Drug Therapy in Patients With Chronic Renal Failure
by Bertram Hartmann, David Czock, and Frieder Keller

SUMMARY

Background: Roughly 20% of patients in hospital have impaired kidney function. This is frequently overlooked because of the creatinine-blind range in which early stages of renal failure are often hidden. Chronic kidney disease is divided into 5 stages (CKD 1 to 5).

Methods: Selective literature search.

Results: Methotrexate, enoxaparin and metformin are examples of drugs that should no longer be prescribed if the glomerular filtration rate (GFR) is 60 mL/min or less. With antidiabetic (e.g. glibenclamide), cardiovascular (e.g. atenolol) or anticonvulsive (e.g. gabapentin) drugs, the advice is to use alternative preparations such as gliquidone, metoprolol or carbamazepine which are independent of kidney function. Drug dose adjustment should be considered with antimicrobial (e.g. ampicillin, cefazolin), antiviral (e.g. aciclovir, oseltamivir) and, most recently, also for half of all chemotherapeutic and cytotoxic drugs in patients with impaired kidney function (with e.g. cisplatin, for instance, but not with paclitaxel).

Conclusion: Decisions concerning drug dose adjustment must be based on the pharmacokinetics but this is an adequate prerequisite only in conjunction with the pharmacodynamics. There are two different dose adjustment rules: proportional dose reduction according to Luzius Dettli, and the half dosage rule according to Calvin Kunin. The latter leads to higher trough concentrations but is probably more efficient for anti-infective therapy.

Prevalence

The prevalence of renal insufficiency with a glomerular filtration rate (GFR) below 60 mL/min was estimated at 11% to 13% in the NHANES study.
Learning objectives
The purpose of this article is to demonstrate that, in renal insufficiency,

- certain drugs are particularly useful, and
- in principle, any drug can be given to any patient, as long as the dose is adjusted in accordance with the relevant pharmacokinetics and pharmacodynamics.

Special aspects of pharmacotherapy in renal insufficiency
If the GFR is below 60 mL/min, i.e., if the patient is in CKD stage 3 or higher, certain drugs should no longer be given, either because they tend to damage the kidneys or because they are insufficiently eliminated by poorly functioning kidneys and will therefore accumulate in the body and cause toxic side effects on other organs (4). Radiological contrast media containing iodine should be given to patients with renal insufficiency only after prophylactic hydration. In order to prevent contrast-medium toxicity, the authors additionally give 1600 mg of MESNA (methoxyethylsulfonate sodium) in 500 mL of normal saline 0.9% intravenously immediately before and during the contrast study (5).

CKD stage 4 and above, magnetic resonance imaging with gadolinium is no longer a risk-free alternative to iodinated media. In these stages of CKD, gadolinium preparations (linear ones more often than cyclic ones) can cause a serious condition known as nephrogenic systemic fibrosis. Thus, if the GFR is under 30 mL/min, gadolinium should be eliminated from the body with hemodialysis immediately after the MRI study (6).

ACE inhibitors, AT, blockers, and the renin inhibitor aliskiren have renally dependent pharmacokinetics, so that it suffices to give these drugs in half the normal dose to patients with renal insufficiency. In this situation, special attention must be paid to the risk of hyperkalemia. To avoid hyperkalemia, spironolactone should be given only in a low dose (no higher than 25 mg/day), or not at all, to patients with renal insufficiency; if it is given, then only in combination with a thiazide diuretic, furosemide, or both.

As there are some drugs that exert their effects independently of renal function, it is often useful to search for one of these drugs as a clinical alternative (Table 2). Cefepime should not be given from CKD stage 4 onward, i.e., when the GFR is below 30 mL/min, because there have been eight reported cases of lethal central nervous system intoxication occurring after five to seven days of parenteral treatment in dialysis-dependent patients (7). Drugs that should not, in general, be given to patients in CKD stage 3 or higher (GFR under 60 mL/min) include methotrexate, enoxaparin, metformin, and lithium (Table 3).

Methotrexate and enoxaparin accumulate in a deep compartment. After four to eight weeks of use, methotrexate can cause long-lasting bone marrow toxicity (8). With enoxaparin (unlike tinzaparin), eight days of treatment—even if the dose is adjusted to the GFR—can lead to hemorrhagic complications, which become more likely each day that the drug is given (9). Lithium is excreted through the kidneys and can induce toxic effects such as athetosis, renal diabetes insipidus, and renal failure (10); therefore, its use in CKD 3 or higher requires expert management (e2). The example of lithium also reminds us, however, that the discontinuation of a drug is always fraught with risk and must always be discussed with the prescribing physician (a psychiatrist, in this particular case). Once all the risks have been taken into consideration, it may turn out to be best to continue giving a nephrotoxic medication despite a low GFR.

