Novel Mechanistic PBPK Model to Predict Renal Clearance in Varying Stages of CKD by Incorporating Tubular Adaptation and Dynamic Passive Reabsorption

Weize Huang¹ and Nina Isoherranen¹,*

Chronic kidney disease (CKD) has significant effects on renal clearance (CLr) of drugs. Physiologically-based pharmacokinetic (PBPK) models have been used to predict CKD effects on transporter-mediated renal active secretion and CLr for hydrophilic nonpermeable compounds. However, no studies have shown systematic PBPK modeling of renal passive reabsorption or CLr for hydrophobic permeable drugs in CKD. The goal of this study was to expand our previously developed and verified mechanistic kidney model to develop a universal model to predict changes in CLr in CKD for permeable and nonpermeable drugs that accounts for the dramatic nonlinear effect of CKD on renal passive reabsorption of permeable drugs. The developed model incorporates physiologically-based tubular changes of reduced water reabsorption/increased tubular flow rate per remaining functional nephron in CKD. The final adaptive kidney model successfully (absolute fold error (AFE) all < 2) predicted renal passive reabsorption and CLr for 20 permeable and nonpermeable test compounds across the stages of CKD. In contrast, use of proportional glomerular filtration rate reduction approach without addressing tubular adaptation processes in CKD to predict CLr generated unacceptable CLr predictions (AFE = 2.61–7.35) for permeable compounds in severe CKD. Finally, the adaptive kidney model accurately predicted CLr of para-amino-hippuric acid and memantine, two secreted compounds, in CKD, suggesting successful integration of active secretion into the model, along with passive reabsorption. In conclusion, the developed adaptive kidney model enables mechanistic predictions of in vivo CLr through CKD progression without any empirical scaling factors and can be used for CLr predictions prior to assessment of drug disposition in renal impairment.

Study Highlights

WHAT IS THE CURRENT KNOWLEDGE ON THE TOPIC? ✔ Physiologically-based pharmacokinetic (PBPK) modeling has been used to simulate the effect of chronic kidney disease (CKD) on renal clearance (CLr) for hydrophilic nonpermeable drugs with active secretion, but not for hydrophobic permeable drugs with passive reabsorption. The overall confidence and consistency in PBPK modeling of CKD is low.

WHAT QUESTION DID THIS STUDY ADDRESS? ✔ This study quantifies the impact of CKD on tubular flow rate, subsequent passive reabsorption, and resulting CLr, using a novel adaptive kidney model.

WHAT DOES THIS STUDY ADD TO OUR KNOWLEDGE? ✔ A novel physiologically-based mechanistic adaptive kidney model was developed to capture the effect of CKD on tubular flow rate, allowing accurate prediction of passive reabsorption and CLr throughout CKD progression using in vitro data without scaling factors. We successfully verified the adaptive model and showed that proportional glomerular filtration rate scaling approach is inadequate for predicting CLr in CKD.

HOW MIGHT THIS CHANGE DRUG DISCOVERY, DEVELOPMENT, AND/OR THERAPEUTICS? ✔ Our adaptive model enables successful CLr prediction of secreted, nonsecreted, permeable, and nonpermeable drugs and metabolites at all CKD stages, facilitating dosing regimen optimization and trial design prior to renal impairment studies.

Chronic kidney disease (CKD) is a progressive illness that is pathologically heterogeneous¹ and systemic.² It is mainly characterized by declining functional nephron mass and glomerular filtration rate (GFR).³ As such, GFR is a critical index for CKD diagnosis, progression, and classification.⁴ Clinically, patients with severe (stage 4, GFR ~ 15–29 mL/min) and end-stage (stage 5, GFR < 15 mL/min) CKD are at high risk for comorbidities, polypharmacy, and adverse drug reactions,⁵–⁸ necessitating careful medication management due to dramatically altered pharmacokinetics (PK) because of CKD. As a result, clinical characterization of drug
disposition in patients with CKD is critically important. However, predicting the disease effects on drug PK and estimating the optimal dosing regimen is challenging and often unreliable prior to dedicated renal impairment studies due to several factors. For instance, renal clearance (CL) may decrease more than proportionally with GFR in CKD stages 4 and 5 for highly renally secreted drugs, due to inhibition of renal transporters by uremic solutes. CL can also decrease less than proportionally with GFR in CKD stages 4 and 5 for highly renally reabsorbed drugs due to tubular adaptation/compensation processes. Nonetheless, the most commonly accepted approach to predict CL in patients with CKD is to follow the Intact Nephron Hypothesis and assume all renal drug handling pathways and CL decrease proportionally with filtration. Although this assumption is supported by clinical data for predominantly filtered drugs without significant secretion or reabsorption, such proportional GFR scaling approach is inadequate to capture active secretion or passive reabsorption in severe CKD due to the complex disease effects on renal drug handling.

Physiologically-based pharmacokinetic (PBPK) models that capture disease-specific characteristics can potentially predict CKD effects on drug PKs more accurately than proportional GFR based scaling. Yet, use of PBPK models in predicting drug disposition in renal impairment scenarios is currently considered a low-confidence application, suggesting that current PBPK models do not fully capture the physiological changes observed in CKD. PBPK modeling has been used to specifically predict the CKD effect on renal active secretion, but the test drugs used for model verification all lack significant permeability and hence passive reabsorption. At present, no PBPK model can quantitatively predict the alterations in renal passive reabsorption or CL for highly reabsorbed drugs in CKD, let alone systematically capture the diverse effects of CKD on renal elimination.

