes tubular atrophy. However, how fatty kidney develops remains not fully understood.

The most well-known function of kidney is to maintain acid-base balance and electrolyte homeostasis, and to remove waste products; however, it also plays important endocrine roles by releasing a number of important factors, such as renin, erythropoietin, prostaglandin and aldosterone, thus involved in regulating bone growth, cardiovascular system, and hematopoiesis. A crosstalk between kidney and white adipose tissue has been revealed in a mouse model of chronic kidney disease (CKD), where renal lesion increases the circulating levels of parathyroid hormone and parathyroid hormone-related protein, which promotes their binding to the receptor PTHR in the white adipose tissue, and stimulates browning. However, connections between kidney and other metabolic organs remain not fully understood.

UTX (ubiquitously transcribed tetratricopeptide repeat on chromosome X, also known as KDM6A), a H3K27 demethylase, plays roles in various biological processes, such as controlling cell fate, embryonic development, and cancers. Recently, upregulation of UTX has been found in the kidneys of patients with diabetic kidney disease or focal segmental glomerulosclerosis, as well as in the kidneys of mouse models of type 1 diabetes (Akita and OVE mouse) and type 2 diabetes (db/db mouse), indicating an important role of UTX in the development of diabetic kidney disease. However, the roles of renal UTX in obesity-related diseases remain unclear.

Here, we report that kidney-specific knockout Utx reduces renal and hepatic steatosis under HFD stress. Notably, renal UTX recruits E3 ligase RNF114, which leads to PHGDH degradation and suppression of serine synthesis. Consistently, oral serine treatment ameliorated HFD-induced obesity-related kidney disease and NAFLD. Together, our results reveal that a renal UTX–PHGDH–serine axis regulates metabolic homeostasis in the kidney and liver.

**Results**

**UTX is upregulated in obesity-related kidney disease.** Previous studies by others and us have reported increased renal UTX expression in patients with diabetic kidney disease or focal segmental glomerulosclerosis, as well as in the kidneys of diabetic mouse models including Akita mice, db/db mice, OVE26 mice, and STZ-treated mice. Herein, we investigated the role of UTX in obesity-related kidney disease. In human tissues, compared to the controls, patients with obesity showed dramatically upregulated UTX in renal tubular and glomerular cells (Fig. 1a and Supplementary Table 1). Consistently, upregulated UTX was observed in the kidney of HFD-fed mouse, further co-immunofluorescence staining demonstrated elevated UTX in the nuclei of proximal tubular cells, distal tubules, mesangial cells and podocytes (Fig. 1b).

Kidney-specific Utx knockout mice show a lean phenotype under HFD stress. To study the role of UTX in the kidney, we generated two strains of kidney-specific Utx knockout mice. Significantly reduced mRNA and protein levels of UTX, as well as increased H3K27me3 level in whole kidney or renal medulla and cortex, but not in the liver and brown adipose tissue (BAT) (Fig. 1c–e, Supplementary Fig. 1a–f), were found in both UtxPax2 KO and UtxKsp KO mice, indicating kidney-specific knockout of Utx. The Utx KO mice were generally healthy and normally born, with no obvious behavioral abnormality.

To study the role of renal UTX in obesity related kidney diseases, UtxPax2 KO and UtxKsp KO mice, as well as their respective WT littermates, were challenged with HFD as indicated (Supplementary Fig. 1g). Under HFD stress, decreased UTX level was observed in the tubular cells of UtxKsp KO mice (Fig. 1f). Surprisingly, under HFD feeding, compared with WT mice, both UtxKsp KO and UtxPax2 KO mice showed significantly lower body weights, reduced fat, increased lean mass, as well as reduced weights of liver, inguinal white adipose tissue (iWAT) and brown adipose tissue (BAT) (Fig. 1g–i, Supplementary Figs. 1h and 2a–c, Supplementary Tables 2 and 3), even though UtxKsp KO mice showed slightly higher amount of food intake at some time points (Supplementary Fig. 1f). Compared to HFD-fed WT mice, HFD-fed UtxPax2 KO mice also showed significantly lower kidney weights (Supplementary Table 3).

Furthermore, consistent with their lean phenotypes under the HFD stress, UtxKsp KO or UtxPax2 KO mice showed significantly decreased serum triglyceride (TG), cholesterol, leptin, insulin, and non-fasting blood glucose (NFBG) levels (Fig. 1j–l and Supplementary Fig. 2d–g), as well as significantly improved glucose tolerance and insulin tolerance (Fig. 1m, n and Supplementary Figs. 1j, k and 2h, i). Meanwhile, the levels of uBUN (urine blood urea nitrogen) and uAUCR (urine albumin/creatinine ratio), which are commonly used to monitor the development and extent of renal damage, were significantly elevated in HFD-fed WT mice. UtxKsp KO mouse showed a trend of inhibition of uAUCR level under HFD-fed conditions, with similar eGFR (estimated glomerular filtration rate) level in these groups (Supplementary Fig. 1f–n). Interestingly, liver or adipose tissue specific knockout Utx in male mice showed no obvious effect on HFD-induced obesity (Supplementary Fig. 2j, k).