Radiological contrast media
Patients with renal insufficiency should undergo prophylactic hydration before they are given radiological contrast media that contain iodine.

Gadolinium
Linear gadolinium preparations can bring about the clinical picture of nephrogenic systemic fibrosis when given to patients in CKD stage 4 or higher.
Metformin is considered to be an insulin sensitizer, and there is a current trend toward increasing use of this drug. Although lactic acidosis is a rare complication of metformin, and is even rarer when the dose is reduced (500 mg b.i.d.), it remains highly lethal when it occurs (11). As the patient’s renal function may be unstable, patients whose GFR is under 60 mL/min should be treated with less dangerous alternative drugs, such as the DPP-IV inhibitor sitagliptin (Table 2), whose dose should be halved if the GFR is under 30 mL/min. Half of therapeutically administered insulin is metabolized in the renal parenchyma (e3); thus, in renal insufficiency, the insulin effect often lasts longer, so that the patient needs less insulin overall. It is a common scenario for a diabetic patient to be admitted to hospital with hypoglycemia as the result of a worsening of renal function.

For patients whose GFR is below 60 mL/min, the proper use of analgesics remains an important issue. In the authors’ view, combined analgesic preparations are contraindicated in patients with renal insufficiency, as they may cause dependency. The kidneys’ major “enemies” are now the nonsteroidal anti-inflammatory drugs (NSAIDs), such as diclofenac, ibuprofen, and indomethacin, and the selective COX-2 inhibitors, which should only be given with monitoring of the serum creatinine level. In patients with preexisting renal damage, the use of NSAIDs can lead to further, permanent renal damage (12). The opioid drug pethidine should not be given to patients with renal insufficiency, because its renal-dependent metabolite, normeperidine, can cause cerebral convulsions (Table 3). Morphine can be given with clinical follow-up at short intervals, but the necessary dose of morphine is lower in the setting of renal insufficiency, and the physician must be aware that respiratory depression can arise in delayed fashion, after a few days of treatment, because of a gradual accumulation of the active metabolite M6-glucuronide. Fentanyl and levomethadone may be advantageous, as these drugs have no active metabolites and are independent of the GFR.

**The treatment of comorbidity in CKD**

The treatment of comorbidity in patients with chronic renal insufficiency becomes an important issue no later than when CKD stage 4 is reached, with a GDR below 30 mL/min. The evidence grade and recommendation strength for the various treatment options can be derived from Cochrane Reviews and published meta-analyses (Table 4).

### TABLE 2

| Drugs that do and do not depend on renal function (e2) |
|---------------------------------|---------------------------------|
| **Class** | **Dependent on renal function** | **Independent of renal function** |
| Analgesics | Morphine (M6-glucuronide), pethidine (norperidine) | Fentanyl, levomethadone |
| Antiarrhythmic drugs | Sotalol | Amiodarone |
| Antibiotics | Ciprofloxacin, levofloxacin | Moxifloxacin |
| Antidiabetic drugs | Glimepiride (hydroxy metabolite) | Gliclazide, gliclazole |
| | Nateglinide | Gliclazide, gliclazole |
| | Sitagliptine | |
| Anticonvulsants | Gabapentin, pregabalin, lamotrigine, levetiracetam | Carbamazepine, phenytoin, valproate |
| Antihypertensive drugs | Atenolol | Bisoprolol, carvedilol, metoprolol, propranolol |
| Cholesterol-lowering drugs | Bezafibrate, fenofibrate | Simvastatin, niacin |
| Drugs for gout and other rheumatological conditions | Methotrexate | Colchicine, hydroxychloroquine, leflunomide |
| Cardiovascular drugs | Digoxin | Digitoxin |
| Psychoactive drugs | Lithium, mirtazapine | Amitriptyline, citalopram (metabolites?), haloperidol, risperidone |
| Antiviral drugs | Acyclovir | Brivudine |
| Cytostatic drugs (20) | Actinomycin D, bleomycin, capetitabine, carboplatin, cisplatin, cyclophosphamide, doxorubicin, epirubicin, etoposide, gemcitabine (dFdU), ifosfamide, irinotecan, melphalan, methotrexate, oxaliplatin, topotecan | Anastrozole, docetaxel, doxorubicin-peg-liposomal, efotinib, fluorouracil, gefitinib, leuprolerin, megestrol, paclitaxel, tamoxifen, terozol, vincristine, trastuzumab |

Because renal insufficiency is associated with sodium retention (13), diuretic treatment is usually necessary; for this purpose, the best strategy is usually sequential nephron blockade with furosemide and hydrochlorothiazide.

