**A catalogue of genetic targets for kidney function from analyses of a million individuals**

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**Abstract**

Chronic kidney disease (CKD) is a public health burden with multi-systemic complications. Through trans-ethnic meta-analysis of genome-wide association studies of estimated glomerular filtration rate (eGFR) and independent replication (n=1,046,070), we identified 264 associated loci (166 novel). Of these, 147 were likely to be relevant for kidney function based on associations with the alternative kidney function marker blood urea nitrogen (n=416,178). Pathway and enrichment analyses, including mouse models with renal phenotypes, support the kidney as the main target organ. A genetic risk score for lower eGFR was associated with clinically diagnosed CKD in 452,264 independent individuals. Co-localization analyses of associations with eGFR among 783,978 European-ancestry individuals and gene expression across 46 human tissues, including tubulo-interstitial and glomerular kidney compartments, identified 17 genes differentially expressed in kidney. Fine-mapping highlighted missense driver variants in 11 genes and kidney-specific regulatory variants. These results provide a comprehensive priority list of molecular targets for translational research.

Chronic kidney disease (CKD) is a major public health issue, with increasing incidence and prevalence worldwide.[1](#_ENREF_1) Its associated burden of disease encompasses metabolic disturbances, end-stage kidney disease, and multi-systemic complications such as cardiovascular disease.[1-4](#_ENREF_1) CKD is a leading cause of death[5](#_ENREF_5) and shows one of the highest increases in disease-attributable mortality over the last decade.[2](#_ENREF_2) Nevertheless, public and clinical awareness remains low.[3](#_ENREF_3) Moreover, clinical trials in nephrology are still underrepresented,[6](#_ENREF_6) resulting in scarce therapeutic options to alter disease progression and high costs for health systems.[7](#_ENREF_7) A major barrier to developing new therapeutics is the limited understanding of the mechanisms underlying kidney function in health and disease, with consequent lack of therapeutic targets.

Genome-wide association studies (GWAS) and exome-chip studies of the glomerular filtration rate estimated from serum creatinine (eGFR), the main biomarker to quantify kidney function and define CKD, have identified nearly one hundred eGFR-associated genetic loci[8](#_ENREF_8) in samples of European[9-15](#_ENREF_9), Asian[16-19](#_ENREF_16), and multiple[20](#_ENREF_20) ancestries. However, similar to other complex traits and diseases, identifying causal genes and molecular mechanisms implicated by genetic associations is challenging and has only been successful for few kidney function-associated loci.[21](#_ENREF_21),[22](#_ENREF_22) Advanced statistical fine-mapping approaches and newly emerging multi-tissue gene expression data provide new opportunities for prioritizing putative causal variants, effector genes, and target tissues from results of large-scale GWAS meta-analyses.

We therefore conducted a trans-ethnic GWAS meta-analysis in the CKD Genetics Consortium (CKDGen, n=765,348), and replicated findings in the Million Veteran Program (MVP, n=280,722),[23](#_ENREF_23) for a combined sample size of >1 million participants. The first aim of this study was to identify novel, globally important loci for kidney function through maximizing statistical power (**Supplementary Figure 1**). Results from GWAS of the complementary kidney function marker blood urea nitrogen (BUN, n=416,178) were used to prioritize eGFR-associated loci most likely to be relevant for kidney function. A genetic risk score (GRS) for low eGFR was used to test relevance for clinically diagnosed CKD among 452,264 independent individuals. The second aim was to characterize replicated eGFR-associated loci through complementary computational approaches, including various enrichment and network analyses, fine-mapping and co-localization with gene expression in 46 tissues and protein levels (**Supplementary Figure 1**). We focused this aim on European ancestry (EA) individuals, as fine-mapping based on summary statistics requires linkage disequilibrium (LD) reference panels whose sample size should scale with that of the GWAS.[24](#_ENREF_24) The resulting list of prioritized variants and genes provides a rich resource of potential therapeutic targets to improve CKD treatment and prevention.

**Results**

*Discovery trans-ethnic meta-analysis*

We performed 121 GWAS encompassing 765,348 individuals of European (n=567,460), East Asian (n=165,726), African American (n=13,842), South Asian (n=13,359), and Hispanic (n=4,961) ancestries (median age: 54 years; 50% females; **Supplementary Table 1**). The median of the study-specific mean eGFR was 89 ml/min/1.73m² (interquartile range, IQR: 81, 94). GWAS were based on genotypes imputed from the Haplotype Reference Consortium[25](#_ENREF_25) or 1000 Genomes Project[26](#_ENREF_26) reference panels (Methods, **Supplementary Table 2**). Following study-specific variant filtering and quality control (QC) procedures, we performed a fixed-effects inverse-variance weighted meta-analysis, showing no evidence of unmodeled population structure (LD Score regression intercept =1.04; λGC=1.05). After variant filtering, 8,221,591 single nucleotide polymorphisms (SNPs) were used for downstream analysis (Methods).

We discovered 308 loci containing at least one eGFR-associated SNP at genome-wide significance (Methods), of which 200 were novel and 108 contained an index SNP reported by previous eGFR GWAS (**Figure 1**; **Supplementary Table 3**). Regional association plots are shown in **Supplementary Figure 2**. The minor alleles across index SNPs showed both decreasing and increasing effects on eGFR, with larger effects observed for lower frequency SNPs (**Figure 1**, inset). The 308 index SNPs explained 7.1% of the eGFR variance, nearly doubling recent GWAS-based estimates,[9](#_ENREF_9) and 19.6% of eGFR genetic heritability (h2=39%, 95% credible interval: 32%, 47%), estimated in a participating general-population-based pedigree-study (**Supplementary Figure 3**;Methods). Index SNPs’ effects were largely homogeneous across studies (**Figure 2A**; **Supplementary Table 3**) and ancestries (**Supplementary Table 4**; **Supplementary Material**).

*Replication and meta-analysis of >1 million individuals*

We assessed replication in an independent trans-ethnic GWAS meta-analysis of eGFR performed among 280,722 MVP participants.[23](#_ENREF_23) Effect estimates, available for 305/308 SNPs, showed almost perfect direction consistency (302/305 SNPs, 99%) and very strong correlation with the discovery results (**Figure 2B**). For these 305 SNPs, we performed a meta-analysis of the 1,046,070 discovery and replication samples. Replication was met by 262 SNPs (**Figure 1**; **Supplementary Table 3**; Methods). Of the three SNPs not available in MVP, the index SNPs at *SHROOM3* (*P*=3.5×10-120) and *SH3YL1* (*P*=1.2×10-11) were also considered replicated based on prior evidence[15](#_ENREF_15),[27](#_ENREF_27), resulting in a total of 264 replicated SNPs (166 novel). Of these, 74 SNPs were genome-wide significant in MVP alone (**Supplementary Table 3**).

*Association of eGFR-associated loci with BUN and CKD*

To evaluate whether associations with creatinine-based eGFR were likely related to kidney function or potentially to creatinine metabolism, we assessed the association of the 264 eGFR-associated index SNPs with BUN, an alternative marker of kidney function that is inversely correlated with eGFR. Trans-ethnic meta-analysis of 65 BUN GWAS (n=416,178; **Supplementary Table 1**) showed no evidence of unmodeled population structure (λGC=1.03; LD Score regression intercept=0.98) and yielded 111 genome-wide significant loci (15 known, 96 novel, **Supplementary Figure 4**, **Supplementary Table 5**).

Of the 264 replicated eGFR index SNPs, 34 and 146 showed genome-wide and nominally (*P*<0.05) significant association with BUN, respectively (**Supplementary Table 6**). SNP effects were inversely correlated (r=-0.65; **Figure 2C**). Their relevance to kidney function was classified as *likely* for 147 eGFR index SNPs with inverse, significant associations with BUN (one-sided *P*<0.05); *inconclusive* for 102 SNPs not associated with BUN (*P*≥0.05); and *unlikely* for 15 eGFR index SNPs showing concordant, significant association with BUN (one-sided *P*<0.05; **Supplementary Table 6**). This comparative analysis of complementary biomarkers supports that signals at the majority of eGFR-associated loci likely reflect kidney function.