Kidney-specific Utx knockout ameliorates obesity-induced renal morphological disorders. In a murine obesity-related kidney disease model, HFD feeding induced glomerular expansion, mesangial matrix expansion and lipid accumulation in the kidney. Compared with respective HFD-fed WT mice, both HFD-fed UtxKsp KO and UtxPax2 KO mice showed significantly decreased glomerular hypertrophy and lipid droplet formation in renal tubules (demonstrated by H&E staining), while decreased mesangial expansion was detected in HFD-fed UtxKsp KO mice (demonstrated by PASH staining), indicating less morphological injury in the kidney under HFD stress (Fig. 2a, b and Supplementary Fig. 3a, b). Furthermore, significantly decreased levels of TG and cholesterol were found in both medulla and cortex, as well as significantly downregulated transcription levels of genes...
involved in TG synthesis and lipid storage, such as Mogat1, Gpam, Cidea and Cidec, in HFD-fed Utx KO or Utx KO mice, compared to their respective controls (Fig. 2c–f and Supplementary Fig. 3c). Downregulation of Cidea and Cidec by knockout Utx in kidney was found in NC-fed mice (Fig. 2e, f and Supplementary Fig. 3c). UTX has been reported to promote gene transcription through removing H3K27me3 from their promoters. Increased H3K27me3 levels on the promoters of Cidea and Cidec were found upon; knockout as demonstrated by ChIP assay (Supplementary Fig. 3d). Moreover, the transcription levels of genes involved in lipid synthesis, lipid transport, β-oxidation, glyconeogenesis and glycolysis pathways, as well as protein levels of several key factors regulating these pathways were examined in mouse kidney. Under HFD stress, Utx KO
resulted in decreased Fasn, Glut4, G6pc and Pc mRNA levels, with no obvious effect on other genes or proteins we examined (Supplementary Fig. 4).

Kidney-specific knockout of Utx reduces hepatic steatosis and adipocyte size under HFD stress. Anatomic analysis indicated significantly reduced liver weight in HFD-fed UtxKsp KO or UtxPax2 KO mice (Supplementary Tables 2 and 3). Furthermore, ameliorated hepatic steatosis was found in dissected liver, and by H&E and Oil Red O staining, in HFD-fed UtxKsp KO mice (Fig. 3a–c and Supplementary Fig. 5a). Furthermore, reduced hepatic TG and cholesterol levels, as well as downregulated TG synthesis and storage related genes, such as Mogat1, Cidea, Cidec, and Gpam were also observed in the liver of HFD-fed UtxKsp KO mice (Fig. 3d, e). Meanwhile, compared with HFD-fed WT mice, HFD-fed UtxKsp KO or UtxPax2 KO mice showed significantly reduced adipocyte size in BAT, iWAT and eWAT (Fig. 4a–c and Supplementary Fig. 5b–d). Significantly upregulated thermogenic genes, including Ucp1, Elovl6, Dio2, Caco5b and Prdm16, were found in iWAT and BAT of HFD-fed UtxKsp KO mice (Fig. 4d, e).

Renal UTX negatively regulates PHGDH and serine levels under HFD stress. By analyzing human kidney datasets for chronic kidney disease (CKD; GSE66494) or diabetic kidney disease (DKD; GSE96804), we examined gene alterations involved in de novo serine synthesis and UTX (UTX) by using RNAseq. As the rate limiting enzyme for de novo serine synthesis, the transcription level of phosphoglycerate dehydrogenase (PHGDH) was downregulated in the kidneys of patients with DKD or CKD (Fig. 5a, b). Consistently, immunohistochemical staining demonstrated decreased PHGDH expression in the kidneys of subjects with obesity or DKD (Fig. 5c and Supplementary Tables 1, 4). Next, eGFR, as well as serum levels of serine and glycine, which are respectively the product of de novo serine synthesis and serine transformation process via one-carbon metabolism, were measured in the individuals with normal body weight (n = 39), overweight (n = 23), obesity (n = 24) and obesity-related renal dysfunction (n = 12) (Supplementary Table 5). Downregulated eGFR was found in subjects with overweight and obesity, and was further downregulated in subjects with obesity-related renal dysfunction (Fig. 5d). Significantly lower serum serine level was found in the subjects with obesity-related renal dysfunction (Fig. 5e). Serum serine level was negatively correlated with CREA and BUN levels, and was positively correlated with eGFR in these clinical samples (Fig. 5f, g, Supplementary Fig. 6, and Supplementary Table 5). Similar glycine levels were found in these groups, and a weak negative correlation with cholesterol was observed (Supplementary Fig. 7 and Supplementary Table 5).

Next, we studied whether knockout of Utx in whole kidney or renal epithelial cells regulates serine and PHGDH levels. Downregulated PHGDH level was found in the kidney of HFD-fed WT mice, while its protein levels, but not mRNA levels, were significantly higher in HFD-fed UtxKsp KO or UtxPax2 KO mice (Fig. 5i, j and Supplementary Fig. 8). In HFD-fed WT mice, decreased serine and glycine levels in renal medulla, serum and urine were found; whereas in HFD-fed UtxKsp KO mice, normal renal medulla and serum serine/glycine levels, as well as partly rescued urine glycine level, were observed (Fig. 5k–n). On the other hand, liver or adipose tissue specific knockout Utx in male mice showed no effect on PHGDH level, and no effect on circulating serine level was observed in liver specific Utx knockout mice (Supplementary Fig. 9). Consistent with in vivo results, reduced lipid accumulation, as demonstrated by Oil red O staining and TG measurement, were found in UTX knockout or knockdown HK-2 cells under palmitic acid (PA) treatment-induced stress that mimics in vivo hyperlipidemia (Fig. 6a, b and Supplementary Fig. 10a–c). Furthermore, knockout or knockdown of UTX in HK-2 cells upregulated PHGDH under NC- and PA-stressed conditions (Fig. 6c and Supplementary Fig. 10d); whereas overexpression of UTX increased lipid accumulation, downregulated PHGDH and upregulated transcription levels of genes involved in TG synthesis/lipid storage upon PA stress (Fig. 6d–g). To explore whether the histone demethylase activity of UTX affects the transcription, plasmids expressing wildtype or a demethylase activity abolished UTX mutant were transfected in UTX knockout HK-2 cells. UTX knockout significantly decreased the mRNA levels of CIDEA and CIDEC, which were restored by overexpression of wildtype UTX but not the mutant (Supplementary Fig. 10e). Furthermore, an in vitro luciferase reporter assay confirmed the binding and regulation of UTX on CIDEA and CIDEC (Supplementary Fig. 10f).
Since knockout or knockdown of UTX upregulated PHGDH level in HK-2 cells, we generated PHGDH knockout cells (Supplementary Fig. 10g). In PA-treated HK-2 cells, knockout of PHGDH increased lipid accumulation, transcription levels of genes involved in TG synthesis/storage and decreased intracellular serine/glycine levels (Fig. 6h–k). Consistent with PHGDH knockout results, in PA-treated HK-2 cells, PHGDH overexpression reduced lipid accumulation and transcription levels of genes involved in TG synthesis/storage (Supplementary Fig. 11).