The TREAT, CREATE, and CHOIR studies have shown that, in pre-dialysis patients, high-dose treatment with drugs that are intended to stimulate erythropoiesis can lead to strokes or tumor progression without protecting the kidneys (e4–e6). Therefore, the new

### The kidney’s main “enemies” are:

- Nonsteroidal anti-inflammatory drugs (NSAIDs) such as diclofenac, ibuprofen, and indomethacin
- Selective COX-2 inhibitors

### Opioids

The opioid drug pethidine should not be given to patients with renal insufficiency, because its renal-dependent metabolite, normeperidine, can cause cerebral convulsions.
**Pharmacokinetics**

Calvin Kunin, in the 1950’s, carried out the first systematic study of the prolongation of drug half-lives in the setting of progressively severe renal insufficiency ([e8]). Drugs for which this is true are excreted more slowly than normal and can, therefore, accumulate to toxic levels if given repeatedly. The dosage of these drugs must be adjusted according to the patient’s degree of renal insufficiency. Not only is every patient different, every drug is different as well. Nonetheless, there are simple rules of pharmacokinetics and pharmacodynamics that apply to every drug and every patient.

Friedrich Hartmut Dost described long ago the practical advantages of using a drug’s half-life ($T_{1/2}$) rather than its clearance ($e9$). The dominant half-life is the most important part of the concentration time curve with respect to the drug’s exerting its intended, primary effect (Luzius Dettli). The renally eliminated fraction of a drug ($f_{\text{ren}}$) is the key to predicting its pharmacokinetics in patients with renal insufficiency. For healthy subjects, $f_{\text{ren}}$ is calculated from the quantity of drug eliminated in the urine ($M_{\text{urine}}$) and the dose after recommendation is for a target hemoglobin value of 10 mg/dL ([14]). Before an erythropoietic hormone is given, iron should be given first. Iron is poorly taken up by patients with renal insufficiency when given orally, because of the elevated hepcidin level ([15]); it is thus best to give iron IV.

Large-scale observational studies have revealed that the additional administration of active vitamin D (e.g., calcitriol) in the predialysis phase is associated with lower mortality ([16]). It is important to monitor the calcium level and the calcium-phosphate product ($Ca \times PO_4$). Paricalcitol and cinacalcet should remain in reserve for the treatment of severe, clinically manifest hyperparathyroidism. Oral neutralization of acidosis with bicarbonate in the predialysis phase protects the bones, lessens malnutrition, and slows the progressive loss of renal function ([17]). A stepwise algorithm for the use of phosphate binders has been published ([e7]). The treatment begins with calcium acetate. If this is ineffective or poorly tolerated, sevelamer (or the newly available sevelamer carbonate) should be given instead as a powder. The authors use lanthanum and phosphate binders containing aluminum only for short times, and only in otherwise intractable cases.

### Table 3

<table>
<thead>
<tr>
<th>Class</th>
<th>Drug</th>
<th>Contraindicated or to be avoided if possible when:</th>
<th>Reason</th>
</tr>
</thead>
<tbody>
<tr>
<td>Analgesics</td>
<td>Pethidine</td>
<td>GFR &lt;60</td>
<td>Convulsions</td>
</tr>
<tr>
<td>Antibiotics</td>
<td>Cefepime</td>
<td>GFR &lt;30</td>
<td>CNS toxicity</td>
</tr>
<tr>
<td>Phase-prophylactic</td>
<td>Lithium</td>
<td>GFR &lt;60</td>
<td>Nephro- and neurotoxicity</td>
</tr>
<tr>
<td>Antidiabetic drugs</td>
<td>Glibenclamide, gimepiride</td>
<td>GFR &lt;60</td>
<td>Hypoglycemia</td>
</tr>
<tr>
<td></td>
<td>Metformin</td>
<td>GFR &lt;60</td>
<td>Lactic acidosis</td>
</tr>
<tr>
<td>Diuretics</td>
<td>Enoxaparin</td>
<td>GFR &lt;60</td>
<td>Risk of hemorrhage</td>
</tr>
<tr>
<td>Immune suppressants</td>
<td>Methotrexate</td>
<td>GFR &lt;60</td>
<td>Myelotoxicity</td>
</tr>
<tr>
<td>Radiological contrast media</td>
<td>Gadolinium</td>
<td>GFR &lt;30</td>
<td>Nephrogenic systemic fibrosis</td>
</tr>
<tr>
<td>LMW-heparin</td>
<td>Enoxaparin</td>
<td>GFR &lt;60</td>
<td></td>
</tr>
</tbody>
</table>