Physiologically, patients with CKD show reduced water reabsorption/increased tubular flow rates per remaining functional nephron as an adaptation mechanism. This is to compensate for the reduced number of functional nephrons and reduced GFR, to maintain critical homeostasis, such as extracellular fluid volume and plasma sodium concentration. We hypothesized that such physiological adaptation in renal tubular water reabsorption and tubular flow rate will dramatically decrease the drug concentration gradient between intratubular filtrate and peritubular blood, leading to reduced passive reabsorption and higher CL than expected from residual GFR alone. Indeed, this hypothesis is supported by observed data for various permeable drugs, such as pefloxacin, metronidazole, and minocycline. For these drugs, the observed CL in patients with CKD was reduced to 30–37% of healthy when residual GFR was only 4–9% of healthy, demonstrating an over 7-fold disproportionality between the change in CL and GFR.

To simulate the effect of CKD on tubular passive reabsorption, we expanded our previously developed and verified mechanistic kidney model to incorporate the physiologically-based tubular changes of reduced water reabsorption/increased tubular flow rate per remaining functional nephron across varying CKD stages. The developed adaptive kidney model enables prediction of changes in passive reabsorption and CL, from in vitro drug permeability data in different CKD stages. The model was globally verified across the progression of CKD from healthy to end-stage CKD using a set of 20 test compounds with differing properties without any compound-specific empirical scaling.

**METHODS**

**Development and sensitivity analyses of proportional and adaptive kidney model for CKD**

Two distinct CKD models were built using MATLAB and Simulink platform (R2018a; MathWorks, Natick, MA) to predict CL in patients with different stages of CKD. Both models were established based on our previously published and verified physiologically-based mechanistic kidney model for healthy humans, but with modified system parameters to reflect physiological changes in CKD. The first model, the proportional model, was built assuming all functions of the nephron, all pathways of renal drug handling, and therefore the values of CL decline proportionately with GFR. The second model, the adaptive model, was built by accounting for the tubular adaptation of reduced water reabsorption/increased water excretion per remaining functional nephron, as observed in many patients with CKD.

In both models, all of the volume (n = 33), surface area (n = 22), peritubular renal blood flow (n = 12), basolateral uptake clearance (n = 3), and apical efflux clearance (n = 3) parameters were reduced proportionally with GFR by multiplying the baseline parameters in healthy subjects by the fraction of GFR remaining (ratio of GFR in a specific CKD stage, over 120 mL/min, the healthy GFR). The major distinction between the proportional model and the adaptive model is the parameterization of renal tubular flow rate (TFR) in patients with CKD. In the proportional model, all TFR parameters were decreased proportionally with GFR like other model parameters. In contrast, the adaptive model incorporates the physiologically-based adaptation of tubular water reabsorption in patients with CKD to parameterize TFR, and therefore TFR decreases less than proportionally in relation to GFR. Quantitatively, a set of tubular subsegment-specific adaptation factors (AF) was calculated based on reported mean urine formation of 0.62 mL/min (62% of healthy urine flow) in patients with CKD (n = 216) with a mean GFR of 10 mL/min (8.3% of healthy GFR). Physiologically and mathematically, the magnitude of adaptation of TFR in patients with CKD must be different in different tubular subsegments. The inflow of the first subsegment of the proximal tubule must equal GFR and as such must decline proportionally with GFR. Similarly, the outflow of the last subsegment of the collecting duct must match the observed urine flow in patients with CKD and hence decline much less than GFR. For example, at a GFR of 10 mL/min, the observed urine flow is 7.5-fold higher than what would be expected from the change in GFR. Based on these boundaries, the tubular subsegment-specific AF (AF) for each TFR was calculated.
after optimization (details shown in **Supplementary Material**) using Eq. 1:

\[
\text{Adaptation Factor}_{i}(\text{AF})_{i} = \min + \frac{\text{max} - \min}{1 + 10^{n(\log GFR - 1)}}
\]

where min and max represent the minimum and maximum adaptation capacities of 0 and 0.57, \(n\) is analogous to Hill coefficient (set as 1.80). TFR_{ij} represents the renal tubular flow rate entering each tubular subsegment (including bladder) in healthy (H) subjects (GFR = 120 mL/min) where \(i\) ranges from 1 to 12.

The calculated AF was incorporated into the subsegment-specific TFR at a given GFR (GFR\(_i\)) using Eq. 2 defined scalar:

\[
\text{Scalar}_{ij} = 1 - \left(1 - \frac{\text{GFR}_{i}}{120}\right) \times (1 - \text{AF}_{i})
\]

where GFR\(_i\) represents the specific GFR value (mL/min) of patients with CKD (\(j\) ranges from 5 to 120 mL/min), GFR/120 represents the remaining renal function in patients with CKD, and AF\(_i\) is as defined by Eq. 1. The TFR\(_{ij,\text{CKD}}\) for each tubular subsegment (\(i\) ranges from 1 to 12) at different CKD stages (\(j\) ranges from 5 mL/min to 120 mL/min) was calculated using Eq. 3:

\[
\text{TFR}_{ij,\text{CKD}} = \text{TFR}_{ij,H} \times \text{Scalar}_{ij}
\]

where TFR\(_{ij,H}\) represents renal tubular flow rate entering each tubular subsegment (including bladder) in healthy subjects and the Scalar\(_i\) is defined by Eq. 2. The full set of calculated AF values and representative TFR values are shown in **Table 1** and **Table S1** and the model file is included as **Supplementary Material**. Each GFR input value results in a different set of TFR values for all 12 subsegments, producing a unique CKD stage-specific kidney model that describes the diseased renal tubular system at that level of renal impairment.