Next, we investigated the effects of the eGFR index SNPs on CKD in CKDGen studies (n=625,219, including 64,164 CKD cases; Methods). GWAS meta-analysis of CKD identified 23 genome-wide significant loci, including 17 likely relevant for kidney function (*SDCCAG8*, *LARP4B, DCDC1*, *WDR72*, *UMOD*-*PDILT*, *MYO19*, *AQP4*, *NFATC1*, *PSD4*, *HOXD8*, *NRIP1*, *SHROOM3*, *FGF5*, *SLC34A1*, *DAB2*, *UNCX*, and *PRKAG2;* **Supplementary Table 6**). The majority of replicated eGFR index SNPs (224/264) was associated with CKD (one-sided *P*<0.05; **Figure 1, inset**), including 130 likely relevant for kidney function (**Supplementary Table 6**).

Lastly, we tested if a GRS based on the combined effect of the 147 eGFR index SNPs likely relevant for kidney function was associated with clinically diagnosed CKD and CKD-related outcomes in the UK Biobank (n=452,264; Methods). A lower GRS, reflecting a genetically lower eGFR, was associated with higher odds ratios (ORs) of chronic renal failure, glomerular diseases, acute renal failure, and hypertensive diseases (**Figure 2D; Supplementary Figure 5**). The OR of chronic renal failure per 10% lower GRS-predicted eGFR was 2.13 (95%CI: 1.90, 2.39; *P*=8.1x10-38). A significant protective association with urolithiasis may reflect a reduced ability to concentrate urine at lower eGFR.

*Genetic correlations of eGFR and BUN with other phenotypes*

We assessed genome-wide genetic correlations (rg) of eGFR associations with each of 748 complex traits and diseases (Methods).[28](#_ENREF_28) We observed 37 significant correlations (*P*<6.7×10-5=0.05/748, **Supplementary Figure 6**; **Supplementary Table 7**). After serum creatinine, the largest negative correlations were observed between eGFR and serum citrate (rg=-0.27) and urate (rg=-0.23), followed by anthropometric traits including lean mass and physical fitness (e.g., rg=-0.20 with left hand grip strength). While the inverse correlation with muscle mass-related traits likely reflects higher creatinine generation leading to lower creatinine-based eGFR, the correlations with citrate and urate levels likely reflect reduced filtration function, as does the positive correlation with GFR estimated from cystatin C (rg=0.53).

A very similar pattern of genetic correlations was observed for BUN (**Supplementary Table 7**), but the genetic correlations with muscle mass-related traits were generally lower than those for eGFR. The largest genetic correlations of BUN was observed with CKD (rg=0.47), as compared to creatinine-based (rg=-0.29) and cystatin-based eGFR (rg=-0.26).

In summary, significant genetic correlations with eGFR reflect the two biological components that govern serum creatinine concentrations: its excretion via the kidney and its generation in muscle. The fact that genetic correlations between BUN and muscle-mass related traits are generally lower than those observed for eGFR underscores the value of genetic associations with BUN to help prioritize eGFR-associated loci most likely to be relevant for kidney function.

*Functional enrichment and pathway analyses*

To identify molecular mechanisms and tissues of importance for kidney function, we assessed the enrichment of the eGFR and BUN genetic associations using tissue-specific gene expression, regulatory annotations, and gene sets and pathways (Methods). First, we used eGFR-associated SNPs (*P*<5×10-8) to explore enriched pathways, tissues and cell types based on gene expression data using DEPICT.[29](#_ENREF_29) We identified 16 significantly enriched physiological systems, cell types and tissues highlighting several aspects of kidney function, physiology and disease. The strongest enrichment was observed for urogenital and renal physiological systems and tissues (kidney, kidney cortex, and urinary tract; false discovery rate (FDR) <0.05; **Supplementary Figure 7A** and **B**). Pathway and gene set enrichment analysis identified three highly correlated and strongly associated meta gene-sets (*P*<1×10-6, FDR<0.05), including some relevant to kidney such as polyuria, dilated renal tubules, and expanded mesangial matrix, as well as signaling and transcription, and energy metabolism (**Supplementary Figure 7C**). Tissue and cell-type enrichment analysis of BUN-associated SNPs with *P*<5×10-8 highlighted a very similar pattern (**Supplementary Figure 8**) but without enrichment for muscle tissues, further supporting the use of BUN to prioritize loci most likely to be related to kidney function.

Second, we used stratified LD Score regression[30](#_ENREF_30) on the genome-wide eGFR and BUN summary statistics to identify cell-type groups with enriched heritability based on data from diverse, cell-type specific functional genomic elements. The strongest enrichment for eGFR was observed for kidney (13.2-fold), followed by liver (7.3-fold) and adrenal/pancreas (5.7-fold enrichment; **Supplementary Table 8**). Kidney was also the most enriched cell-type group for BUN (11.5-fold enrichment; **Supplementary Table 8**).

Lastly, using a complementary approach, we assessed enrichment of eGFR-associated variants in genes resulting in kidney phenotypes in genetically manipulated mice.[31](#_ENREF_31) From the Mouse Genome Informatics database, we selected all genes causing abnormal GFR (n=24), abnormal kidney physiology (n=453), or abnormal kidney morphology (n=764), and interrogated their human orthologs in the eGFR summary statistics (Methods). We identified significant associations in 10 genes causing abnormal GFR in mice (enrichment p-value=8.9×10-4), 55 causing abnormal kidney physiology (enrichment p-value=1.1×10-4) and 96 causing abnormal kidney morphology (enrichment p-value=1.8×10-5; **Figure 3**; Methods). Of these, 25 genes represent novel eGFR candidate genes in humans, i.e. they were not previously reported to contain genome-wide significant eGFR-associated SNPs or map near known loci (**Supplementary Table 9**). The existing mouse models may pave the way for experimental confirmation of these findings.

*Fine-mapping and second signal analysis in EA individuals*

Conditional and fine-mapping analyses were restricted to EA participants, for whom data to construct a large enough LD reference panel was publicly available (Methods). The meta-analysis of 85 EA CKDGen GWAS identified 256 genome-wide significant loci (**Supplementary Table 10**). Replication among 216,518 EA MVP participants confirmed 228 SNPs, including 227 index SNPs that met replication criteria and the *SHROOM3* index SNP (Methods, **Supplementary Table 10**). Of these 228 SNPs, 221 mapped into one of the 264 replicated loci from the trans-ethnic analysis (≤±500 kb from the trans-ethnic index SNP), and the remaining 7 showed *P*≤3.3×10-6 in the trans-ethnic discovery analysis. BUN GWAS meta-analysis of CKDGen EA studies (n=243,029) allowed to classify 122 SNPs as likely relevant for kidney function, 90 as inconclusive, and 16 as unlikely (**Supplementary Table 10**).

To perform statistical fine-mapping of the 228 eGFR loci, we first performed summary-statistics based conditional analysis and identified 253 independent genome-wide significant SNPs (**Supplementary Table 11**) mapping into 189 regions (Methods). For each independent variant, we computed a 99% credible set,[32](#_ENREF_32) with median set size of 26 SNPs (IQR: 6, 60). We observed 58 *small credible sets* (≤5 SNPs), including 20 single-SNP sets: *EDEM3*, *CACNA1S*, *HOXD11*, *CPS1*, *DAB2*, *SLC34A1*, *LINC01512*, *LARP4B*, *DCDC1*, *SLC25A45*, *SLC6A13*, *GATM*, *CGNL1*, *CYP1A1*, *NRG4*, *RPL3L*, *UMOD-PDILT,* *SLC47A1,* and two independent sets at *BCL2L14* (**Supplementary Table 11;** **Figure 4**). Of the 58 small credible sets, 33 were likely relevant for kidney function and contain genes and SNPs that can now be prioritized for further study (**Supplementary Table 11**).