*1 Some of the Cochrane Reviews and Cochrane Meta-Analyses cited (e18, e20, e22) do not consider the more recent studies ([e4–e6, 16, 17]).

### Table 4

<table>
<thead>
<tr>
<th>Condition</th>
<th>Treatment</th>
<th>Evidence level and recommendation strength</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sodium retention</td>
<td>Diuretics</td>
<td>3A</td>
<td>(13, e17)</td>
</tr>
<tr>
<td>Target Hb 10 rather than 12–13 g/dL</td>
<td>Epo</td>
<td>1A</td>
<td>(14, e4–e6, e17)</td>
</tr>
<tr>
<td>Inadequate iron resorption</td>
<td>Iron IV</td>
<td>1A</td>
<td>(15, e19)</td>
</tr>
<tr>
<td>1α-hydroxylase deficiency</td>
<td>Calcitriol</td>
<td>2C</td>
<td>(15, e20)</td>
</tr>
<tr>
<td>Hyperparathyroidism</td>
<td>Paricalcitol</td>
<td>3C</td>
<td>(e7, e20)</td>
</tr>
<tr>
<td>Hyperparathyroidism</td>
<td>Cinacalcet</td>
<td>1B</td>
<td>(e7, e21)</td>
</tr>
<tr>
<td>Neutralization of acidosis</td>
<td>Bicarbonate</td>
<td>3B</td>
<td>(17, e22)</td>
</tr>
<tr>
<td>$Ca \times PO_4$-product</td>
<td>Sevelamer</td>
<td>2B</td>
<td>(e7, e23)</td>
</tr>
<tr>
<td>$Ca \times PO_4$-product</td>
<td>Lantham 3C</td>
<td>3C</td>
<td>(e7, e23)</td>
</tr>
</tbody>
</table>

*1 Some of the Cochrane Reviews and Cochrane Meta-Analyses cited (e18, e20, e22) do not consider the more recent studies ([e4–e6, 16, 17]).

### Problematic drugs whose use in patients with renal insufficiency is nephrologically contraindicated:

Pethidine, cefepime, lithium, glibenclamide, gimepiride, metformin, spironolactone, eplerenone, methotrexate, gadolinium, enoxaparin.

### Pharmacokinetics

The renally eliminated fraction of a drug is a key to predicting its pharmacokinetics in patients with renal insufficiency.
intravenous administration (D), or else from the half-life of the drug in patients with normal (T½norm) and failing (T½fail) renal function.

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\[ f_{\text{ren}} = \frac{M_{\text{urine}}}{D} = 1 - \frac{T_{1/2\text{norm}}}{T_{1/2\text{fail}}} \]

Luzius Dettli (e10) had the innovative idea of correlating the elimination rate constant, which is inversely proportional to the half-life \( (k_e = 0.693 / T_{1/2}) \), linearly with renal function (Figure 1). To estimate the expected half-life of a drug based on the patient’s individual GFR, we need to know the two extreme values of the half-life (T½norm und T½fail), which, in turn, tell us what fraction of the drug is normally eliminated renally (fren):

\[ T_{1/2} = \frac{T_{1/2\text{norm}}}{1 - f_{\text{ren}} \cdot \left(1 - \frac{\text{GFR}}{\text{GFR}_{\text{norm}}} \right)} \]

The renally eliminated fraction can also be derived from its counterpart, the non-renally eliminated fraction Q0 (fren = 1 – Q0), and can be obtained online on the “dosing” website (www.dosing.de). Like the serum creatinine level, the half-life has a hyperbolic dependence on renal function (Figure 2). There is a range of non-critical values for the half-life when the GFR is above 60 mL/min; this is analogous to the creatinine-blind range (18).