To further investigate whether UTX-deficiency prevents PA-induced lipid accumulation via upregulating PHGDH, a stable UTX knockout and a UTX/PHGDH double knockout HK-2 cell (DKO) were generated (Supplementary Fig. 10h). Under PA stress, compared to UTX KO per se, DKO cells showed increased TG level and upregulated transcription levels of genes involved in TG synthesis/lipid storage (Fig. 7a, b). While knockout UTX normalized PA-induced reduction of intracellular serine and glycine levels, this normalization was abolished in DKO cells (Fig. 7c). Similar results were found for the medium serine level,
but not glycine level, in these groups (Fig. 7d). Next, we administrated NCT-503, a PHGDH inhibitor24, to PA-treated UTX KO HK-2 cells. Significantly upregulated TG levels, transcriptional levels of genes involved in TG synthesis/lipid storage, as well as downregulated cellular/medium serine level and cellular glycine level, were observed (Fig. 7e–h). A control study indicated the specificity of NCT-503 on the enzymatic activity of PHGDH (Supplementary Fig. 12).

UTX negatively regulates PHGDH by promoting ubiquitination at K310 and K330. Ubiquitin-proteasome and autophagy mediated proteolysis are two major eukaryotic protein degradation pathways25. To investigate how UTX negatively regulates PHGDH protein level in renal cells, we first determined the major PHGDH degradation pathway affected by UTX. Inhibitors for autophagosome (3-methyladenine, 3-MA), lysosome (chloroquine, CQ) and proteasome (MG132) were administrated to the control or UTX overexpressing HK-2 cells, only MG132 abolished the negative regulation of UTX on PHGDH levels (Fig. 8a), suggesting UTX proteasome-dependently mediates PHGDH degradation.

To identify possible binding partners of UTX, immunoprecipitation (IP) was performed in 293 T cells transfected with UTX, following mass spectrometromic analysis suggested that RNF114, an E3 ligase, may bind to UTX (Supplementary Fig. 13a). Under...
overexpression conditions, co-IP experiments demonstrated binding of UTX with RNF114, and PHGDH binds to the UTX/ RNF114 complex (Fig. 8b–d and Supplementary Fig. 14a–c). Endogenous co-IP experiments indicated that PHGDH was weakly associated with UTX and RNF114 at the normal conditions, and their association was dramatically increased after PA stress, partly due to upregulated UTX and RNF114 expression (Fig. 8e). Increased RNF114 level was also found in the kidney of HFD-fed mice (Supplementary Fig. 13b). Knockout of UTX abolished the binding between endogenous PHGDH and RNF114 (Fig. 8f); moreover, overexpression of either UTX or RNF114 promoted the degradation of PHGDH in HK-2 cells (Fig. 8g). Since UTX upregulates the transcription of CIDEA and CIDEC (Supplementary Figs. 3c, d and 10e, f), while UTX is the only component in the PHGDH/UTX/RNF114 complex with known direct effect on transcription, we explored whether PHGDH or RNF114 affects gene transcription by regulating UTX protein level. However, no change in UTX level was observed in PHGDH KO or RNF114 KO HK-2 cells (Supplementary Fig. 14d).

To address whether UTX regulates PHGDH stability by ubiquitination, we performed ubiquitination assays, and overexpression of UTX markedly increased ubiquitination of PHGDH (Fig. 8i). We next screened and identified that the K48-linked polyubiquitin chain was removed from PHGDH by the UTX/RNF114 complex (Fig. 8j). To determine the ubiquitination site(s), based on bioinformatic prediction (https://www.genecards.org) and a recent mass-spectrometry study,24 we determined six lysines (K146, K289, K310, K330, K364, and K389) as ubiquitination sites, based on bioinformatic prediction (https://www.genecards.org) and a recent mass-spectrometry study. Therefore, we performed ubiquitination assays, and overexpression of UTX markedly increased ubiquitination of PHGDH (Fig. 8i). We next screened and identified that the K48-linked polyubiquitin chain was removed from PHGDH by the UTX/RNF114 complex (Fig. 8j). To determine the ubiquitination site(s), based on bioinformatic prediction (https://www.genecards.org) and a recent mass-spectrometry study,24 we determined six lysines (K146, K289, K310, K330, K364, and K389) as ubiquitination sites, based on bioinformatic prediction (https://www.genecards.org) and a recent mass-spectrometry study.
K384) as potential ubiquitination site(s). Six respective lysine to arginine PHGDH mutants were constructed. Compared to the wildtype, K310R and K330R mutations dramatically abolished UTX-promoted PHGDH ubiquitination (Fig. 8k), and markedly restored PHGDH level (Fig. 8l). These results indicate K310 and K330, which respectively reside on the surface of substrate binding domain and regulatory domain (Fig. 8m), are sites for UTX/RNF114 mediated ubiquitination and degradation of PHGDH.