Credible set SNPs were annotated with respect to their functional consequence and regulatory potential. Missense SNPs with >50% posterior probability (PP) of driving the association and/or mapping into a small credible set are of particular interest because they directly implicate the affected gene. Such missense SNPs were identified in 11 genes (*SLC47A1*, *RPL3L*, *SLC25A45*, *CACNA1S*, *EDEM3*, *CPS1*, *KLHDC7A*, *PPM1J*, *CERS2*, *C9,* and *SLC22A2;* **Supplementary Table 12**), of which *CACNA1S*, *RPL3L*, *CERS2*, and *C9* were likely relevant for kidney function(**Figure 4A**). The majority of the 11 variants had CADD score>15, indicating potential deleteriousness.[33](#_ENREF_33) Several identified genes are plausible biological candidates for driving the association signal (**Table 1**). For example, the missense p.(Ala465Val) SNP in *SLC47A1* (PP>99%) alters the encoded multidrug and toxin extrusion protein (MATE1), a transport protein responsible for the secretion of cationic drugs, toxins and internal metabolites including creatinine across brush border membranes including kidney proximal tubules. The fact that MATE1 knockout mice have higher blood levels of both creatinine and BUN[34](#_ENREF_34) argues against a sole effect on creatinine transport.

 To evaluate the regulatory potential of small credible set SNPs in kidney, we annotated them to open chromatin regions identified from primary human tubular and glomerular cell cultures,[35](#_ENREF_35) as well as from publicly available kidney cells types (Methods). We identified 72 SNPs mapping into one of these annotations, which may thus represent causal regulatory variants (**Supplementary Table 12**). A particularly interesting finding was the intronic rs77924615 in *PDILT*, which showed PP>99% of driving the association at the *UMOD* locus, and mapped into open chromatin in all evaluated resources (native kidney cells, ENCODE and Roadmap kidney cell types; **Figure 4B**).

*Gene prioritization: co-localization with gene expression*

We performed co-localization analyses for each eGFR-associated locus with gene expression in *cis* across 46 tissues including kidney glomerular and tubulo-interstitial compartments (Methods). A PP>80% of co-localization in at least one kidney tissue was observed for 17 transcripts mapping into 16 of the 228 replicated loci (**Figure 5**), pointing towards a shared underlying SNP associated with both eGFR and gene expression, and implicating the gene encoding the co-localized transcript as the locus’ effector gene(s).

Novel insights emerged on several levels: first, *UMOD* is a well-established causal gene for CKD and can therefore be used to evaluate our workflow. In the tubulo-interstitial compartment, we observed a shared underlying variant associated with higher *UMOD* gene expression and lower eGFR (**Figure 5**), consistent with previous GWAS of urinary uromodulin concentration, in which alleles associated with lower eGFR at *UMOD*[*15*](#_ENREF_15) were associated with higher urinary uromodulin concentrations.[36](#_ENREF_36) The lead SNP at this locus was rs77924615, highlighted above as the candidate causal regulatory variant mapping into the intron of *PDILT* (upstream of *UMOD*). The association with differential *UMOD* but not *PDILT* gene expression supports *UMOD* as the causal gene and rs77924615 as a regulatory SNP.

Second, novel, biologically plausible candidates emerged. For example, our results suggest *KNG1* and *FGF5* as effector genes in the respective eGFR-associated loci (**Figure 5**, **Supplementary Table 13**). *KNG1* encodes for high-molecular weight kininogen, which is cleaved to bradykinin. Bradykinin influences blood pressure, natriuresis and diuresis, and can be linked to kidney function via the renin-angiotensin-aldosterone system.[37](#_ENREF_37) *FGF5* encodes for Fibroblast Growth Factor 5, and the index SNPs for eGFR or highly correlated SNPs (r2>0.9) have been identified in multiple GWAS of blood pressure, atrial fibrillation, coronary artery disease, hematocrit and multiple kidney-function related traits (**Supplementary Table 13**). The eGFR index SNP rs1458038 (PP>50%, CADD score=14.8; **Supplementary Table 13**), co-localized with the eGFR signal only in tubulo-interstitial kidney portions (**Figure 5**), supporting its regulatory potential on the expression levels of *FGF5* in this compartment. Both *KNG1* and *FGF5* index SNPs were associated with BUN and CKD, and are thus likely related to kidney function.

Third, co-localization of eGFR with gene expression across multiple tissues revealed that for kidney-co-localized transcripts, some showed the same direction on transcript levels across all tissues with lower eGFR (e.g. *METTL10*), while others showed higher transcript levels in some tissues but lower levels in others (e.g. *SH3YL1*;**Figure 5**). These observations were also reflected broadly across all transcripts with evidence of co-localization in any tissue (**Supplementary Figure 9**), and highlight tissue-shared and tissue-specific signals.[38](#_ENREF_38),[39](#_ENREF_39)

Finally, *trans*-eQTL annotation of the index SNPs in whole and peripheral blood identified a reproducible link of rs10774625 (12q24.11) with several transcripts (Methods; **Supplementary Material**; **Supplementary Tables 14** and **15**).

*Co-localization with uromodulin protein levels in urine*

The *UMOD* locus is of particular clinical interest for CKD research:[21](#_ENREF_21) rare *UMOD* mutations cause autosomal-dominant tubulo-interstitial kidney disease[40](#_ENREF_40), and common variants at *UMOD* give rise to the strongest eGFR and CKD GWAS signals.[15](#_ENREF_15) We therefore performed conditional analyses based on the EA-specific summary statistics and found two independent variants: rs77924615, mapping into upstream *PDILT*, and rs34882080 mapping into an intron of *UMOD* (**Figure 6A**). SNP association with the urinary uromodulin-to-creatinine ratio (UUCR) in one participating cohort (**Figure 6B**) matched the eGFR association pattern. Co-localization of the conditional eGFR and UUCR associations was evaluated separately for the rs34882080 (**Figure 6C**) and rs77924615 (**Figure 6D**). Both regions showed high probability of a shared underlying variant driving the respective associations with eGFR and UUCR levels (PP=0.97 and 0.96, respectively), further supporting rs77924615 as a causal regulatory variant and *UMOD* as its effector gene.

A summary of the various gene characterization results for replicated loci from the EA analysis is shown in **Supplementary Table 16**, to facilitate selection of the most promising candidates for further experimental studies.

**Discussion**

This trans-ethnic study is 5-fold larger than previous eGFR GWAS meta-analyses and identified 264 replicated loci, 166 of which are reported here for the first time. By also analyzing BUN, an established complementary kidney function marker, we highlight eGFR-associated loci that are likely to be important for kidney function as opposed to creatinine metabolism, and provide a comprehensive annotation resource. Clinical relevance is supported by associations of a GRS for low eGFR with higher odds of clinically diagnosed CKD, CKD-related phenotypes, and hypertension. Enrichment analyses confirm the kidney as the main target organ. Co-localization of associations with eGFR and gene expression in the kidney implicates specific target genes for follow-up. Conditional analyses, fine-mapping and functional annotation at 228 replicated eGFR-associated loci among EA participants implicate single potentially causal variants at 20 loci*.*

 Most previous eGFR GWAS meta-analyses were limited to a single ancestry group[8](#_ENREF_8) and did not prioritize causal variants or effector genes in associated loci. While underpowered to uncover novel loci, one previous trans-ethnic study employed fine-mapping, resolving one signal to a single variant,[20](#_ENREF_20) rs77924615 at *UMOD-PDILT*, also identified in our study. At this locus, we further characterized the relationship between the causal variant, *UMOD* expression in the target tissue, and uromodulin protein levels. This increase in resolution - from locus to single potentially causal variant with its effector gene, protein and target tissue, represents a critical advance over 10 years of eGFR GWAS,[15](#_ENREF_15) and is a prerequisite for translational research.