To ensure model integrity and evaluate sensitivities of CL\(_r\) to drug properties and physiological parameters, five sets of local sensitivity analyses were conducted to evaluate the impact of drug permeability (\(P_{app} = 1–100 \times 10^{-6} \text{cm/s}\)), plasma unbound fraction (\(f_{u,p} = 0.1–1\)), peritubular renal blood flow (300–1,000 mL/min), unbound intrinsic basolateral uptake transport clearance (CL\(_{\text{uptake}} = 10–3,000 \text{mL/min}\)), and unbound intrinsic apical efflux transport clearance (CL\(_{\text{efflux}} = 10–3,000 \text{mL/min}\)) on simulated CL\(_r\) using both proportional and adaptive models across varying levels of renal impairment.

**Prediction of CL\(_r\) for 20 test compounds in varying CKD stages using proportional and adaptive kidney models**

To assess the performance of the proportional and adaptive models in predicting CL\(_r\) with declining GFR, the CL\(_r\) of 20 test compounds were simulated and compared with observed values. The 20 test compounds included three parent-metabolite pairs, six nonpermeable drugs, six highly permeable drugs, all of which lacked significant secretion (CL\(_r < 1.25 \times f_{u,p} \times \text{GFR}\), **Table S2**) and two renally secreted compounds. All the permeable nonsecreted drugs are highly renally reabsorbed (1-CL\(_r/ \text{f}_{u,p} \times \text{GFR}\) ~ 72% to 97%). The collected pK\(_a\) values, in vitro permeability data, \(f_{u,p}\), and observed CL\(_r\) values in

| **Model subsegment** | **Healthy (stage 1)** GFR = 120 mL/min | **Adaptation factor** | **Mild stage CKD (stage 2)** GFR = 80 mL/min | **Moderate stage CKD (stage 3)** GFR = 40 mL/min | **Severe stage CKD (stage 4)** GFR = 20 mL/min | **End-stage CKD (stage 5)** GFR = 5 mL/min |
|----------------------|----------------------------------------|-----------------------|---------------------------------------------|---------------------------------------------|---------------------------------------------|---------------------------------------------|
| Proximal tubule\(_A\) | 120                                    | 0                     | 80.00                                       | 80.00                                       | 40.00                                       | 40.00                                       |
| Proximal tubule\(_B\) | 94                                     | 0.0099                | 62.67                                       | 62.98                                       | 31.33                                       | 31.96                                       |
| Proximal tubule\(_C\) | 68                                     | 0.018                 | 45.33                                       | 45.73                                       | 22.67                                       | 23.46                                       |
| Loop of Henle\(_A\)  | 43                                     | 0.038                 | 28.67                                       | 29.22                                       | 14.33                                       | 15.44                                       |
| Loop of Henle\(_B\)  | 24                                     | 0.098                 | 16.00                                       | 16.78                                       | 8.00                                        | 9.56                                        |
| Distal tubule         | 24                                     | 0.098                 | 16.00                                       | 16.78                                       | 8.00                                        | 9.56                                        |
| Collecting duct\(_A\)| 11                                     | 0.26                  | 7.33                                        | 8.29                                        | 3.67                                        | 5.58                                        |
| Collecting duct\(_B\)| 9                                      | 0.31                  | 6.00                                        | 6.94                                        | 3.00                                        | 4.87                                        |
| Collecting duct\(_C\)| 7                                      | 0.37                  | 4.67                                        | 5.54                                        | 2.33                                        | 4.08                                        |
| Collecting duct\(_D\)| 5                                      | 0.44                  | 3.33                                        | 4.07                                        | 1.67                                        | 3.14                                        |
| Collecting duct\(_E\)| 3                                      | 0.51                  | 2.00                                        | 2.51                                        | 1.00                                        | 2.02                                        |
| Bladder (urine)       | 1                                      | 0.56                  | 0.67                                        | 0.85                                        | 0.33                                        | 0.71                                        |

**Table S1** Representative renal TFR for the proportional and adaptive models at different stages of CKD

The tubular subsegment-dependent adaptation factors (AF) were calculated according to Eq. 1 and implemented for the adaptive model. The renal TFRs were calculated using either proportional model (TFR\(_p\)) by direct scaling to GFR, as described in Methods, or using the adaptive model (TFR\(_A\)) according to Eqs. 2 and 3. The renal TFR (mL/min) shown here indicates the inflow rate of entering each renal subsegment, which equals the outflow rate exiting from the previous renal subsegment. The inflow rate of the first proximal tubule subsegment always equals the GFR. The outflow rate of the last subsegment of collecting duct always equals the inflow rate of bladder and the urine formation rate (as no reabsorption occurs within bladder). All flows are presented in mL/min.

CKD, chronic kidney disease; GFR, glomerular filtration rate; TFR, tubular flow rate.
healthy subjects and patients with CKD stages 4 and 5 are summarized in Table S2. Overall, the 20 test compounds had a wide range of pKₐ values (2.9–11.5), in vitro permeabilities (0.1–120 × 10⁻⁸ cm/s), fₚ,u (0.05–1), and CLₑ (0.42–599 mL/min in healthy subjects and 0.19–76.4 mL/min in patients with CKD stages 4 and 5), providing a robust dataset to verify the adaptive model and to differentiate the performance of the adaptive and proportional models.