**Dose adjustment**

Many drugs must have their doses adjusted to renal function in patients with CKD stage 3 or higher, i.e., GFR lower than 60 mL/min. In the absence of dose adjustment, aciclovir and valaciclovir can be neurotoxic, ganciclovir and valganciclovir myelotoxic, foscavir nephrotoxic, and cidofovir both myelo- and nephrotoxic. Dose adjustment has proved to be outstandingly useful for these vitally indicated drugs. Further drugs whose doses need to be adjusted are oseltamivir, which is effective against the H1N1 virus, and intravenous zanamivir.

Drugs for which dose adjustment is often forgotten include the newer antiepileptic drugs, such as gabapentin (19), pregabalin, lamotrigine, and levetiracetam. For these drugs, half the normal dose or an even smaller amount will often suffice (Table 2).

### FIGURE 1

**Drug elimination and renal function.** According to Luzius Dettli, the elimination rate constant \( (ke = 0.693 / T_{1/2}) \) depends linearly on the glomerular filtration rate (GFR) (e10). The non-renal elimination rate constant \( (k_{enonren}) \) is calculated as the difference of the normal and renal elimination rate constants \( (k_{enonren} = ke - ke_{ren}) \). The drug dose (D) is adjusted to renal function proportionally to the elimination rate constant, and therefore inversely proportionally to the half-life (e10). D % norm, dose change expressed as a percentage of the normal dose.

### FIGURE 2

**Half-life and renal function.** The half-life (T½) depends in hyperbolic fashion on the glomerular filtration rate (GFR). The dependence is stronger with a higher renal elimination fraction \( (f_{\text{ren}}) \). There is, however, a range of non-critical values for the half-life when the GFR is above 60 mL/min; this is analogous to the creatinine-blind range (18).

**Dose adjustment**

For patients in CKD stage 3 or above whose GFR is below 60 mL/min, the dose of many drugs must be adjusted in accordance with renal function.

**Cytostatic agents that depend on renal function**

When a drug of this type is given to a dialysis patient, its dose must be reduced to 40% to 80% of the standard dose.
More and more commonly, cancer patients with renal insufficiency undergo treatment with cytostatic and other chemotherapeutic drugs. It is helpful to distinguish cytostatic agents that depend on, or are independent of, renal function (20). Those that depend on renal function must be given to dialysis patients in 40% to 80% of the standard dose (21). Those that are independent of renal function do not require dose adjustment for renal function, but rather for the patient’s general condition. Unlike the authors of an article on this topic that was published in 2007 (20), the authors of the current article consider gemcitabine to depend on renal function because of its active metabolite dFdU (Table 2). The dose of gemcitabine, platinum derivatives, or of cyclophosphamide need not be lowered if the patient undergoes hemodialysis on three consecutive days, one to twelve hours after the administration of either of these dialyzable drugs. For carboplatin and cisplatin, dialysis must begin one to two hours after the infusion in order to be effective. Etoposide and epirubicin are not dialyzable.

Cumulation kinetics implies that there are separate considerations for the starting dose and the maintenance dose, as Augsberger already pointed out with respect to the digitalis glycosides (e11). If the effect of the drug must have a rapid onset, then the treatment should begin with a normal starting dose, so that an effective drug concentration can be established at once. With antibiotics, in particular, one should never commit the capital error of forgetting the starting dose. One always begins with the normal dose as the starting dose; only the maintenance dose is adjusted to the patient’s renal function, as reflected by the drug’s half-life.

The second innovative idea of Luzius Dettli was to realize (e10) that the necessary dose adjustment could be calculated proportionally to the changed elimination rate (and thus inversely proportionally to the half-life) (Figure 1). Either the dose (D) of a drug must be reduced in inverse proportion to the half-life $T_{1/2}$ (the Dettli-1 rule), or else the interval (Tau) between doses must be prolonged proportionally to the half-life $T_{1/2}$ (the Dettli-2 rule). These two rules can also be applied in combination.

$$D = \frac{D_{norm}}{\frac{Tau}{T_{1/2 norm}}}$$

If, for example, the half-life of a drug is five times longer because of renal insufficiency, its dose should be reduced to 20% of the normal dose ($1/5 = 0.20$). If the dose is thus reciprocally lowered and the interval between doses is unchanged, then the area under the curve will be the same as in the normal case, but with higher trough values (Figure 3), which are what is customarily measured in drug monitoring. This can easily lead physicians unfamiliar with the relevant pharmacokinetics to draw false conclusions and then reduce the dose of the drug still further, e.g., in the case of vancomycin or gentamicin. In order to achieve effective levels in patients with renal failure, the trough values of vancomycin should lie between 10 and 15 mg/L, and those of gentamicin between 2 and 4 mg/L.