The complementary multi-tissue approaches including enrichment analyses based on gene expression, regulatory annotations, and gene-sets and pathways highlight the kidney as the most important target organ. However, relatively few kidney-specific experimental datasets are publicly available. For example, the kidney is not well represented in the GTEx Project and not included in its tissue-specific eQTL datasets,[38](#_ENREF_38) which emphasizes the value of open access resources and in depth characterization of uncommon tissues and cell types. We were able to specifically investigate the kidney by using a recently published eQTL dataset from glomerular and tubulo-interstitial portions of micro-dissected human kidney biopsies,[41](#_ENREF_41) kidney-specific regulatory information from the ENCODE and Roadmap resources, and by obtaining regulatory information from primary cultures of human glomerular and tubulo-interstitial cells.[35](#_ENREF_35)

Functional follow-up studies of potentially causal variants will benefit from prioritized loci that show clear evidence supporting one or a few SNPs driving the association signal. The fine-mapping workflow allowed us to prioritize several SNPs at single- or ≤5-SNP resolution, some of which may have broader clinical relevance. For example, the OCT2 protein encoded by *SLC22A2* transports several cationic drugs such as metoprolol, cisplatin, metformin and cimetidine across the basolateral membrane of renal tubular cells.[42](#_ENREF_42) The prioritized missense SNP encodes p.(Ser270Ala), a known pharmaco-genomic variant that alters the transport of these drugs and their side-effects, such as cisplatin-induced nephrotoxicity.[43](#_ENREF_43) Along the same lines, the prioritized SNP encoding the p.(Ala465Val) substitution in the transporter MATE1 encoded by *SLC47A1* may affect the ability to secrete drugs and other toxins from proximal tubular cells into the urine[44](#_ENREF_44) and hence alter CKD risk.

 Strengths of this project include the large sample size with dense genotype imputation, standardized and automated phenotype generation and QC, independent replication, as well as advanced and comprehensive downstream bioinformatics analyses. Further strengths are the use of BUN to prioritize eGFR-associated loci likely relevant for kidney function, and to provide genome-wide BUN summary statistics as a annotation resource for other studies of eGFR. Moreover, we evaluated a GRS for eGFR for association with clinically diagnosed CKD in a large independent study. Among the limitations, non-European populations are still underrepresented in our study, like many other genomic efforts.[45](#_ENREF_45) Statistical fine-mapping using trans-ethnic data with different LD structures can potentially narrow down association signals. However, a sufficiently large reference dataset to compute ancestry-matched LD structure for summary-statistics based fine-mapping was only available for EA, highlighting the potential of future large-scale efforts with trans-ethnic fine-mapping and the need to generate data from non-EA populations enabling such endeavors. Lastly, several SNPs had small effective sample sizes in some subpopulations, which might have affected the ability to assess between-ancestry heterogeneity and potentially underestimated true heterogeneity.

We estimated GFR from serum creatinine, as done in clinical practice and observational studies, because direct measurement of kidney function is invasive, time-consuming, and burdensome. Under the assumption that genetic associations supported by multiple markers are less likely to reflect marker metabolism, we used BUN to prioritize eGFR-loci likely to be relevant to kidney function. Blood creatinine, urea and cystatin C concentrations are influenced not only by glomerular filtration, but also by their synthesis, active secretion, or reabsorption, as illustrated by loci detected in our study: for example, the *GATM* locus was associated with eGFR but not with BUN, consistent with the function of the encoded protein as a rate-limiting enzyme in creatine synthesis.[46](#_ENREF_46) Conversely, the *SLC14A2* locus was associated with BUN but not with eGFR, consistent with the function of the encoded protein as a urea transporter.[47](#_ENREF_47) Even so, lack of a SNP’s association with one kidney function marker based on a combination of p-value and effect direction may not necessarily mean that the locus is not relevant to kidney function. Our categorization of the eGFR loci into three classes based on effect direction and significance of the BUN associations should be interpreted with caution, with “*likely*” and “*unlikely*” reflecting uncertainty of the assignment. Factors complicating the comparison of eGFR and BUN associations at the locus level are differential statistical power, differential ancestry distribution, and potential allelic heterogeneity. Further large-scale studies with multiple kidney function markers measured in the same individuals are therefore warranted.

To identify broadly representative and generalizable association signals, we focused on SNPs that were present in the majority of the participating studies. This choice might have limited our ability to uncover novel or to fine-map low-frequency or population-specific variants, which represents a complementary avenue of research. Moreover, even with well-powered fine-mapping approaches, potentially causal SNPs need to be confirmed as functional variants in experimental studies. Although co-localization with gene expression can help prioritize effector genes, these associations are based on measures from a single time point and hence cannot answer whether changes in gene expression precede or follow changes in kidney function.

In summary, we identified and characterized a large number of loci associated with eGFR and prioritized potential effector genes, driver variants and target tissues. These findings will help direct functional studies and advance the understanding of kidney function biology, a prerequisite to develop novel therapies to reduce the burden of CKD.

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**Online Methods**

*Overview*

We set up a collaborative meta-analysis based on a distributive data model and QC procedures. To maximize phenotype standardization across studies, an analysis plan and a command line script (<https://github.com/genepi-freiburg/ckdgen-pheno>) were created centrally and provided to all participating studies (mostly population-based, **Supplementary Table 1**). Data processing, analysis and troubleshooting instructions were distributed to all studies via a Wiki system ([https://ckdgen.eurac.edu/mediawiki/index.php/
CKDGen\_Round\_4\_EPACTS\_analysis\_plan](https://ckdgen.eurac.edu/mediawiki/index.php/CKDGen_Round_4_EPACTS_analysis_plan)). Automatically generated summary files were checked centrally. Upon phenotype approval, studies run their GWAS and uploaded results and imputation quality (IQ) information to a common calculation server. GWAS QC was performed using GWAtoolbox[48](#_ENREF_48" \o "Fuchsberger, 2012 #59) and custom scripts to assess ancestry-matched allele frequencies and variant positions. All studies had their own research protocols approved by the respective local ethics committees. All participants in all studies provided written informed consent.

*Phenotype definition*

Each study measured serum creatinine and BUN as described in **Supplementary Table 1**. Creatinine values obtained with a Jaffé assay before 2009 were calibrated by multiplying by 0.95.[49](#_ENREF_49) Studies on >18 year-old adults estimated GFR with the Chronic Kidney Disease Epidemiology Collaboration (CKD-EPI) equation,[50](#_ENREF_50) using the R package ‘nephro’.[51](#_ENREF_51) Studies on ≤18 year-old individuals used the Schwartz formula.[52](#_ENREF_52) eGFR was winsorized at 15 and 200 ml/min/1.73m2. CKD was defined as eGFR<60 ml/min/1.73m2. In studies reporting blood urea measurements, BUN was derived as blood urea×2.8, with units expressed as mg/dl.

*Genotyping and genotype imputation*

Genotypes imputed based on the Haplotype Reference Consortium (HRC) v1.1 or the 1000 Genomes Project phase 3 v5 (1000Gp3v5) ALL or phase 1 v3 (1000Gp1v3) ALL panels. Imputed variants were coded as allelic dosages accompanied by the corresponding IQ scores (IMPUTE2 info score, MACH/minimac RSQ, or as applicable), and annotated on the NCBI b37 (hg19) reference build (see **Supplementary Table 2** for study-specific genotyping arrays, haplotype phasing and genotype imputation methods).