The CLₑ of all 20 test compounds were simulated as described before using both proportional and adaptive models with GFR input value decreasing from 120 mL/min to 5 mL/min with a decrement of 1 mL/min. Permeability and fₛ,u were compound-specific input parameters and uptake/exflux transport clearances were set as 0 for all test compounds except para-aminohippuric acid (PAH) and memantine for which previously published transport clearance values were used. The simulated CLₑ using both proportional and adaptive models were compared with the observed values at different CKD stages. To quantitatively evaluate the performance of proportional and adaptive models, absolute fold-error (AFE; Eq. 4) between the simulated and observed mean CLₑ in patients with severe/end-stage (i.e., stages 4 and 5) CKD along with a twofold acceptance criterion was used.

\[
AFE = 10^{\log_{10}\left(\frac{\text{Simulated Mean}}{\text{Observed Mean}}\right)}
\]  
(4)

RESULTS

Development of proportional and adaptive kidney models for CKD

To predict CLₑ of drugs in patients with CKD, the proportional and adaptive models were developed and tested. In the proportional model, all TFR are reduced proportionally with GFR (Figure 1) resulting in a predicted urine flow that is reduced by the same percentage as GFR. For GFR of 10 mL/min (92% decrease from 120 mL/min), the proportional model predicts a urine flow of 0.083 mL/min, which is 87% lower than the observed urine flow of 0.6 mL/min. In contrast, in the adaptive model, the TFR values are reduced (Eq. 3) by scalars (Eq. 2) defined by the remaining GFR and the adaptation factors (Eq. 1), resulting in a urine flow of 0.6 mL/min (Figure 1) when GFR = 10 mL/min, matching the observed data. This suggests that the adaptive model successfully captures the physiologic changes in renal tubules of remaining functional nephrons in patients with CKD. Table 1 summarizes the representative TFR values predicted using both adaptive and proportional models at GFRs of 80, 40, 20, and 5 mL/min. At mild CKD (GFR = 80 mL/min), the difference of TFR values between proportional and adaptive models was small, with a maximum of 28% difference in the final urine flow. However, the difference in urine flow between proportional and adaptive models increased with declining GFR reaching an ~200% difference in moderate CKD (GFR = 40 mL/min), ~400% difference in severe CKD (GFR = 20 mL/min), and ~1,400% difference in end-stage CKD (GFR = 5 mL/min).

Sensitivity analyses of proportional model and novel adaptive kidney model for CKD

To further test which drug-specific or system-specific parameters impact the CLₑ predictions across varying levels of renal impairment, sensitivity analyses were conducted to examine the effects of Pₚ,u, fₛ,u, and peri-tubular renal blood flow. For the proportional model, the fold-decrease in CLₑ with decreasing GFR was unaffected by Pₚ,u (Figure 2a), whereas for the adaptive model the fold-decrease in CLₑ with decreasing GFR was highly sensitive to Pₚ,u (Figure 2b). Using the adaptive model, when GFR decreased by 96% to 5 mL/min, the simulated CLₑ for drugs with low permeability (e.g., 1 × 10⁻⁶ cm/s) also decreased by 96%, whereas the simulated CLₑ for drugs with high permeability (e.g., 100 × 10⁻⁶ cm/s) only decreased by 32%, resulting in considerable difference in the predicted effect of CKD on CLₑ (Figure 2b). Overall, the adaptive model predicts dramatically different effects of CKD on CLₑ than the proportional model for drugs with moderate to high permeability (> 10 × 10⁻⁶ cm/s) in severe/end-stage CKD (GFR < 30 mL/min; Figure 2c). As expected, fₛ,u correlated negatively with simulated CLₑ (Figure 2d,e), and had no impact on the difference in simulated CLₑ between proportional and adaptive models (Figure 2f). Further, peritubular renal blood flow had no effect on simulated CLₑ regardless of CKD stage in both proportional and adaptive models for nonsecreted drugs (Figure 2g–i).

Prediction of CLₑ for nonsecreted compounds in varying CKD stages and verification of the adaptive kidney model

To verify the adaptive model for different CKD stages, the CLₑ of 18 nonsecreted compounds at different CKD stages (GFR 120 mL/min to 5 mL/min) were predicted and compared with observed CLₑ. First, the CLₑ of all compounds were simulated in healthy subjects (GFR = 120 mL/min) based on reported fₛ,u and in vitro permeability (experimentally determined or optimized in healthy subjects) and all predicted CLₑ values were within twofold of the observed (Table S2), confirming satisfactory baseline simulation in healthy subjects and appropriate model parameterization. In healthy subjects, the adaptation was 0, therefore, both adaptive and proportional models resulted in identical simulation results when GFR input was 120 mL/min (Figures 3–5 and Figure S1). To evaluate the performance of adaptive and proportional models with decreasing GFR (advancing CKD), three pairs of permeable drugs and their respective nonpermeable glucuronide metabolites studied concomitantly were used as the first test set. For all three permeable drugs, the adaptive model successfully predicted the CLₑ across the entire range of GFRs considered (Figure 3a–c) whereas the proportional model only predicted the CLₑ acceptably in mild CKD where physiological changes are not drastic. In stages 4 and 5 CKD (Figure 3d–f), the calculated AFEₑ ranged from 1.03 to 1.28 using adaptive model whereas the calculated AFEₑ were all > 2 using the proportional model. This suggests successful verification of the adaptive model in predicting CLₑ at all stages of CKD, and inadequate performance of proportional...
Figure 1 Schematic presentation of the mechanistic kidney model structure, together with the corresponding renal tubular flow rate (TFR; in mL/min) for each individual tubular subsegment (a total of 12) of the model. Three sets of physiologically-based TFR shown here are for healthy subjects (glomerular filtration rate (GFR) 120 mL/min, in green) and for the representative patients with chronic kidney disease (CKD) who have residual GFR of 10 mL/min using the adaptive model (in red) and the proportional model (in blue). The dynamic physiologically-based mechanistic kidney model shown here is parameterized by 33 volume parameters, 22 surface area parameters, 12 peritubular renal blood flow parameters, 12 renal TFR parameters, 3 basolateral uptake clearance parameters, and 3 apical efflux clearance parameters to fully capture the disposition of drugs/metabolites between renal tubules, cells, and vasculature. CL, clearance.
model for severe/end-stage of CKD. For the nonpermeable glucuronide metabolites measured in the same studies, both adaptive and proportional models successfully predicted the CLr (Figure 3g–i) across varying levels of renal impairment, including CKD stages 4 and 5 (AFE 1.14–1.38, all < 2; Figure 3j–l).