Serine inhibits PA-induced lipid accumulation in cultured cells. We further investigated whether serine, the product catalyzed by PHGDH, regulates lipid accumulation. Serine (200 or
400 μM) significantly downregulated TG accumulation and genes involved in TG synthesis/storage in PA-treated HK-2 cells (Fig. 9a, b). These beneficial effects were significantly inhibited by knockout of serine transporter ASCT1 and/or ASCT2 (Fig. 9c–f). Moreover, serine treatment and PHGDH overexpression increased the NAD+/NADH ratio in PA-treated HK-2 cells (Supplementary Fig. 15).

The attenuated hepatic steatosis found in HFD-fed UtxKsp KO mice (Fig. 3) let us hypothesize that knockout of Utx in renal epithelial cells may result in releasing factors that affect liver lipid metabolism. To test this possibility, culture medium from differently treated HK-2 cells were collected and added to primary mouse hepatocytes (Supplementary Fig. 16a, d). Under PA stress, compared to cells treated with medium from respective control cells, reduced lipid accumulation was found in hepatic cells treated with medium from UTx KO cells; whereas increased lipid accumulation was found in those treated with medium from UTx overexpressing cells (Supplementary Fig. 16b, c, e, and f).

In UTx KO (Fig. 2d), a larger amount of serine was released to the medium (Fig. 7d), we next tested whether exogenous serine accumulation was found in those treated with medium from UTx KO HK-2 cells; whereas increased control cells, reduced lipid accumulation was found in hepatic cells treated with medium from UTx KO cells; whereas increased lipid accumulation was found in those treated with medium from UTx overexpressing cells (Supplementary Fig. 16b, c, e, and f).

Serine prevents HFD-induced renal dysfunction and hepatic steatosis. Next, by supplementing serine in drinking water for 18 weeks (Fig. 10a), we investigated whether serine affects lipid accumulation in the kidney and liver of HFD-fed mice. At 8 weeks after HFD feeding, similar glucose tolerance and insulin tolerance were found in mice with or without serine treatment (Supplementary Fig. 19a, b). At 18 weeks after HFD feeding, increased serum serine and glycine levels, as well as reduced serum TG and cholesterol levels, were found after serine treatment (Fig. 10b, c). Decreased glomerular hypertrophy and lipid droplet formation in renal tubules (demonstrated by H&E staining), increased renal serine and glycine levels, decreased renal TG and cholesterol levels, downregulated genes involved in TG synthesis/storage, and similar insulin signaling related p-AKT level were found in the kidney of serine-treated HFD-fed mice (Fig. 10d–h, and Supplementary Fig. 19c, d). Similarly, decreased liver weights, hepatic steatosis (demonstrated by H&E staining and scoring), increased hepatic serine level, decreased hepatic TG and cholesterol levels, downregulated genes involved in TG synthesis/storage, as well as unchanged insulin signaling pathway, were found in the liver of serine-treated HFD-fed mice (Fig. 10i–m, Supplementary Fig. 19e, f and Supplementary Table 6). Whereas serine treatment showed no obvious effect on adipose tissue weights and adipose hypertrophy (Supplementary Table 6 and Supplementary Fig. 20).

**Discussion**

In metabolic diseases, multiple organs, as well as their releasing factors, are simultaneously affected to generate an altered signaling network. Elucidating such tissue crosstalk is vital for understanding and preventing/treating diseases. Kidney, besides its well-known urinary function, also releases several circulating hormones, indicating its endocrinal role. Kidney also releases some amino acids into the renal venous, for example, it is a major organ for serine synthesis. Here, we demonstrated that renal synthesized serine, which is regulated by UTx, affects body metabolic homeostasis, especially the liver.

Increased UTx has been found in multiple renal cell types in diabetic patients and murine models by others and us. This phenomenon was also observed in individuals with obesity and HFD-fed mice (Fig. 1a, b). Therefore, we first generated whole kidney Utx knockout mice by crossing Utxfl/fl with Pax2-cre. To further explore the role of UTx in renal tubular cells, we generated the renal tubular specific Utx knockout mouse by crossing Utxfl/fl with Ksp-cre, which has previously been reported to also express in renal proximal tubular cells. Since the proximal tubular cells are highly involved in lipid and glucose metabolism in the kidney, therefore it will be interesting as a future study to use a renal proximal tubular specific Cre, such as SGLT2-cre, to generate a proximal tubular specific Utx knockout mouse, and compare its phenotype with that of UtxKsp/Pax2 KO mice.