*Genome-wide association studies* (GWAS)

Each study fitted sex- and age-adjusted linear regression models to log(eGFR) and BUN. Regression residuals were regressed on SNP dosage levels, assuming an additive genetic model. Study site, genetic principal components (PCs), relatedness, or other study-specific features, were accounted for in the study-specific models as appropriate (**Supplementary Table 2**). Logistic regression models were fitted for CKD.

*Trans-ethnic GWAS meta-analysis*

Studies contributed 121 GWAS summary statistics files for eGFR (total post-QC n=765,348), 60 GWAS files for CKD (total post-QC n=625,219, including 64,164 CKD cases), and 65 GWAS files for BUN (total post-QC n=416,178). Ancestry-specific details for eGFR, CKD and BUN are given in **Supplementary Table 1**.

Before meta-analysis, study-specific GWAS files were filtered to retain only variants with IQ score>0.6 and minor allele count (MAC)>10, and genomic control (GC) correction applied in case of GC factor λGC>1. Fixed effects inverse-variance weighted meta-analysis was performed using METAL,[53](#_ENREF_53) which was adapted to increase the precision of effect estimates and their standard errors (SE; seven decimal places instead of four).

After meta-analysis of 43,994,957 SNPs, only SNPs present in ≥50% of the GWAS files and with total MAC≥400 were retained. Across ancestries, this yielded 8,221,591 variants for eGFR (8,834,748 in EA), 8,176,554 for BUN (8,358,347 in EA), and 9,585,923 for CKD. Post-meta-analysis GC correction was not applied (LD Score regression intercept≈1 in all analyses of eGFR, BUN, and CKD).[54](#_ENREF_54) The genome-wide significance level was set at 5×10-8. Between-study heterogeneity was assessed using the *I2* statistic.[55](#_ENREF_55) For CKD, variants with *I2*≥95% were removed to moderate influence of single large studies. Variants were assigned to loci by selecting the SNP with the lowest p-value genome-wide as the index SNP, defining the corresponding locus as the 1 Mb-segment centered on the index SNP, and repeating the procedure until no further genome-wide significant SNPs remained. The extended MHC region was considered as a single locus. A locus was considered novel if not containing any variant identified by previous GWAS of eGFR.

*Meta-regression analysis of trans-ethnic GWAS*

For eGFR, we evaluated ancestry-related heterogeneity using the software Meta-Regression of Multi-Ethnic Genetic Association (MR-MEGA v0.1.2)[56](#_ENREF_56) using study-specific GWAS results. Meta-regression models included three axes of genetic variation. GC correction was applied to the meta-regression results. The 308 genome-wide significant index SNPs from the trans-ethnic GWAS meta-analysis were tested for ancestry-related heterogeneity of the allelic effects at a significance level of 0.05/308=1.6×10-4 (referring to the corresponding p-value as p-anc-het).

*Variance explained and genetic heritability*

The proportion of phenotypic variance explained by the index SNPs was estimated as $β^{2}\left(\frac{2p\left(1-p\right)}{var}\right)$, with *β* being the SNP effect, *p* the effect allele frequency, and *var* the variance of the sex- and age-adjusted log(eGFR) residuals (assumed as 0.016 based on data from 11,827 EA participants of the population-based ARIC study).[9](#_ENREF_9) Genetic heritability of age- and sex-adjusted log(eGFR) was estimated using the R package ‘MCMCglmm’[57](#_ENREF_57) on the Cooperative Health Research In South Tyrol (CHRIS) study,[58](#_ENREF_58) a participating pedigree-based study with 186 up-to-5 generation pedigrees (n=4373).[59](#_ENREF_59) We fitted two models, with and without the inclusion of the identified index SNPs (304/308), running 1,000,000 MCMC iterations (*burn in*=500,000).[59](#_ENREF_59)

*Comparison with and replication of results in the MVP*

The eGFR-associated SNPs identified in the discovery GWAS meta-analyses, were tested for replication in a GWAS from the MVP,[23](#_ENREF_23) an independent trans-ethnic study with participants recruited across 63 U.S. Veteran’s Administration (VA) medical facilities. Written informed consent was obtained and all documents and protocols were approved by the VA Central Institutional Review Board. After genotyping and QC, genotypes were phased and imputed on the 1000Gp3v5 reference panel. Serum creatinine was assessed up to one year prior to MVP enrollment using isotope dilution mass spectrometry. GFR was estimated using the CKD-EPI equation[50](#_ENREF_50) after excluding subjects on dialysis, transplant patients, amputees, individuals on HIV medications, and those with creatinine values of <0.4 mg/dl. GWAS of eGFR on SNP dosage levels were performed by fitting linear regression models adjusted for age at creatinine measurement, age2, sex, body mass index, and the first 10 genetic PCs, using SNPTEST v2.5.4-beta.[60](#_ENREF_60) All GWAS were stratified by self-reported ethnicity (79.6% White non-Hispanic; 20.4% Black non-Hispanic), diabetes, and hypertension status. Results were combined across strata using fixed effects inverse-variance weighted meta-analysis in METAL.[53](#_ENREF_53) This analysis encompassed a total of 280,722 individuals across all strata, of whom 216,518 were non-Hispanic Whites (EA). The MVP is described more extensively in the **Supplementary Material**.

Of the 308 eGFR index SNPs identified in the CKDGen trans-ethnic analysis, 305 variants or their good proxies were available in the MVP GWAS (proxies had to have r2≥0.8 with the index SNP and were selected by maximum r2 followed by minimum distance in case of ties). Replication testing of the 256 EA-specific index SNPs was restricted to the MVP EA GWAS. CKDGen and MVP meta-analysis results were pooled via sample size weighted meta-analysis of z-scores using METAL.[53](#_ENREF_53) In both the trans-ethnic and EA-specific analyses, replication was defined as a one-sided p-value<0.05 in the MVP and genome-wide significance of the CKDGen and MVP meta-analysis result.

*Assessment of kidney function relevance using BUN*

We used genetic associations with BUN to assess replicated eGFR-associated SNPs with respect to their potential kidney function relevance. Support for kidney function relevance was categorized as “likely” (1) for all eGFR index SNPs with an inverse, significant (one-sided *P*<0.05) association with BUN for a given reference allele, “inconclusive” (2) for eGFR index SNPs whose effect on BUN was not different from 0 (*P≥*0.05), and “unlikely” (3) for all eGFR index SNPs with a concordant, significant (one-sided *P*<0.05) association with BUN for a given reference allele.

*Genetic risk score (GRS) analysis in the UK Biobank dataset*

To test the combined effect of eGFR-associated SNPs on clinically diagnosed CKD related outcomes, a GRS-based association analysis was conducted based on summary GWAS results, as described before.[61](#_ENREF_61),[62](#_ENREF_62) The genetic association results with the diseases were obtained for 452,264 UK Biobank participants available in the GeneAtlas[63](#_ENREF_63" \o "Canela-Xandri, 2018 #114) database for glomerular diseases (ICD-10 codes N00-N08; 2289 cases); acute renal failure (N17; 4913 cases); chronic renal failure (N18; 4905 cases); urolithiasis (N20-N23; 7053 cases); hypertensive diseases (I10-I15; 84,910 cases); and ischemic heart diseases (I20-I25; 33,387 cases). Asthma (J45; 28,628 cases) was included as a negative control. The log(estimated OR) provided by the GeneAtlas phewas interface was used as effect size, and its SE was calculated from the corresponding effect size and p-value. When OR=1, the SE was imputed by the median value of the remaining associations of the trait. Of the 147 eGFR index SNPs from the trans-ethnic GWAS meta-analysis that were replicated and showed likely kidney function relevance, 144 were available in the UK Biobank dataset, and 259 out of all 264 replicated trans-ethnic GWAS meta-analysis SNPs. The effect *beta* of the GRS association corresponds to the OR of the disease depending on the relative change in eGFR, e.g. OR=1.10*beta* for a 10% change in eGFR. Alternatively, exp(*beta*) can be interpreted as the OR of the disease per unit change of log(eGFR).