For further verification, the adaptive and proportional models were evaluated using six nonpermeable drugs that are predominantly filtered without significant reabsorption or secretion (Figure 4) and six permeable drugs that are significantly renally reabsorbed (Figure 5). Both adaptive and proportional models successfully predicted the CLr for all 6 nonpermeable drugs across varying levels of renal impairment, including CKD stages 4 and 5 (Figure 4a–c,g–i), with calculated AFE_A of 1.00–1.76 (adaptive model) and AFE_p of 1.02–1.65, (proportional model; Figure 4d–f,j–l).
In addition, the ratio between AFE_A and AFE_P for individual drugs was within the 0.8 to 1.25 range for all nonpermeable test drugs, suggesting an equivalent performance between adaptive and proportional models for CL_r prediction for nonpermeable and nonsecreted compounds. In contrast, the performance of the adaptive model was considerably better than the proportional model (Figure 5) for advanced CKD stages for permeable compounds that undergo significant renal reabsorption (83–97%; Table S2). The adaptive model successfully (AFE_A = 1.05–1.73, all < 2) predicted the CL_r across varying levels of renal impairment, including stages 4 and 5 CKD (Figure 5), suggesting high confidence in the use of the adaptive model to predict CL_r in patients with CKD. In contrast, discrepancy between the proportional model-simulated CL_r and observed CL_r was pronounced as the GFR decreased (Figure 5). At stages 4 and 5 CKD, the proportional model dramatically underpredicted the CL_r (AFE_P = 2.61–7.35, all > 2) with 83% of drugs having AFE_P > 3 (Figure 5). This demonstrates alarming inappropriateness of applying proportional model (or proportional GFR scaling of CL_r) in patients with CKD for drugs with medium-to-high permeability.

Prediction of CL_r for secreted compounds in varying stages of CKD
To test whether the adaptive model could incorporate tubular secretion and predict impact of CKD on the CL_r of secreted compounds as well as in varying stages of CKD in addition to nonsecreted drugs. Sensitivity analyses and CL_r predictions for PAH and memantine in CKD were also done using the proportional model and are shown in Figure S2 and Figure S3.

DISCUSSION
PBPK modeling has been proposed as a promising tool to predict drug disposition in complicated and unknown scenarios, including patients with renal impairment, due to its mechanistic capability of integrating multiple complex interactions. Indeed, progress has been made in PBPK modeling of CKD effects on drug disposition, with special attention focused on altered hepatic metabolism and renal active secretion. In contrast, the effect of CKD on renal passive reabsorption has not been considered in existing CKD models. This study is the first to systematically explore the observed effects of CKD on highly renally reabsorbed drugs, and to investigate the adaptive physiological changes in CKD, to establish a PBPK modeling framework to predict CL_r through CKD progression. We expanded our previously developed and verified mechanistic kidney model to incorporate the physiologically-based tubular changes of reduced water reabsorption/increased TFR per remaining functional nephron across varying CKD stages. Incorporation of the adaptive changes in water reabsorption is necessary to address the significant discrepancy in urine flow between the proportional model-derived urine flow of 0.083 mL/min and the observed mean urine flow of 0.6 mL/min in patients with CKD with GFR = 10 mL/min. Our final physiologically-based adaptive model developed here is consistent with the observed urine flow in patients with CKD (i.e., urine flow = 0.6 mL/min when GFR = 10 mL/min), and successfully predicted CL_r of a wide variety of compounds at different stages of CKD using f_ulp and in vitro permeability without any optimization or empirical scaling.

Using 20 test compounds representing different degrees of permeability and extent of passive renal reabsorption, our analyses unequivocally show that passive reabsorption and hence renal clearance may not decrease proportionally with GFR in CKD (Figures 3–5) due to the adaptation in tubular water reabsorption. Instead, the permeable and renally reabsorbed drugs had much lower (by 261–735%) reduction of CL_r than the reduction of GFR (Figure 5), whereas nonpermeable and predominantly filtered drugs...
did not show such disproportionality (Figure 4). Our model, which considers physiologically-based tubular surface area and dynamic tubular flow, allows prediction of the extent of passive reabsorption based on $P_{app}$, conferring appropriate model sensitivity to $P_{app}$ that reflects the observed $CLr$ in humans. Further, only the adaptive model shown here that accounts for the tubular flow adaptation (decreased water reabsorption/increased TFR per
remaining functional nephron) due to CKD allowed simulation of the decrease in drug concentration gradient between intratubular filtrate and peritubular blood, and subsequently reduced passive reabsorption and higher CLr than expected from residual GFR alone. The proportional model or the empirical GFR scaling approach that assumes all renal handling pathways are reduced proportionally (linearly) with GFR was shown to be inappropriate for drugs with moderate-to-high permeability (Figure 5) and thus only applicable to low permeability, non-reabsorbed drugs (Figure 4).