Our study suggested that UtxKsp/Pax2 KO activated serine biosynthesis by promoting PHGDH level in the kidney. Moreover, renal Utx knockout not only ameliorated the HFD-induced kidney steatosis, but also prevented HFD-induced obesity and NAFLD due to increasing circulating serine level; renal-released serine prevented hepatocytes and renal tubular cells from lipid accumulation upon HFD stress (Figs. 1–3, 5, and 10 and Supplementary Figs. 1–3, 5). In contrast, liver or adipose tissue...
Fig. 6 UTX and PHGDH regulated lipid level in cultured renal cells. a–c UTX knockout prevented PA-induced lipid accumulation in HK-2 cells. Oil Red O staining (a), TG level (b) and PHGDH level (c) in UTX knockout HK-2 cells with or without PA stress. b n = 3 biological samples (mean ± SD), " P (Ctrl+PA vs Ctrl+CT) < 0.0001; " P (Ctrl+PA vs UTX KO+PA) < 0.0001 (one-way ANOVA); c n = 2 biological samples per group. d–g UTX overexpression exacerbated PA-induced lipid accumulation in HK-2 cells. Oil Red O staining (d), TG level (e), PHGDH level (f) and transcriptional levels of TG synthesis/lipid storage related genes (g) in UTX overexpressing HK-2 cells. e, g n = 3 biological samples (mean ± SD), " P (Vec+CT vs Vec+PA) = 0.0063, " P (Vec+PA vs Ctrl+PA) = 0.0119; " P (MOGAT1(Vec+CT vs Vec+PA) = 0.0038, " P (CIDEA(Vec+PA vs Vec+CT)) = 0.0009; " P (CIDEA(Vec+PA vs Vec+CT)) = 0.0009, " P (CIDEA(Vec+PA vs Ctrl+PA)) = 0.0076 (one-way ANOVA); f n = 2 biological samples per group. h–k PHGDH knockout aggravated PA-induced lipid accumulation and reduced cellular serine/glycine levels in HK-2 cells. TG level (h), transcriptional levels of TG synthesis/storage related genes (i), Nile Red staining (j) and cellular serine (k, top) or glycine (k, bottom) level in PHGDH knockout HK-2 cells. Scale bar, 50 μm. h–j n = 3 biological samples per group, k n = 6 biological samples per group (mean ± SD). " P (Ctrl+PA vs Ctrl+CT) = 0.0006, " P (Ctrl+PA vs PHGDH KO+PA) < 0.0001, " P (CH1+PA vs CH1+CT) = 0.0034; " P (MOGAT1(CH1+CT vs CH1+PA)) = 0.0351, " P (MOGAT1(CH1+PA vs PHGDH KO1+PA)) = 0.0408; " P (CIDEA(CH1+CT vs CH1+PA)) = 0.0230, " P (CIDEA(CH1+PA vs PHGDH KO2+PA)) = 0.0495, " P (CIDEA(CH1+PA vs PHGDH KO1+PA)) = 0.0289; " P (CIDEA(CH1+CT vs CH1+PA)) = 0.0158; " P (CIDEA(CH1+PA vs PHGDH KO1+PA)) = 0.0092, " P (CIDEA(CH1+PA vs PHGDH KO2+PA)) = 0.0002; " P (Glycine(CH1+CT vs CH1+PA)) < 0.0001, " P (Glycine(CH1+PA vs PHGDH KO1+PA)) = 0.0036. " P (Glycine(CH1+CT vs CH1+PA)) < 0.0001, " P (Glycine(CH1+PA vs PHGDH KO1+PA)) = 0.0101 (one-way ANOVA). a–k at least three independent experiments were performed and similar results were obtained. Source data are provided in the Source Data file.
specific knockout Utx in male mice, showed no effect on circulation serine and PHGDH levels (Supplementary Figs. 2j, k and 9), which may due to the much lower RNF114 level in the liver and adipose tissues compared to that in the kidney (https://www.proteinatlas.org/ENS0000124226-RNF114/tissue). Interestingly, renal knockout Utx also increased intestinal probiotics in HFD-fed mice (Supplementary Fig. 2f). There are some differences observed between UtxKO and UtxPAx2KO mice, such as the mildly increased fat weight in UtxKO mice vs. no obvious change in fat weight in UtxPAx2KO mice under NC conditions and dramatically decreased fat weight in HFD-fed UtxKO vs UtxPAx2KO mice (Fig. 1g, h, Supplementary Figs. 1h and 2a-c, Supplementary Tables 2 and 3). Such difference may be caused by the varied renal cell types knockout of Utx.
using different Cre strains, similar phenomena have been previously observed in other genes knockout mediated by different Cre strains in the same tissue32,33.

Important functions of serine have attracted attention from different angles. Neuroprotective role of serine has been revealed34,35, impaired serine biosynthesis has been found in Alzheimer’s disease (AD)36, therefore, serine is now being investigated as a therapy for neurological disorders including polyneuropathy, amyotrophic lateral sclerosis and AD34,36,37. As a critical one-carbon unit donor involved in methionine cycle and folate cycle which contribute to nucleotide synthesis, cancer cells utilize serine as the major source of one-carbon units to accelerate...
Serine also plays an important role in metabolic diseases. Significantly decreased serum serine level is found in patients with diabetes, CKD or NAFLD, as well as in ob/ob and db/db mice, indicating that serine may play an important role in metabolic diseases. In the present study, decreased serine level was also found in obesity related kidney disease, while renal knockout Utx increased circulating serine level (Figs. 5c and 5m). Whether eGFR affects circulating serine level is unclear. There was no signif i cant difference in eGFR between the WT and UtxKsp KO mice under NC- and HFD-fed conditions (Supplementary Fig. 1n), which indicates no correlation in our experimental setting; whereas a mild negative correlation between eGFR and circulating serine level was suggested in collected clinical samples (Fig. 5h). Therefore, it will be interesting to explore in a larger cohort with different renal disease groups.

Previous studies have indicated lipotoxicity as a major mechanism involved in HFD-induced glomerular hypertrophy. Here, we found that serum treatment inhibited lipid accumulation in the tubular cells which may benefit glomerular hypertrophy (Fig. 10d, e). Meanwhile, glomerular hypertrophy in obesity-related nephropathy is associated with tubule-glomerular feedback induced glomerular hyperfiltration, altered glomerular hypertrophy by serum treatment (Fig. 10c, e) may also be due to tubule-glomerular feedback, similar to the effects of SGLT2 inhibitors, which will be interesting for future investigation.

Additionally, as a rate limiting enzyme of serine synthesis, PHGDH also play important roles in metabolic diseases and tumors. For example, decreased hepatic PHGDH level has been found in the HFD- or MCD-induced steatosis animals, and in patients of alcoholic and non-alcoholic liver diseases, which is associated with lower hepatic serine levels. Furthermore, transgenic mice show significantly reduced hepatic triglyceride accumulation and lipogenic gene levels fed on HFD diet, indicating regulation of hepatic lipid level by PHGDH. However, specific KO of Phgdh mice in adipose tissues show no effect on body weight and glucose intolerance on males, but a mild reduced body weight and glucose intolerance on females upon a high-fat high sucrose diet, indicating tissue specific and gender specific effects of PHGDH.