*Genetic correlations with other complex traits and diseases*

Genome-wide genetic correlation analysis was performed to investigate evidence of co-regulation or shared genetic bases between eGFR and BUN and other complex traits and diseases, both known and not known to correlate with eGFR and BUN. We estimated pairwise genetic correlation coefficients (rg) between the results of our trans-ethnic meta-analyses of eGFR and BUN and each of 748 pre-computed and publicly available GWAS summary statistics of complex traits and diseases available through LD Hub v1.9.0 using LD Score regression.[28](#_ENREF_28) An overview of the sources of these summary statistics and their corresponding sample sizes is available at <http://ldsc.broadinstitute.org>. Statistical significance was assessed at the Bonferroni corrected level of 0.05/748=6.7×10-5.

*Pathway and tissue enrichment analysis*

We used DEPICT v1 release 194 to perform Data-Driven Expression Prioritized Integration for Complex Traits analysis,[29](#_ENREF_29) including pathway/gene-set enrichment and tissue/cell type analyses as described previously.[9](#_ENREF_9),[10](#_ENREF_10) All 14,461 gene sets were reconstituted by identifying genes that were transcriptionally co-regulated with other genes in a panel of 77,840 gene expression microarrays,[64](#_ENREF_64) from mouse knock-out studies, and molecular pathways from protein-protein interaction screening. In the tissues and cell type enrichment analysis, we tested whether genes in associated regions were highly expressed in 209 MeSH annotation categories for 37,427 microarrays (Affymetrix U133 Plus 2.0 Array platform). For both eGFR and BUN, we included all variants associated with the trait at *P*<5×10-8 in the trans-ethnic meta-analysis. Independent variant clumping was performed using Plink 1.9[65](#_ENREF_65) with 500 kb flanking regions and r²>0.01 in the 1000Gp1v3 dataset. After excluding the MHC region, DEPICT was run with 500 repetitions to estimate the FDR and 5000 permutations to compute p-values adjusted for gene length by using 500 null GWAS. All significant gene sets were merged into meta gene sets by running an affinity propagation algorithm[66](#_ENREF_66) implemented in the Python ‘scikit-learn’ package (<http://scikit-learn.org/>). The resulting network was visualized using Cytoscape (<http://cytoscape.org/>).

*Enrichment of heritability by cell type group*

We used stratified LD Score regression to investigate important tissues and cell types based on the trans-ethnic eGFR and BUN meta-analysis results. Heritability enrichment in 10 cell type groups was assessed using the default options of stratified LD Score regression described previously.[30](#_ENREF_30) The 10 cell type groups were collapsed from 220 cell-type specific regulatory annotations for the four histone marks H3K4me1, H3K4me3, H3K9ac, and H3K27ac. The enrichment of a cell type category was defined as the proportion of SNP heritability in that group divided by the proportion of SNPs in the same cell type group.

*Analysis of genes causing kidney phenotypes in mice*

A nested candidate gene analysis was performed using GenToS[67](#_ENREF_67" \o "Hoppmann, 2016 #16) to identify additional genetic associations that were not genome-wide significant. Candidate genes that when manipulated cause kidney phenotypes in mice were selected using the comprehensive Mouse Genome Informatics (MGI) phenotype ontology in September 2017 (abnormal renal glomerular filtration rate [MP:0002847]; abnormal kidney morphology [MP:0002135]; abnormal kidney physiology [MP:0002136]). The human orthologs of these genes were obtained, when available, using the Human-Mouse: Disease Connection webtool (<http://www.informatics.jax.org/humanDisease.html>). Statistical significance was defined as a Bonferroni correction of a type I error level of 0.05 for the number of independent common SNPs across all genes in each of the three candidate gene lists plus their flanking regions, derived from an ancestry-matched reference population. The GWAS meta-analysis summary statistics for eGFR were queried for significantly associated SNPs mapping into the selected candidate genes. Enrichment of significant genetic associations in genes within each candidate list was computed from the complementary cumulative binomial distribution.[67](#_ENREF_67) GenToS was used with default parameters on each of the three candidate gene lists, using the 1000 Genomes phase 3 release 2 ALL dataset as reference.

*Independent variant identification in the EA meta-analysis*

To identify additional, independent eGFR-associated variants within the EA-specific and replicated loci, approximate conditional analyses were performed based on genome-wide discovery summary statistics that incorporated LD information from an ancestry-matched reference population. These analyses were restricted to participants of EA, because an LD reference sample scaled to the size of our meta-analysis could only be constructed from publicly available data for EA individuals,[24](#_ENREF_24) for which we randomly selected 15,000 UK Biobank participants (dataset ID 8974). Individuals who withdrew consent and those not meeting data cleaning requirements were excluded, keeping only those who passed sex-consistency check, had ≥95% call rate, and did not represent outliers with respect to SNP heterozygosity. For each pair of individuals, the proportion of variants shared identical-by-descent (IBD) was computed using PLINK.[68](#_ENREF_68) Only one member of each pair with IBD coefficient ≥0.1875 was retained. Individuals were restricted to those of EA by excluding outliers along the first two PCs from a PC analysis seeded with the HapMap phase 3 release 2 populations as reference. The final dataset to estimate LD included 13,558 EA individuals and 16,969,363 SNPs.

The basis for statistical fine-mapping were the 228 1-Mb genome-wide significant loci identified in the EA meta-analysis, clipping at chromosome borders. Overlapping loci as well as pairs of loci whose respective index SNPs were correlated (r² >0.1 in the UKBB LD dataset described above) were merged. A single SNP was chosen to represent the MHC region, resulting in a final list of 189 regions prior to fine-mapping. Within each region, the GCTA COJO Slct algorithm[69](#_ENREF_69) was used to identify independent variants employing a step-wise forward selection approach. We used the default collinearity cut-off of 0.9 (sensitivity analyses showing no major influence of alternative cutoff values; data not shown). We deemed an additional SNP as independently genome-wide significant if the SNPs’ p-value conditional on all previously identified SNPs in the same region was <5×10-8.

*Fine-mapping and credible sets in the EA meta-analysis*

For each region containing multiple independent SNPs and for each independent SNP in such regions, approximate conditional analyses were conducted using the GCTA COJO-Cond algorithm to generate approximate conditional association statistics conditioned on the other independent SNPs in the region. Using the Wakefield’s formula implemented in the R package ’gtx’,[70](#_ENREF_70) we derived approximate Bayes factors (ABF) from conditional estimates in regions with multiple independent SNPs and from the original estimates for regions with a single independent SNP. Given that 95% of the SNP effects on log(eGFR) fell within -0.01 to 0.01, the standard deviation prior was chosen as 0.0051 based on formula 8 in the original publication.[32](#_ENREF_32) Sensitivity analyses showed that results were robust when higher values were used for the standard deviation prior (data not shown). For each variant within an evaluated region, the ABF obtained from the association betas and their SEs of the marginal (single signal region) or conditional estimates (multi-signal regions) was used to calculate the PP for a SNP of driving the association signal (“causal variant”). We derived 99% credible sets, representing the SNP sets containing the variant(s) driving the association signal with 99% probability, by ranking variants by their PP and adding them to the set until reaching a cumulative PP>99% in each region.

*Variant annotation*

Functional annotation of SNPs mapping into credible sets was performed with SNiPA v3.2 (March 2017),[71](#_ENREF_71) based on the 1000Gp3v5 and Ensembl v87 datasets. SNiPA was also used to derive the Combined Annotation Dependent Depletion (CADD) PHRED-like score,[72](#_ENREF_72) based on CADD v1.3. The Ensembl VEP tool[73](#_ENREF_73) was used for SNP’s primary effect prediction.