Dettli’s proportionality rule often implies the administration of absurdly low doses, or much too wide intervals between doses, in patients who are under treatment in intensive care units. In the case of ampicillin, for example, one would have to reduce the dose of ampicillin from 1000 mg every 8 hours to 100 mg every 8 hours, or perhaps 1000 mg every 80 hours. This illustrates the importance of a second dosing rule, the halving rule of Calvin Kunin (e12), which generally yields higher doses than Dettli’s rule and thus provides severely ill patients with a higher chance of effective treatment (Figure 3).
states that the starting dose should be the same as the normal dose ($D_{\text{start}} = D_{\text{norm}}$), and thereafter half the starting dose should be given at intervals equal to one half-life ($T = T_{1/2}$). The main benefit of this scheme is to achieve effective peak levels.

$$\frac{D}{T} = \frac{1}{2} \cdot \frac{D_{\text{start}}}{T_{1/2}}$$

Kunin’s halving rule makes it immediately clear that patients with life-threatening conditions generally do not need dose adjustment when the half-life of a drug is shorter than the dosing interval.

The halving rule results in peak drug levels that are the same as those in patients with normal renal function, but markedly higher trough levels; the area under the curve is greater than with the Dettli rule (Figure 3). This larger area under the curve can, however, be associated with more frequent adverse effects. The authors prefer to use the Kunin rule in the intensive care unit for pharmacodynamic reasons (4), and we have published corresponding dosing recommendations in tabular form (www.uni-ulm.de/nephrologie). Underdosing of antibiotics, in comparison to overdosing, can have much worse, indeed potentially fatal effects.

The importance of pharmacodynamics also becomes clear in the use of loop diuretics such as furosemide and torasemide. Starting in CKD stage 5, i.e., when the GFR is 15 mL/min or less, these drugs must be given in higher rather than lower doses to be effective.

**Pharmacodynamics**

Dose adjustment is a question not just of pharmacokinetics, but also of pharmacodynamics, i.e., of the effect that one intends to achieve. Pharmacodynamics involves a formulation of the relationship between the concentration of a drug and its effect, most commonly described in terms of the so-called sigmoid $E_{\text{max}}$ model. In this mode, the effect is a non-linear function of the concentration of the drug at its site of action (C) and of the concentration that produces a half-maximal effect ($C_{50}$). The parameter $E_{\text{max}}$ is associated with the maximal effect, and the Hill coefficient $H$ describes the degree of sigmoidicity (“S-shapedness”) of the relationship. When the Hill coefficient $H$ equals 1, the $E_{\text{max}}$ model yields the Michaelis-Menten equation.

$$E = \frac{E_{\text{max}}}{1 + (C_{50}/C)^H}$$

The examples of furosemide and torasemide show that pharmacodynamic parameters can also change in renal insufficiency. The required concentration of these drugs to achieve a half-maximal effect becomes higher in renal insufficiency ($C_{\text{50}}_{\text{fail}} > C_{\text{50}}_{\text{norm}}$); both must often be dosed 10 times higher in order to achieve an adequate diuretic effect.

The sigmoid $E_{\text{max}}$ model (Figure 4) involves the definition of a threshold concentration $C_{E_{05}}$ (18) at which only 5% of the maximal effect is achieved ($E_{05} = 0.05 \times E_{\text{max}}$).

$$C_{E_{05}} = C_{E_{50}} \cdot 19^{-1}$$

Similarly, the so-called ceiling concentration $C_{E_{95}}$ is that at which 95% of the maximal effect is achieved ($E_{95} = 0.95 \times E_{\text{max}}$).

$$C_{E_{95}} = C_{E_{50}} \cdot 19^{-1}$$

For a Hill coefficient of 1.0, the threshold concentration is only $1/19$ of that which produces a half-maximal effect, while the ceiling concentration is 19 times as high as that which produces a half-maximal effect (Figure 4). On the other hand, when the Hill coefficient equals 4.25 (for example), the threshold concentration is half that which produces a half-maximal effect, and the ceiling concentration is only twice the concentration that produces a half-maximal effect (Figure 4). The therapeutically effective range of a drug thus depends on its Hill coefficient: the higher the Hill

**Antibiotic effect**

Antibiotics whose effect depends on their concentration have a low Hill coefficient ($H<1.5$), while antibiotics whose effect is time-dependent have a high Hill coefficient ($H>1.5$).
coefficient, the narrower the desired range is, and the more closely it centers on CE_{50}.