Although the biological function of UTX in regulating gene transcription has been intensively studied, its function in protein stability is unknown. Accumulating evidence has suggested that histone-modification enzymes may regulate the ubiquitination of target proteins. Here, we report an important role of UTX on the regulation of PHGDH stability, through interacting with a E3 ligase RNF114, the UTX/RNF114 complex ubiquitinates PHGDH at K310 and K330 and promotes its degradation (Fig. 8). Consistently, in vitro and in vivo results all suggested that stabilization of PHGDH and the resulting increase of serine synthesis contribute to the protective effects on renal lipid accumulation after Utx knockout. These results thus reveal an important mechanism underlying the regulatory role of UTX in metabolic diseases. Both our results (Supplementary Fig. S13b) and a single-cell sequencing study suggested upregulated RNF114 under obesity or diabetic conditions in the kidney, therefore, upregulated expression of UTX and RNF114 in the kidney may contribute to the enhanced degradation of PHGDH. Furthermore, enhanced binding between UTX and RNF114 was found under PA stress (Fig. 8e). Since UTX forms a complex with RNF114 that leads to PHGDH degradation, thus enhanced binding may also induce increased PHGDH degradation. However, other factors that trigger the changes in HFD conditions still await further investigation.

In the present study, we showed that kidney-specific knockout UTX affected lipid accumulation in the liver and adipose tissue. Consistently, primary mouse hepatocytes and stromal vascular fraction (SVF) cells cultured with medium from differently treated HK-2 cells suggested that under PA stress, reduced lipid accumulation was found in hepatocytes and SVF cells treated with medium from UTX KO cells (Supplementary Figs. 16, 22). Renal UTX knockout increased PHGDH and circulation serine levels, while serine treatment also regulated lipid accumulation in liver and primary mouse hepatocytes (Figs. 5, 9–10), but not affected in vivo adipogenesis and in vivo adipose hypertrophy under HFD stress (Supplementary Figs. 18, 20). Together, these results suggested that renal knockout UTX increased serine level which attenuated hepatic steatosis. On the other hand, renal knockout UTX also released unknown factor(s) that affect adipose tissue, however, serine was not the one. The effect of renal knockout UTX on adipose tissue warrants future investigations.

In summary, our findings reveal a renal UTX-PHGDH-serine axis regulates metabolic homeostasis in the kidney and liver (Fig. 10n). Dietary serine supplement may be further explored as a preventive/therapeutic approach against high-fat diets induced metabolic disorders.

Methods

Study approval. Human serum samples were collected by Affiliated Wuhan Hospital of Traditional Chinese and Western Medicine of Tongji Medical School (Supplementary Table 5). Normal (CT), overweight or subjects with obesity were defined as BMI < 23 kg/m², 23 ≤ BMI < 27.5 kg/m², and BMI ≥ 27.5 kg/m², respectively as described. Paraffin embedded human renal biopsy samples, including diabetic kidney disease samples or subjects with obesity, were collected by the Affiliated Union Hospital and Hubei Cancer Hospital of Tongji Medical College (Supplementary Tables 1, 4). Collection and use of patient samples and patient characteristics were approved by the Institutional Review Boards of affiliated Wuhan Hospital of Traditional Chinese and Western Medicine, Hubei Cancer Hospital and Union Hospital of Tongji Medical College (approval numbers 2020 16, LHBCH2022015, and 2018)S196, respectively), all subjects included in the study provided written informed consent. Mice were handled according to the Guidelines of the China Animal Welfare Legislation, as approved by the Committee on Ethics in the Care and Use of Laboratory Animals of College of Life Sciences, Wuhan University.
Animals. Mouse strains, including *Utx*<sup>fl</sup> mice (JAX stock No.021926), *Pax2*-Cre mice (Cre activity expressing in glomeruli, renal tubules, collecting ducts, and nephric duct in the kidney<sup>28</sup>) and *Ksp*-Cre mice (*CDH16*-Cre, Cre activity expressing in renal tubules<sup>29</sup>) were used. Male *Pax2*-Cre or *Ksp*-Cre mice were crossed with *Utx*<sup>fl</sup>/+ females to generate whole kidney specific or renal epithelial cell specific *Utx* knockout mice, respectively. Additionally, male *Adiponectin*-Cre (*Adi*-Cre, JAX, stock No.010803) or *Alb*-Cre (JAX, stock No.003574) mice were crossed with *Utx*<sup>fl</sup>/+ females to generate liver or adipose tissue specific *Utx* knockout mice. Genotyping was performed (primers listed in Supplementary Table 7). As an X-linked gene, males genotyped as *Utx*<sup>fl</sup>/y (WT) and *Utx*<sup>fl</sup>/y *Pax2*-Cre (*UtxPax2* KO), *Utx*<sup>fl</sup>/y *Ksp*-Cre (*UtxKsp* KO), *Utx*<sup>fl</sup>/y *Adi*-Cre (*UtxAdi* KO) or *Utx*<sup>fl</sup>/y *Alb*-Cre (*UtxAlb* KO) were used in the present study.

Mice were maintained in a specific-pathogen-free, temperature controlled (22 ± 1 °C) animal facility with a 12-h light/dark cycle, and free access to water/food. Male mice were used in this study, euthanasia was performed by CO2 inhalation. Mice were maintained on normal chow (NC, #1025, Beijing Huafukang, ...
Beijing, China). For diet-induced obesity-related kidney disease, mice were fed with a high-fat diet (60% Kcal fat, #D12492, Research Diets, New Brunswick, NJ) at 4 weeks-old for indicated period of time. For serine treatments, L-serine (Sigma, St. Louis, MO) was supplemented in drinking water (3%, v/v) for indicated period of time.