Over the recent decade, PBPK modeling has been used to model CKD effects on renal elimination, primarily for highly secreted drugs that do not have significant permeability or reabsorption.
passive reabsorption. Although these models recovered the drug disposition in CKD, inconsistent scaling factors, such as relative activity factor (scalars ranging from 0.28 to 0.40) and proximal tubular cells per gram of kidney (scalars up to 15) have been applied, to allow the model to recover the observed data in healthy subjects and patients.
with CKD. This suggests low confidence on both in vitro-to-in vivo extrapolation of renal transport and understanding of CKD effect on renal drug handling. This is concerning as many renally secreted drugs, such as amphetamines, also have considerable permeability and reabsorption. For such drugs, the practice of empirically optimizing active secretion using data from healthy subjects and extrapolating to patients with CKD may result in erroneous parameter optimization and misleading prediction of unstudied scenarios, as recently demonstrated in the context of full-body PBPK modeling.43 As such, the adaptive model was entirely developed based on physiologic knowledge independent of drug molecules, and was collectively verified against 20 test compounds with different permeabilities throughout CKD stages without any optimization or empirical scaling for any of the test compounds. Our work demonstrates successful simulations of CL in CKD for a wide variety of drugs, establishing confidence on disease effect on renal passive reabsorption and laying a foundation for incorporating the impact of CKD on renal active secretion.

It is well-established that transporter-mediated renal active secretion is reduced in CKD due to declined number of functional nephrons and accumulating uremic solutes that may inhibit OAT1/3 activity. In this study, we used the developed and verified adaptive model to simulate CL for the two secreted test compounds, PAH and memantine, and showed successful CL prediction (AFE = 1.10–1.12; Figure 6) in varying stages of CKD. We demonstrated that our adaptive model can effectively incorporate specific transporter (OAT1/3 and OCT2) mediated renal secretion into CL prediction. Based on the simulation results, the novel adaptive model together with proportional decline in transporter-mediated secretion clearance adequately captures the changes in CL with progression of CKD for PAH and memantine. Uremic solutes have been shown to inhibit OAT1/3-mediated renal uptake but this is unlikely to impact CL prediction for PAH and memantine, as CL (and the vectorial secretion) of PAH is renal plasma flow limited and memantine is a substrate of OCT2, which is less affected by uremic solutes. Further studies on OAT1/3 and OCT2 activity and expression in CKD and clinical data on secreted

Figure 6 Sensitivity analyses of simulated renal clearance (CLr in mL/min) at multiple stages of chronic kidney disease (CKD) reflected by varying glomerular filtration rates (GFR in mL/min) using adaptive model (shown in yellow-red). (a) The sensitivity analyses of adaptive model-simulated CLr of neutral unbound permeable drugs (f_u,p = 1, P_app = 30 × 10^-6 cm/s) with a constant unbound intrinsic apical efflux transport clearance (CL_efflux = 150 mL/min) and different unbound intrinsic basolateral uptake transport clearances (CL_uptake = 10–3,000 mL/min) across a range of GFRs (5–120 mL/min). (b) The sensitivity analyses of adaptive model-simulated CLr of neutral unbound permeable drugs (f_u,p = 1, P_app = 30 × 10^-6 cm/s) with a constant unbound intrinsic basolateral uptake transport clearance (CL_uptake = 150 mL/min) and different unbound intrinsic apical efflux transport clearances (CL_efflux = 10–3,000 mL/min) across a range of GFRs (5–120 mL/min). The sensitivity analyses using proportional model and the comparison between the two models are shown in the Figure S2. (c, d) The simulations of CLr of para-amino-hippuric (PAH) and memantine in red curves, respectively, using adaptive model, at multiple stages of CKD, and comparison to the observed data (Table S2) shown in black open circles with calculated absolute folderror (AFE) shown in the insets. The simulation results for PAH and memantine using proportional model are shown in Figure S3.
drugs that have moderate-to-high permeability in CKD are warranted for future model refinement.

The approach described in this study also has several assumptions and limitations. Disease effect on the \( f_{u,p} \) was assumed to be insignificant, as the plasma albumin concentration is only 7% lower in mild and 16% lower in severe CKD than in healthy subjects,\(^{42}\) and alterations in \( f_{u,p} \) due to uremic solute displacement likely only apply to acidic compounds with a reported median (\( n = 16 \)) increase of 35% in \( f_{u,p} \) in severe CKD. For the 20 test compounds included here, only 3 are highly protein bound (Table S2) and digitoxin is the only acid, although the \( f_{u,p} \) of digitoxin is unchanged in patients with CKD.\(^{46}\) As illustrated by the sensitivity analyses (Figure 2), the small changes in \( f_{u,p} \) in CKD cannot explain the dramatic (mean = 458% (261–735%)) discrepancy between the observed CL\(_r\) and the expected CL\(_r\) from the proportional model for permeable drugs (Figure 5). Yet, the modeling framework established here could readily incorporate changes in \( f_{u,p} \) during CKD if observed/expected. Another challenge for model verification is the paucity of data on GFR, urine flow, and CL\(_r\) measurements for individual subjects in CKD studies. It would be ideal to compare simulated and observed CL\(_r\) for each individual subject with known GFR and urine flow. However, most clinical studies only report group mean CL\(_r\) for individual subjects in CKD studies. It would be ideal to compare simulated and observed CL\(_r\) for each individual subject with known GFR and urine flow. However, most clinical studies only report group mean CL\(_r\) for individual subjects in CKD studies. It would be ideal to compare simulated and observed CL\(_r\) for each individual subject with known GFR and urine flow.