*Co-localization of eGFR and gene expression in cis*

As the great majority of gene expression datasets is generated based on EA ancestry samples, co-localization analysis was based on the genetic associations with eGFR in the EA sample and with gene expression (eQTL) quantified from micro-dissected human glomerular and tubulo-interstitial kidney portions from 187 individuals from the NEPTUNE study,[41](#_ENREF_41) as well as from the 44 tissues included in the GTEx Project v6p release.[38](#_ENREF_38) The eQTL and GWAS effect alleles were harmonized. For each locus, we identified tissue gene pairs with reported eQTL data within ±100 kb of each GWAS index SNP. The region for each co-localization test was defined as the eQTL *cis* window defined in the underlying GTEx and NephQTL studies. We used the ‘coloc.fast’ function, using default setting, from the R package ‘gtx’ (https://github.com/tobyjohnson/gtx), which is an adaption of Giambartolomei’s co-localization method.[74](#_ENREF_74) ‘gtx’ was also used to estimate the direction of effect over the credible sets as the ratio of the average PP-weighted GWAS effects over the PP-weighted eQTL effects.

*Trans-eQTL analysis*

We performed *trans*-eQTL annotation through LD mapping based on the 1000Gp3v5 European reference panel (r2 cut-off >0.8). We limited annotation to replicated index SNPs with a fine-mapping PP≥1%. Due to expected small effect sizes, only genome-wide *trans*-eQTL studies of either peripheral blood mononuclear cells or whole blood with n≥1000 individuals were considered, resulting in five non-overlapping studies[75-79](#_ENREF_75) (**Supplementary Table 14**). For one study,[79](#_ENREF_79) we had access to an update with larger sample size (n=6645) obtained by combining two non-overlapping studies (LIFE-Heart[80](#_ENREF_80) and LIFE-Adult[81](#_ENREF_81)). To improve stringency of results, we focused the analysis on inter-chromosomal *trans*-eQTLs with *P*<5×10-8 in ≥2 studies.

*Co-localization with urinary uromodulin concentrations*

Association between concentrations of the urinary uromodulin-to-creatinine ratio with genetic variants at the *UMOD-PDILT* locus was evaluated in the German Chronic Kidney Disease (GCKD) study.[82](#_ENREF_82) Uromodulin concentrations were measured from frozen stored urine using an established ELISA assay with excellent performance.[36](#_ENREF_36) Concentrations were indexed to creatinine to account for urine dilution. Genetic associations were assessed using the same software and settings as for the eGFR association (**Supplementary Table 2**). Co-localization analyses were performed using identical software and settings as described above for the association with gene expression.

*Life Sciences Reporting Summary*

Further information on research methods are available in the Nature Research Life Sciences Reporting Summary linked to this article.

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**Data Availability**

Genome-wide summary statistics for this study are made publicly available through dbGaP accession number phs000930.v7.p1 and at <http://ckdgen.imbi.uni-freiburg.de>.

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**Figure Legends**

**Figure 1 – Trans-ethnic GWAS meta-analysis identifies 308 loci associated with eGFR**

Circos plot: Red band: –log10(*P*) for association with eGFR, by chromosomal position. Blue line indicates genome-wide significance (*P*=5×10-8). Black gene labels indicate novel loci, blue labels known loci. Non-replicating loci are colored in gray (novel) or light blue (known). Green band: Measures of heterogeneity related to the index SNPs associated with eGFR. Dot sizes are proportional to I² or ancestry-related heterogeneity (p-anc-het). Blue band: –log10(*P*) for association with CKD, by chromosomal position. Red line indicates genome-wide significance (*P*=5×10-8). Radial lines mark regions with p-anc-het <1.6×10-4 = 0.05/308 or I² >25%. Inset: Effects of all 308 index SNPs on log(eGFR) by minor allele frequency, color-coded by the associated odds ratio (OR) of CKD (red scale for OR≤1, blue scale for OR>1). The largest effects on CKD were observed for rs77924615 at *UMOD-PDILT* (odds ratio (OR) = 0.81, 95%CI: 0.80, 0.83), rs187355703 at *HOXD8* (OR=0.82, 95%CI: 0.77, 0.87), and rs10254101 at *PRKAG2* (OR=1.11, 95%CI: 1.09, 1.11). Triangles highlight SNPs that were associated with CKD (one-sided *P*<0.05).

**Figure 2 – Generalizability with respect to other populations and other kidney function markers**

**Panel A: Measures of heterogeneity for 308 eGFR-associated index SNPs.** Comparison of each variant’s heterogeneity quantified as I² from the trans-ethnic meta-analysis (Y-axis) vs. ancestry-related heterogeneity from meta-regression (-log10(p-anc-het), X-axis). Histograms summarize the distribution of the heterogeneity measures on both axes. SNPs with ancestry-related heterogeneity (p-anc-het<1.6×10-4 = 0.05/308) are marked in blue and labeled; SNPs with I²>50% are labeled. **Panel B: Comparison of genetic effect estimates between CKDGen Consortium discovery (X-axis) and MVP replication (Y-axis).** Blue font indicates one-sided *P*<0.05 in the MVP. Error bars indicate 95% confidence intervals (CIs). Dashed line: line of best fit. Pearson’s correlation coefficient r=0.92, 95%CI: 0.90, 0.94. **Panel C: Comparison of the magnitude of genetic effects on eGFR (X-axis) vs. BUN (Y-axis). for the 264 replicated eGFR-associated index SNPs**. Color coding reflects the evidence of kidney-function relevance (see Methods), which is coded as *likely* (blue), *inconclusive* (gray), or *unlikely* (green). Error bars indicate 95% CIs. Dashed line: line of best fit. Correlation: r=-0.65, 95%CI: -0.72, -0.58. **Panel D**: **Association** **between lower genetically-predicted eGFR based on a genetic risk score (GRS) of 147 SNPs likely to be most relevant for kidney function and ICD-10 based clinical diagnoses from 452,264 individuals from the UK Biobank**. Asthma was included as a negative control. Displayed are odds ratios and their 95% CIs per 10% lower GRS-predicted eGFR (Methods).

**Figure 3 – Human orthologs of genes with renal phenotypes in genetically manipulated mice are enriched for association signals with eGFR**

Signals in candidate genes identified based on the murine phenotypes abnormal GFR (**Panel** **A**),abnormal kidney physiology (**Panel** **B**), and abnormal kidney morphology (**Panel** **C**). Y-axis: –log10(*P*) for association with eGFR in the trans-ethnic meta-analysis for the variant with the lowest p-value in each candidate gene. Dashed line indicates genome-wide significance (*P*=5×10-8), solid gray line indicates the experiment-wide significance threshold for each nested candidate gene analysis (included in lower right corner in each panel). Orange color indicates genome-wide significance, red color experiment-wide but not genome-wide significance, and blue color indicates genes with no significantly associated SNPs. Genes are labeled when reaching experiment- but not genome-wide significance; black font for genes not mapping into loci reported in the main analysis, gray font otherwise. Enrichment p-value reported for observed number of genes with association signals below the experiment-wide threshold against the expected number based on the complementary cumulative binomial distribution (Methods).

**Figure 4** **– Credible set size (X-axis) against variant posterior probability (Y-axis) of 3655 SNPs in 253 99% credible sets by annotation**

**Panel** **A: Exonic variants.** SNPs are marked by triangles, with size proportional to their CADD score. Red triangles and variant labeling indicate missense SNPs mapping into small (≤5 SNPs) credible sets or with high individual posterior probability of driving the association signal (>0.5). **Panel** **B: Regulatory potential.** Symbol colors identify variants with regulatory potential as derived from DNAse hypersensitivity analysis in target tissues (Methods). Variant annotation was restricted to variants with variant PP>1%; SNPs with PP≥90% contained in credible sets with ≤10 SNPs were labeled.