Biochemical measurements. Serum levels of leptin, insulin, triglyceride and cholesterol were measured using mouse leptin ELISA kit (Millipore, Billerica, MA), rat/mouse insulin ELISA kit (Millipore), triglyceride kit (Jiangcheng, Nanjing, China), and total cholesterol kit (Kehua Biotech, Shanghai, China), respectively.68 Triglyceride and cholesterol levels in the kidney or liver were measured using triglyceride kit (Jiangcheng) and total cholesterol kit (Kehua Biotech), respectively. Glucose tolerance test (GTT) and insulin tolerance test (ITT) were performed69 for GTT, mice were fasted overnight, then intraperitoneally injected with D-glucose (1.5 g/kg body weight; Amresco, Solon, OH). For ITT, mice were fasted for 6 h, then intraperitoneally injected with insulin (0.75 unit/kg body weight; Lilly, France). Blood glucose levels were measured with a OneTouch blood glucose meter (LifeScan, Milpitas, CA) at 0, 15, 30, 60, 90 and 120 min after glucose or insulin injection. Blood composition was scanned with a miniscan LF50 analyzer (Bruker, Rheinstetten, Germany). Twenty-four hour urine samples from WT or UtxKsp KO mice, fed on NC and HFD diets, were collected in metabolic cages (Tecniplast, Italy) 1 day before sacrifice, and the volume was measured. Serum creatinine (CREA) level was analyzed on a Siemens ADVIA 2400 automatic biochemical analyzer using a creatinine reagent kit, while urine creatinine, albumin and BUN were measured with an Olympus AU2700 automatic biochemical analyzer using creatinine reagent kit, albumin reagent kit and urine BUN reagent kit, respectively (all from Fuxing Changzheng Medical, Shanghai, China). Mouse eGFR was calculated by the creatinine clearance with the formula Ccr = CREA (urine)/CREA (serum) × 144/24 (h).64,65 Cellular NAD+ and NADH levels were determined using a NAD+/NADH kit (Beyotime, Suzhou, China).

Histological and immunohistochemical studies. Paraffin embedded kidney, liver and adipose samples were sectioned and stained with hematoxylin & eosin (H&E), analyses for glomerular area, PAS positive stained area and adipocyte size were performed.62

Paraffin-embedded sections were deparaffinized and rehydrated as previously reported63. Sections were incubated with 3% H2O2 for 5 min to quench endogenous peroxidase activity. After blocking with 2% goat serum, primary antibodies including PHGDH (1:2000 dilution, Proteintech, Wuhan, China) was applied for the sections overnight at 4°C. After washing with PBS-T, sections were incubated with biotinylated anti-rabbit (Vector laboratories, Burlingame, CA) for 1 h at room temperature. Positive staining was visualized using DAB substrate following the protocol (Vector laboratories). Sample pictures were taken using an Olympus BX60 microscope with a digital camera.

For co-immunofluorescent staining, sections were incubated overnight with respective primary antibodies (Supplementary Table 8). For renal tubular staining, sections were incubated with 1 μg/mL PNA (peanut agglutinin, detecting distal tubules and collecting ducts), or 1 μg/mL LTL (lotus tetragonolobus lectin, detecting proximal tubules) (all from Vector Laboratories). While, a-SMA or WT1 was used to detect renal mesangial cells or podocytes, respectively. After washing, sections were incubated with respective Alexa Fluor secondary antibody (Thermo Fisher, Waltham, MA). Sections were covered with DAPI and anti-fading medium. Images were taken by a confocal microscope (Leica SP8, Germany).

Cell culture. Human renal tubular cell line HK-2 (GDC0152, from CCTCC, China Center for Type Culture Collection, Wuhan, China) was cultured in DMEM/F12 media (HyClone, Palo Alto, CA) plus 10% FBS. A human hepatocellular carcinoma cell line HepG2 (CL-013000, from Procell Biotech, Wuhan, China) and HEK293T cells (CL-0005, Procell Biotech) were cultured in DMEM media (HyClone) containing 10% FBS. HK-2 cells, HepG2 and HEK293T were analyzed with authenticated STR locus and tested for mycoplasma contamination, by CCTCC or Procell Biotech.

Primary mouse hepatocytes were isolated as described.64 Briefly, mouse liver was perfused with calcium-free solution, digested with collagenase (Sigma), dispersed hepatocytes were collected and plated in collagen-coated plates with DMEM media plus 10% FBS. Plates were washed 4 h later and the adherent hepatocytes were used for further experiments. For medium treatment, medium was freshly collected and filtered from HK-2 cells treated with/without 400 μM palmitic acid (PA), then mixed with equal volume of fresh DMEM medium containing 10% FBS, and applied to 100% confluent primary hepatocytes or HepG2 cells for additional 48 h.

Mouse stromal vascular fraction (SVF) was isolated as reported.65 Briefly, minced iWAT was digested at 37 °C in digestion buffer containing collagenase D and dispase II (Sigma), then filtered. Mature adipocytes and SVF cells were separated by centrifugation. Mouse SVFs were grown in DMEM/F12 media (HyClone) supplemented with 10% FBS. Differentiation of primary white adipocytes was performed as described65,66. Briefly, confluent SVFs (Day 0) were induced by differentiation media (containing 0.5 mM IBMX, 1 μM dexamethasone and 10 μg/mL insulin) for 4 days, then changed to media containing 10 μg/mL insulin until Day 6.