Supporting Information. Supplementary information accompanies this paper on the CPT: Pharmacometrics & Systems Pharmacology website (www.psp-journal.com).

Funding. This work was supported by National Institutes of Health grant P01 DA032507. W.H. was supported by Warren G. Magnuson Scholarship and William E. Bradley Fellowship from the University of Washington, Seattle, WA.

Conflict of Interest. Both authors declared no competing interests for this work.

Author Contributions. W.H. and N.I. designed the research, performed the research, analyzed data, and wrote the manuscript.

1. Levey, A.S. & Coresh, J. Chronic kidney disease. Lancet 379, 165–180 (2012).
2. Zoccali, C. et al. The systemic nature of CKD. Nat. Rev. Nephrol. 13, 344–358 (2017).
3. Bricker, N.S., Morrin, P.A.F. & Kime, S.W. The pathologic physiology of chronic Bright’s disease. Am. J. Med. 28, 77–93 (1960).
4. KDIGO 2012 clinical practice guideline for the evaluation and management of chronic kidney disease. Am. J. Med. 129, 1–10 (2013).
5. Chapin, E. et al. Adverse safety events in chronic kidney disease: the frequency of ‘multiple hits’. Clin. J. Am. Soc. Nephrol. 5, 95–101 (2010).
6. Secora, A., Alexander, G.C., Ballew, S.H., Coresh, J. & Grams, M.E. Kidney function, polymyxin, and potentially inappropriate medication use in a community-based cohort of older adults. Drugs Aging 35, 735–750 (2018).
7. Schmidt, J.M. et al. Patterns of medication use and the burden of polypharmacy in patients with chronic kidney disease: the German Chronic Kidney Disease study. Clin. Kidney J. 12, 663–672 (2019).
8. Thomas, R., Kanse, A. & Sedor, J.R. Chronic kidney disease and its complications. Prim. Care 35, 329–344 (2008).
9. Hseu, C.H. et al. Identification and quantitative assessment of uremic solutes as inhibitors of renal organic anion transporters, OA1T1 and OA1T3. Mol. Pharm. 13, 3130–3140 (2016).
10. Chapron, A. et al. Does secretory clearance follow glomerular filtration rate in chronic kidney diseases? Reconsidering the intact nephron hypothesis. Clin. Transl. Sci. 10, 395–403 (2017).
11. Houghton, G., Dennis, M. & Gabriel, R. Pharmacokinetics of metronidazole in patients with varying degrees of renal failure. Br. J. Clin. Pharmacol. 19, 203–209 (1985).
12. Welling, P.G., Shaw, W.R., Uman, S.J., Tse, F.L. & Craig, W.A. Pharmacokinetics of minocycline in renal failure. Antimicrob. Agents Chemother. 8, 532–537 (1975).
13. Montay, G., Jacquot, C., Barriety, J. & Cunci, R. Pharmacokinetics of pefloxacin in renal insufficiency. Eur. J. Clin. Pharmacol. 29, 345–349 (1985).
14. Bricker, N.S., Morrin, P.A. & Kime, S.W. The pathologic physiology of chronic Bright’s disease. An expositions of the ‘intact nephron hypothesis’. Am. J. Med. 28, 77–93 (1960).
15. Blum, R.A. et al. Pharmacokinetics of gabapentin in subjects with various degrees of renal function. Clin. Pharmacol. Ther. 56, 154–159 (1994).
16. Blair, A.D., Burgess, E.D., Maxwell, B.M. & Cutler, R.E. Sotalol kinetics in renal insufficiency. Clin. Pharmacol. Ther. 29, 457–463 (1981).
17. US Food and Drug Administration (FDA). Development of Best Practices in Physiologically Based Pharmacokinetic Modeling to Support Clinical Pharmacology Regulatory Decision-Making (2019). <https://www.fda.gov/drugs/news-event>.
18. Grimstein, M. et al. Physiologically based pharmacokinetic modeling in regulatory science: an update from the U.S. Food and Drug Administration’s Office of Clinical Pharmacology. J. Pharm. Sci. 108, 21–25 (2019).
19. Hsueh, C.H. et al. Towards quantification of the effects of renal impairment and probenecid inhibition on kidney uptake and efflux transporters, using physiologically based pharmacokinetic modeling and simulations. Clin. Pharmacokinet. 53, 263–293 (2014).
20. Hseu, C.H. et al. PBPK modeling of the effect of reduced kidney function on the pharmacokinetics of drugs excreted renally by organic anion transporters. Clin. Pharmacol. Ther. 103, 485–492 (2018).
21. Yee, K.L. et al. Evaluation of model-based prediction of pharmacokinetics in the renal impairment population. J. Clin. Pharmacol. 50, 364–376 (2010).
22. Lindberg, A.A., Nilsson, L.H.S., Bucht, H. & Kallings, L.O. Concentration of chloramphenicol in the urine and blood in relation to renal function. Br. Med. J. 2, 724–728 (1966).
23. Sharpsone, P. The renal handling of trimethoprim and sulphaethoxazole in man. Postgrad. Med. J. 4 (suppl.), 38–42 (1969).
24. Fukuda, M. et al. Polynocturia in chronic kidney disease is related to natriuresis rather than to water diuresis. Nephrol. Dial. Transplant. 21, 2172–2177 (2006).
25. Yeh, B.P.Y. et al. Factors influencing sodium and water excretion in uremic man. Kidney Int. 7, 103–110 (1975).
26. Welling, P.G., Craig, W.A., Amidon, G.L. & Kunin, C.M. Pharmacokinetics of trimethoprim and sulfaethoxazole in normal subjects and in patients with renal failure. J. Infect. Dis. 128 (suppl. 3), S556–S566 (1973).
27. Nechita, A.M. et al. Determining factors of diuresis in chronic kidney disease patients initiating hemodialysis. J. Med. Life 8, 371–377 (2015).
28. Pennell, J.P. & Bourgeois, J.J. Adaptive changes of juxtamedullary glomerular filtration in the remnant kidney. Philos. Trans. R. Soc. B. 363, 131–135 (1981).
29. McNay, J.L. & Miyazaki, M. Regional increases in mass and flow during compensatory renal hypertrophy. Am. J. Physiol. 224, 219–222 (1973).
30. Biber, T.U.L. et al. A study by micropuncture and microdissection of acute renal damage in rats. Am. J. Med. 44, 684–705 (1968).
31. Huang, W. & Isoherranen, N. Development of a dynamic physiologically based mechanistic kidney model to predict renal clearance. CPT Pharmacometrics Syst. Pharmacol. 7, 593–602 (2018).
32. Rowland Yeo, K., Arabi, M., Jameel, M. & Rostami-Hodjegan, A. Modeling and predicting drug pharmacokinetics in patients with renal impairment. Expert Rev. Clin. Pharmacol. 4, 261–274 (2011).
33. Sayama, H., Takubo, H., Komura, H., Kogayu, M. & Iwaki, M. Application of a physiologically based pharmacokinetic model informed by a top-down approach for the prediction of pharmacokinetics in chronic kidney disease patients. AAPS J. 16, 1018–1028 (2014).
34. Grillo, J.A. et al. Utility of a physiologically-based pharmacokinetic (PBPK) modeling approach to quantitatively predict a complex drug-drug-disease interaction scenario for rivaroxaban during the drug review process: implications for clinical practice. BioPharm. Drug Dispos. 33, 99–110 (2012).
35. Zhao, P. et al. Evaluation of exposure change of nonrenally eliminated drugs in patients with chronic kidney disease using physiologically based pharmacokinetic modeling and simulation. J. Clin. Pharmacol. 52, 915–1085 (2012).
36. Tan, M.L. et al. Use of physiologically based pharmacokinetic modeling to evaluate the effect of chronic kidney disease on the disposition of hepatic CYP2C8 and OATP1B drug substrates. Clin. Pharmacol. Ther. 105, 719–729 (2019).