Antibiotics provide a good illustration of the explanatory power of pharmacodynamics. Antibiotics can be classified into those whose effect depends on concentration, and those whose effect depends on time (22). Aminoglycosides such as gentamicin and fluoroquinolones such as levofloxacin are examples of antibiotics whose effect is concentration-dependent and for which the ceiling concentration CE_{95} should be reached to make the treatment effective—even if the treating physician often does not know the value of CE_{95} or CE_{50} with any precision. Beta-lactams, such as cefazolin, and antiviral drugs, such as valganciclovir, are examples of antibiotics whose effect is time-dependent and for which the concentration should not become any lower than the threshold concentration CE_{05}. These two classes of antibiotics, distinguished by the concentration- or time-dependency of their effects, differ in the height of their Hill coefficients (Figure 4): antibiotics of the former type have a low Hill coefficient (H<1.5), while those of the latter type have a high Hill coefficient (H>1.5) (22).

The starting dose is important for both types of antibiotic, those whose effect is concentration-dependent and those whose effect is time-dependent. After the starting dose is given, the important consideration for antibiotics of the former type is the drug concentration at the beginning of each dosing interval, while that for the latter type is the drug concentration at the end of each dosing interval. Thus, for drugs of the former type, such as ciprofloxacin or levofloxacin, there is less reason to reduce the dose (23); it would seem to be better to prolong the interval between doses for patients with renal insufficiency, while keeping the individual dose the same as usual (Figure 5). Further examples of drugs of this pharmacodynamic type are cytostatic agents such as cyclophosphamide, melphalan, and daunoblastin (18). For cytostatic agents with a narrow therapeutic window, careful consideration of pharmacodynamics is needed, because adverse effects may be correlated with the area under the curve and might arise even after the administration of a single dose.

On the other hand, for antibiotics whose effect depends on time, such as ceftazidime (a beta-lactam) and the virustatic agents, it is less important to achieve peak levels that are well above CE_{50}, but it is absolutely necessary to ensure that the trough level at the end of each dosing interval does not drop below CE_{05}, as this might lead to loss of the antimicrobial effect. It follows that a continuous infusion might be a better administration strategy for beta-lactams, as has been shown in the case of oxacillin (24). The same consideration often applies in antiviral HIV therapy: the trough levels must not drop below the therapeutically effective concentration, as this will allow the pathogen to replicate, leading to the danger that resistance will develop (25). For antibiotics whose effect depends on time, such as cefotaxime or aciclovir, it makes more sense to lower the dose rather than prolong the interval between doses, as is done in the case of ciprofloxacin or daptomycin.

**Conclusion**

In the field of gerontopharmacology, one often hears the principle “Start low, go slow,” which was formulated by Jerry Gurwitz and applies mainly to drugs that exert an effect in the central nervous system (e13). On the other hand, there is a much older principle that has been, to some extent, forgotten over the years: “Go fast, start high”—in the original German, “Sofort und hoch dosieren,” as stated by Paul Ehrlich in 1913 with regard
to anti-infective therapy (e14). This remains an important principle in nephropharmacology today, particularly for antibiotics and antiviral drugs, but also for oncological cytostatic drugs. “Go fast, start high” applies not only in the intensive care unit, but also in general medical practice. Above all, one must not make the mistake of dosing anti-infective or cytostatic drugs too low at the start, or of forgetting the starting dose entirely. This mistake should be avoided in all patients, including those with renal insufficiency.

**Conflict of interest statement**

Dr. Hartmann and Professor Keller state that they have received payment of travel expenses and research support from Wyeth and Novartis. PD Dr. Czock states that he has no conflict of interest as defined by the guidelines of the International Committee of Medical Journal Editors.

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**REFERENCES**


Please answer the following questions to participate in our certified Continuing Medical Education program. Only one answer is possible per question. Please select the answer that is most appropriate.