Plasmids, transfections, treatments. Plasmids for human full length UTX (pFLAG-UTX), and shUTX were kind gifts from Dr. Min Gyu Lee (MD Anderson Cancer Center). CRISPR-Cas9 vectors containing gRNA targeting UTX (5'-ACCG AAAAGCGACGGGAGGAGGCGC-3'), PHGDH (5'-ACCGGCTCTAGGGCC-3') were transfected into HK-2 and HEK293T cells.
TTGGTG-3' or 5'-ACCGAAAGCAGAACCTTAGCAAAG-3'), RNF114 (5'-CA
CCGGCCGCTTACACGTGTCCGCA-3'), ASCT1 (5'-CACCGTTGCAGCT
TTCCGTACGGTA-3'), ASCT2 (5'-CACCGCAGCGCCACACCAAAGACGA-3'),
were used to knockout UTX, PHGDH, RNF114 or ASCT1/2, respectively.
Expression plasmids for HA-tagged Ubi (WT, K6, K11, K27, K29, K33, K48, K63,
AKR) and myc-tagged UTX were constructed. HA-tagged RNF114, Flag-tagged
wildtype PHGDH and seven lysine to arginine mutants, pGL3-basic-CIDEA,
pGL3-basic-CIDEC plasmids were constructed with standard procedures (the
primers used are provided in Supplementary Table 9).
Most cells were transfected with indicated plasmids for 6 h, then treated with/
without 400 μM PA for 48 h before collection. Some cells were transfected with
indicated plasmids for 24 h, and treated with different inhibitors (CQ, chloroquine,
0.05 mM; 3-MA, 3-methyladenine, 3.35 mM; MG132, 10 μM) for 6 h. For serine
treatments, normally cultured or PA-stressed HK-2 cells, primary mouse
Fig. 10 Serine suppressed renal and hepatic lipid accumulation in HFD-fed mice. a Experiment design for serine administration via drinking water. b, c Concentrations of serine, glycine, TG and cholesterol in the serum of indicated groups, n = 9, 6 independent animals for HFD or HFD + Ser group, respectively (mean ± SD). d–f, g–i Serine and glycine concentration (μmol/L) and transcriptional levels of TG synthesis/lipid storage related genes (h) in the kidney of indicated groups, data shown as mean ± SD. e n = 9, 5 independent animals for HFD or HFD + Ser group, respectively, “pHSerine = 0.0053; “pTG = 0.0363 (unpaired, two-tailed t-test). d–h H&E staining (d) with quantitative data for relative glomerular area (e), serine and glycine concentration (f), renal TG and cholesterol levels (g), and transcriptional levels of TG synthesis/lipid storage related genes (h) in the kidney of indicated groups, data shown as mean ± SD. n = 9, 6 independent animals for HFD or HFD + Ser group, respectively, “pHSerine = 0.0073; “pTG < 0.0001; “pCholesterol < 0.0001; (h) n = 8, 6 independent animals for HFD or HFD + Ser group, respectively, “pMogat1 = 0.0005; “pCidea = 0.0049; “pCidec = 0.0495 (unpaired, two-tailed t-test). i–m H&E staining (i), with quantitative data for relative liver steatosis score (j), serine and glycine concentration (k), hepatic TG and cholesterol levels (l) and transcriptional levels of TG synthesis/lipid storage related genes (m) in the liver of indicated groups, data shown as mean ± SD. j n = 9, five independent animals HFD or HFD + Ser group, respectively, “pHSerine = 0.197; “pTG = 0.0437; “pCholesterol < 0.0001; (m) n = 8, 6 independent animals for HFD or HFD + Ser group, respectively, “pMogat1 = 0.0008; “pCidea = 0.0495; “pCidec = 0.0072 (unpaired, two-tailed t-test). n Possible mechanisms in present study. Scale bar, 50 μm. Source data are provided in the Source Data file.

Integrated analyzing tool (GEO2R) or R2 fold change of serine metabolism related genes and UTX were visualized.

Homologous modeling. Human PHGDH structure was homologous modeled by the Swiss model (https://swissmodel.expasy.org/), with the crystal structure of Mycobacterium tuberculosis PHGDH (PDB 3DC2) as a reference, and illustrated by PyMol.

16S rDNA sequencing and data analysis for gut flavicolimicrobial. 16S rDNA amplicon sequencing was performed on an Ion S5 XL platform (Thermo Fisher, Waltham, MA). Cluster analysis was performed using unweighted pair-group method with arithmetic means by QiIME (Version 1.7.0), phylogenetic investigation of communities by reconstruction of unobserved states was performed25. Sequencing data is deposited in the NCBI Sequence Read Archive (SRA) database.

Statistics. Data were analyzed with GraphPad Prism (version 8). Statistical analysis was performed using two-tailed Student’s t-test for two experimental groups, and one-way ANOVA for multiple experimental groups without adjustment. Data are reported as the mean values with error bars showing the standard deviation (SD). For cell culture experiments, at least three independent experiments were performed with similar results. For animal study, the number of biological independent animals in different groups are provided in figure legends. A p-value of <0.05 was considered statistically significant.

Data availability. Source data are provided with this paper. Sequencing dataset described in this work is available in NCBI SRA database under accession code SRP375452 (https://www.ncbi.nlm.nih.gov/sra/?term=SRP375452). Microarray datasets reused in this work are from NCBI Gene Expression Omnibus (GSE66494 and GSE96804). Source data are provided with this paper.

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Author contributions

H.C., L.Z., and K.H. designed the study and analyzed the data. H.C., Q.W., C.L., X.Z., M.X., D.Y., Y.X., Y.Z., Y.H., C.L., performed the experiments. J.Y., H.S., and S.W. provided materials. Y.C., H.C., L.Z. and K.H. wrote the manuscript.

Competing interests

The authors declared no competing interests.

Additional information

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