37. Huang, W., Nakano, M., Sager, J., Ragueneau-Majlessi, I. & Isoherranen, N. Physiologically based pharmacokinetic model of the CYP2D6 probe atomoxetine: extrapolation to special populations and drug–drug interactions. Drug Metab. Dispos. 45, 1156–1165 (2017).

38. Tan, M.L. et al. Effect of chronic kidney disease on nonrenal elimination pathways: a systematic assessment of CYP1A2, CYP2C8, CYP2C9, CYP2C19, and OATP. Clin. Pharmacol. Ther. 103, 854–867 (2018).

39. Yeung, C.K., Shen, D.D., Thummel, K.E. & Himmelfarb, J. Effects of chronic kidney disease and uremia on hepatic drug metabolism and transport. Kidney Int. 85, 522–528 (2014).

40. Bergman, A. et al. Effect of hepatic organic anion-transporting polypeptide 1B inhibition and chronic kidney disease on the pharmacokinetics of a liver-targeted glucokinase activator: a model-based evaluation. Clin. Pharmacol. Ther. 106, 792–802 (2019).

41. You, X. et al. Development of a physiologically based pharmacokinetic model for prediction of pramipexole pharmacokinetics in Parkinson’s disease patients with renal impairment. J. Clin. Pharmacol. 60, 999–1010 (2020).

42. Huang, W., Czuba, L.C. & Isoherranen, N. Mechanistic PBPK modeling of urine pH effect on renal and systemic disposition of methamphetamine and amphetamine. J. Pharmacol. Exp. Ther. 373, 466–501 (2020).

43. Huang, W. & Isoherranen, N. Sampling site has a critical impact on physiologically based pharmacokinetic modeling. J. Pharmacol. Exp. Ther. 372, 30–45 (2020).

44. Cheung, K.W.K. et al. The effect of uremic solutes on the organic cation transporter 2. J. Pharm. Sci. 106, 2551–2557 (2017).

45. Brater, D.C. Drug dosing in patients with impaired renal function. Clin. Pharmacol. Ther. 86, 483–489 (2009).

46. Kirch, W., Ohnhaus, E.E., Dylewicz, P., Pabst, J. & Storstein, L. Bioavailability and elimination of digoxin in patients with hepatorenal insufficiency. Am. Heart J. 111, 325–329 (1986).