**Question 1**
Chronic renal insufficiency is divided into how many CKD stages?
- a) One
- b) Two
- c) Three
- d) Four
- e) Five

**Question 2**
According to epidemiological studies, what is the prevalence of renal insufficiency with a GFR below 60 mL/min?
- a) 5–10%
- b) 10–15%
- c) 15–20%
- d) 20–25%
- e) 25–30%

**Question 3**
With what glomerular filtration rate does it usually become necessary to adjust drug doses to renal function?
- a) GFR <15 mL/min
- b) GFR <30 mL/min
- c) GFR <60 mL/min
- d) GFR <90 mL/min
- e) GFR <120 mL/min

**Question 4**
Which of the following drugs can cause long-lasting bone marrow toxicity four to six weeks after it is given to a patient with renal insufficiency?
- a) Methotrexate
- b) Hydroxychloroquine
- c) Colchicine
- d) Leflunomide
- e) Levomethadone

**Question 5**
Which of the following drugs is unsuitable for the treatment of comorbidity in patients with renal insufficiency?
- a) Captopril
- b) Bicarbonate
- c) Calcitriol
- d) Diclofenac
- e) Phosphate binders

**Question 6**
Which of the following variables is linearly related to the GFR?
- a) Serum creatinine concentration
- b) Elimination rate constant
- c) Half-life
- d) Non-renally eliminated fraction Q₀
- e) Threshold concentration CE₀₅

**Question 7**
Which of the following anti-infective drugs exerts its effect in time-dependent fashion?
- a) Gentamicin
- b) Cefazolin
- c) Fluoroquinolone
- d) Levofloxacin
- e) Ciprofloxacin

**Question 8**
Which of the following drugs must be given at a higher dose than normal in the setting of renal insufficiency?
- a) Atenolol
- b) Enoxaparin
- c) Etoposide
- d) Furosemide
- e) Gabapentin

**Question 9**
The threshold concentration produces what percentage of the maximal effect?
- a) 5%
- b) 20%
- c) 35%
- d) 50%
- e) 75%

**Question 10**
How must the dosage of ciprofloxacin be adjusted in a patient with renal insufficiency if the half-life of the drug is doubled?
- a) The dose should be halved.
- b) The interval between doses should be doubled.
- c) The drug concentration should be monitored.
- d) The dose of the drug should be gradually increased.
- e) The duration of therapy should be doubled.
CONTINUING MEDICAL EDUCATION

Drug Therapy in Patients With Chronic Renal Failure

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eReferences

CONTINUING MEDICAL EDUCATION

Drug Therapy in Patients With Chronic Renal Failure

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Case Illustration

A 33-year-old woman is admitted to hospital in preparation for peritoneal dialysis and implantation of a Tenckhoff catheter. She suffers from chronic transplant failure 12 years after renal transplantation. She developed end-stage renal failure at age 19 because of focal segmental glomerulosclerosis.

10 months ago, she underwent a biopsy of the transplanted kidney, which revealed interstitial fibrosis with tubular atrophy and calcineurin-inhibitor damage. The serum creatinine was 1220 micromoles/L, corresponding to a GFR of 5.7 mL/min. There were no other important laboratory abnormalities. She categorically refused hemodialysis, the creation of a dialysis fistula, or the implantation of a Shaldon catheter.

Her general condition is surprisingly good; indeed, she is asymptomatic except for labial herpes (Figure). She weighs 59 kg and is 170 cm tall. The systolic blood pressure is 130 mmHg and the pulse 72/min. The heart, lungs, abdomen, and limbs are all unremarkable on examination. Her current medications are prednisolone, mycophenolate, and sirolimus.

The surgeons and anesthetists are ready to implant a Tenckhoff catheter, but the patient’s labial herpes must be treated first with an antiviral drug. She refuses IV drug therapy and is therefore given valacyclovir by mouth. The normal dosage is two 500-mg tablets taken three times daily.

What dose would you have given?
The authors gave valganciclovir in a reduced dose, a single 500-mg tablet taken twice daily. This adjusted dose was still twice as high as the dose recommended in our tables, namely, 500 mg once daily (4). Today, we would give brivudine instead, in a single daily dose of 125 mg, as this drug is converted outside the kidneys to its inactive metabolite bromovinyluracil.

Labial herpes resolved completely within 48 hours, whereupon the patient was transferred to the surgery ward and operated on under a brief general anesthetic the next day. Two days after surgery, we were called to see the patient because she could not speak and could only sit up with difficulty. We suspected a neurotoxic adverse effect of valaciclovir and reassured the patient that her condition would improve within 24 hours. Valaciclovir was stopped and her neurotoxic manifestations promptly resolved.