**Figure 5 – Co-localization of eGFR-association signals with gene expression in kidney tissues**

All eGFR loci were tested for co-localization with all eQTLs where the eQTL cis-window overlapped (±100 kb) the sentinel genetic variants. Genes with ≥1 positive co-localization (posterior probability of one common causal variant, H4, ≥0.80) in a kidney tissue are illustrated with the respective sentinel SNP (Y-axis). Co-localizations across all tissues (X-axis) are illustrated as dots, where the size of the dots indicates the posterior probability of the co-localization. Negative co-localizations (posterior probability of H4 <0.80) are marked in gray, while the positive co-localizations are color-coded based on the predicted change in expression relative to the allele associated with lower eGFR.

**Figure 6 – Co-localization of independent eGFR-association signals at the *UMOD*-*PDILT* locus with urinary uromodulin concentrations supports *UMOD* as the effector gene.**

Association plots: association –log10(p-value) (Y axis) vs. chromosomal position (X axis). Approximate conditional analyses among EA individuals support the presence of two independent eGFR-associated signals (**Panel A**). The association signal with urinary uromodulin/creatinine levels looks similar (**Panel B**); r2=0.93 between rs34882080 and rs34262842. Co-localization of association with eGFR (upper sub-panel) and urinary uromodulin/creatinine levels (lower sub-panel) for the independent regions centered on *UMOD* (**Panel C**) and *PDILT* (**Panel D**) support a shared underlying variant in both regions with high posterior probability.

**Table 1** – **Genes implicated as causal via identification of missense SNPs with high probability of driving the eGFR association signal.** Genes are included if they contain a missense SNP with posterior probability of association of >50% or mapping into a small credible set (≤5 SNPs).

|  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- |
| **Gene** | **SNP1** | **Credible set size**  | **SNP PP**2 | **Functional consequence** | **CADD score**3 | **DHS**4**, tissue** | **Brief summary of the gene’s function and relevant literature** (OMIM entries are indicated as #number) |
| *CACNA1S* | **rs3850625** | 1 | 1.00 | p.(Arg1539Cys)(NP\_000060.2) | 34.0 | - | Encodes a subunit of the slowly inactivating L-type voltage-dependent calcium channel in skeletal muscle. Reports of altered expression in kidney cancer (PMID 28781648) and after indoxyl sulfate treatment (PMID: 27550174). Rare variants can cause autosomal dominant hypokalemic periodic paralysis, type 1 (#170400) or malignant hyperthermia susceptibility (#601887). Common variation at this locus has been reported as associated with eGFR in previous GWAS (PMID: 24029420, PMID: 26831199). |
| *CPS1* | rs1047891 | 1 | 1.00 | p.(Thr1406Asn) (NP\_001866.2) | 22.1 | - | Encodes a key mitochondrial enzyme of the urea cycle that catalyzes the synthesis of carbamoyl phosphate from ammonia and bicarbonate to remove excess urea. Rare mutations cause autosomal recessive carbamoylphosphate synthetase I deficiency (#237300). GWAS locus for eGFR (PMID: 20383146), serum metabolites (PMID: 23378610), and urinary glycine (PMID: 26352407), as well as for many other quantitative biomarkers. This variant has been reported to associate with hyperammonemia after valproate therapy (PMID: 23997965).  |
| *EDEM3* | rs78444298 | 1 | 1.00 | p.(Pro746Ser)(NP\_079467.3) | 24.6 | - | The gene product accelerates the glycoprotein ER-associated degradation by proteasomes by catalyzing mannose trimming from Man8GlcNAc2 to Man7GlcNAc2 in the N-glycans. This variant has been identified by a previous exome chip association study with eGFR (PMID: 27920155). |
| *KLHDC7A* | rs11261022 | 7 | 0.71 | p.(Arg160Ser)(NP\_689588.2) | 1.1 | Roadmap, ENCODE kidney | Kelch Domain Containing 7A is a protein coding gene and a paralog of *KBTBD11*. No specific entry in relation to kidney disease in PubMed.  |
| *RPL3L* | **rs113956264** | 1 | 1.00 | p.(Val262Met)(NP\_005052.1) | 27.2 | - | The gene product shares sequence similarity with ribosomal protein L3. It has a tissue-specific expression pattern, with highest levels in skeletal muscle and heart. |
| *SLC25A45* | rs34400381 | 1 | 1.00 | p.(Arg285Cys)(NP\_001070709.2) | 26.0 | ENCODE kidney | Belongs to the SLC25 family of mitochondrial carrier proteins and is an orphan transporter. This variant has already been identified in a GWAS of symmetric dimethylarginine levels (PMID: 24159190) and in a whole-genome sequence (WGS) analysis of serum creatinine (PMID: 25082825). *SLC25A45* may play a role in biosynthesis of arginine, which is involved in the synthesis of creatine.  |
| *SLC47A1* | rs111653425 | 1 | 1.00 | p.(Ala465Val)(NP\_060712.2) | 24.6 | - | Encodes the multidrug and toxin extrusion protein (MATE1), a transport protein responsible for the secretion of cationic drugs and creatinine across brush border membranes. This variant has already been identified in a WGS analysis of serum creatinine from Iceland (PMID: 25082825). Rare and common variants in the locus have been identified in exome chip (PMID: 27920155) and in GWAS (PMID: 20383146) studies of eGFR, respectively. MATE1 knockout (KO) mice show higher levels of serum creatinine and BUN (PMID: 19332510), arguing against a sole effect on creatinine transport and supporting an effect on kidney function.  |
| *PPM1J* | rs34611728 | 5 | 0.02 | p.(Leu213Phe)(NP\_005158.5) | 13.1 | ENCODE kidney | This gene encodes the serine/threonine protein phosphatase. The variant has been reported in association with eGFR in an exome chip association study (PMID: 27920155). |
| *CERS2* | **rs267738** | 5 | 0.46 | p.(Glu115Ala)(NP\_071358.1) | 32.0/ 28.2 | - | Encodes Ceramide Synthase 2, which may be involved in sphingolipid synthesis. Changes in ceramides were reported as essential in renal Madin-Darby Canine Kidney (MDCK) cell differentiation (PMID: 28515139). *CERS2* KO mice show strongly reduced ceramide levels in the kidney and develop renal parenchyma abnormalities (PMID: 19801672). This variant has been reported as associated with the rate of albuminuria increase in patients with diabetes (PMID: 25238615). |
| *C9* | **rs700233** | 5 | 0.32 | p.(Arg5Trp)(NP\_001728.1) | 6.6 | - | Encodes a constituent of the membrane attack complex that plays a key role in the innate and adaptive immune response. Rare mutations can cause C9 deficiency (#613825). *C9* is mentioned in several kidney disease case reports, including patients with congenital factor 9 deficiency showing IgA nephropathy (PMID: 1453611).  |
| *SLC22A2* | rs316019 | 4 | 0.04 | p.(Ser270Ala)(NP\_003049.2) | 12.7 | - | Encodes the polyspecific organic cation transporter (OCT2) that is primarily expressed in the kidney, where it mediates tubular uptake of organic compounds including creatinine from the circulation. Many publications relate *SLC22A2* to kidney function. rs316019 is a known pharmacogenomics variant associated with response to metformin and other drugs such as cisplatin. Carriers of the risk allele have a higher risk of cisplatin-induced nephrotoxicity (PMID: 19625999), indicating that this transporter is essential in excreting toxins. The locus has been reported in previous GWAS of eGFR (PMID: 20383146). |

1Boldface indicates SNPs most likely to be relevant for kidney function based on the combined effects on eGFR and BUN;  2PP: posterior probability. 3CADD score: Combined Annotation Dependent Depletion (CADD) PHRED-like score (Methods); 4DHS: DNAse Hypersensitivity Site

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